Factors controlling temporal variability of near-ground atmospheric ²²²Rn concentration over central Europe

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16 Abstract

Concentration of ²²²Rn in near-ground atmosphere has been measured quasi-continuously 17 from January 2005 to December 2009 at two continental sites in Europe: Heidelberg (south-18 19 west Germany) and Krakow (southern Poland). The atmosphere was sampled at ca. 30m and 20 m, respectively, above the local ground. Both stations were equipped with identical 20 instruments. Regular observations of ²²²Rn were supplemented by measurements of surface 21 fluxes of this gas in Krakow urban area, using two different approaches. The measured 22 concentrations of ²²²Rn varied at both sites in a wide range, from less than 2.0 Bg m⁻³ to 23 approximately 40 Bq m⁻³ in Krakow and 35 Bq m⁻³ in Heidelberg. The mean ²²²Rn content in 24 Krakow, when averaged over entire observation period, was 30% higher than in Heidelberg 25 $(5.86\pm0.09 \text{ Bg m}^{-3} \text{ and } 4.50\pm0.07 \text{ Bg m}^{-3}, \text{ respectively})$. Distinct seasonality of ²²²Rn signal 26 is visible in the obtained time series of ²²²Rn concentration, with higher values recorded 27 generally during late summer and autumn. The surface ²²²Rn fluxes measured in Krakow also 28 29 revealed a distinct seasonality, with broad maximum observed during summer and early

autumn and minimum during the winter. When averaged over 5-year observation period, the 1 night-time surface 222 Rn flux was equal 46.8±2.4 Bq m⁻² h⁻¹. Although the atmospheric 222 Rn 2 levels at Heidelberg and Krakow appeared to be controlled primarily by local factors, it was 3 possible to evaluate the "continental effect" in atmospheric ²²²Rn content between both sites, 4 related to gradual build-up of ²²²Rn concentration in the air masses travelling between 5 Heidelberg and Krakow. The mean value of this build-up was equal 0.78±0.12 Bq m³. The 6 measured minimum ²²²Rn concentrations at both sites and the difference between them was 7 8 interpreted in the framework of a simple box model coupled with HYSPLIT analysis of air mass trajectories. The best fit of experimental data was obtained for the mean ²²²Rn flux over 9 the European continent equal 52 Bq $m^{-2} h^{-1}$, the mean transport velocity of the air masses 10 within convective mixed layer of PBL on their route from the Atlantic coast to Heidelberg 11 and Krakow equal 3.5 m s⁻¹, the mean rate constant of ²²²Rn removal across the top of PBL 12 equal to the ²²²Rn decay constant and the mean height of the convective mixed layer equal 13 1600 m. 14

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16 **1** Introduction

Radon (²²²Rn) is an alpha-emitting radioactive inert gas with the half-life of 3.8 days. It is a 17 product of the decay of ²²⁶Ra which belongs to ²³⁸U-decay series. Uranium (²³⁸U) and its 18 decay product, ²²⁶Ra, are ubiquitous in the Earth's crust and in the soils. Radon is being 19 released into the pore space of the soils and diffuses into the atmosphere where it decays to 20 lead ²¹⁰Pb via a chain of intermediate decay products. Under specific conditions (heavy rain 21 events), ²²²Rn decay products sticking to aerosol particles can be washed-out from the 22 23 atmosphere, resulting in underestimation of the measured radon concentrations when the radon progeny method is employed. The flux of ²²²Rn into the atmosphere is controlled by the 24 source term (²²⁶Ra content in the soil and its vertical distribution), by physical properties of 25 the upper soil layer (mineral structure, porosity, water content) and to some extent by short-26 27 term variations of physical parameters characterizing the soil-atmosphere interface (mainly 28 atmospheric temperature and pressure) (e.g. Greeman and Rose, 1996; Levin et al., 2003, 29 Taguchi et al., 2011).

First measurements of atmospheric ²²²Rn were performed in the late 1920s (Wigand and Wenk, 1928). Recent summary by Zhang et al. (2011) identifies 41 stations worldwide where ²²²Rn was measured regularly for periods longer that one year (USA - 8; Europe - 19; Asia - 1 12; South America, Australia, Africa - 5; remote ocean and polar regions - 7). In Europe, the 2 earliest datasets originate from Paris (1955-1960) and Saclay (1956-1960) (Servant and 3 Tanaevsky, 1961). Two major categories of ²²²Rn detection techniques have been employed 4 in those studies: (i) various designs of ionization chambers measuring directly the alpha 5 particles of ²²²Rn, and (ii) indirect methods based on measurements of radon decay products.

Due to lack of important sinks apart of radioactive decay, ²²²Rn is an excellent tracer for 6 7 atmospheric processes. Nowadays, major applications of radon in atmospheric research 8 include: (i) tracing of horizontal air mass transport (e.g. Dörr et al., 1983; Gerasopoulos et al. 9 2005), (ii) investigating vertical mixing in the lower atmosphere (Williams et al., 2008, 10 Zahorowski et al., 2008), (iii) evaluation of atmospheric chemistry and transport models (e.g. 11 Jacob et al., 1997; Chevillard et al., 2002; Gupta et al., 2004; Bergamaschi et al., 2006; Zhang 12 et al., 2008), (iv) validation of the parameterisation schemes in numerical weather forecasting 13 models (Jacob et al., 1997), and (v) assessing surface emissions of major greenhouse gases 14 such as CO₂, N₂O and CH₄ (e.g. Schmidt et al., 1996, Schmidt et al., 2003; Levin et al., 1999; Levin et al., 2003; Biraud et al., 2000; Conen et al., 2002; van der Laan et al., 2009; van der 15 16 Laan et al., 2010, Wilson et al., 1997).

17 Here we present an in-depth evaluation of two five-year records of quasi-continuous nearground atmospheric ²²²Rn concentration measurements performed at two continental sites in 18 19 Europe: Heidelberg (south-west Germany) and Krakow (southern Poland). The records were 20 obtained using identical instruments located in similar settings (urban environment). The primary objective of our work was the identification of major factors controlling temporal 21 variability of atmospheric ²²²Rn concentration observed at the both sites at diurnal, synoptic 22 23 and seasonal time scales. This included, among others, the measurements of local surface fluxes of this gas in Krakow urban area using two different approaches. The second objective 24 of the study was linked with the specific location of the two ²²²Rn monitoring sites (similar 25 latitudinal position with different distance to the Atlantic coast, en-route of major directions 26 of air mass transport across Europe). It was aimed at quantification of the "continental effect" 27 associated with build-up of ²²²Rn in the atmosphere over the European continent and its 28 interpretation in the framework of a simple model. 29

1 2 Measurement sites

The measurement sites are located at the same latitudinal band (ca. 50°N), in a similar urban settling, with the distance to the Atlantic Ocean equal approximately 600 km for Heidelberg and 1600 km for Krakow. Both measurement sites were equipped with identical instruments (Levin et al., 2002) and the atmosphere was sampled at a comparable level. The meteorological parameters (wind speed, wind direction and temperature) were monitored at the same elevation at which the inlet systems of the instruments measuring radon concentration were installed.

9 Krakow (approx. 800 000 inhabitants) is located in southern Poland. Characteristic features of the local climate are generally weak winds and frequent atmospheric temperature 10 inversion situations, sometimes extending over several days. The average wind speed 11 calculated for the period 2005-2007 was 1.9 m s⁻¹. West and south-west direction of surface 12 winds prevails. Westerly circulation is generally connected with stronger winds (wind speeds 13 above 4 m s⁻¹). Periods characterized by low wind speeds (<1 m s⁻¹), favouring accumulation 14 of ²²²Rn in near-ground atmosphere, constituted 34% of the total time considered. Monthly 15 16 mean air temperature at the site reveals a distinct seasonal cycle, with summer maximum (July-August) reaching 19-24°C and winter minimum (January-February) between -5 and 17 18 $+2^{\circ}$ C. Monthly precipitation rates are more irregular, with a broad maximum during summer 19 and minimum during winter months. The radon monitoring site (50°04'N, 19°55'E, 220 m a.s.l.) was located on the campus of the AGH University of Science and Technology, situated 20 in the western sector of the city, bordering recreation and sports grounds. Air intake for ²²²Rn 21 measurements was located on the roof of the Faculty of Physics and Applied Computer 22 Science building, 20 meters above the local ground. The site where surface fluxes of ²²²Rn 23 24 were measured, was located on the premises of the Institute of Nuclear Physics, Polish Academy of Sciences, situated in the western outskirts of the city, approximately 3 km north-25 west from the location of atmospheric ²²²Rn measurements. The soil type at the chamber 26 location was Endogleyic Cambisol (IUSS, 2007), dominated by silty clay loam. The mean 27 concentration of ²²⁶Ra in the soil profile, the precursor of ²²²Rn, was equal 22±3 Bq/kg 28 29 (Mazur, 2008).

Heidelberg (approximately 130 000 inhabitants) is located in the upper Rhine valley, in south-west Germany. Monthly mean surface air temperatures vary between 1-3 °C during winter months and 18-22 °C during the summer. The local atmospheric circulation patterns in

Heidelberg are dominated by alternate north/south flow along the Rhine valley, but also by 1 frequent easterly winds from the Neckar valley (Levin et al., 1999). The average wind speed 2 calculated for the period 2005-2007 was 3.0 m s⁻¹ while low wind speed (<1 m s⁻¹) periods 3 constituted only 9.8% of the total time considered. In contrast to local, near-surface wind 4 5 direction, backward air trajectories calculated for Heidelberg site clearly demonstrate predominance of westerly air masses. Air inlet for ²²²Rn measurements was installed on the 6 roof of the Institute of Environmental Physics building (49°24'N, 8°42'E, 116 m a.s.l.), 30 m 7 8 above the local ground.

9

10 3 Methods

11 **3.1.** Measurements of ²²²Rn content in near-ground atmosphere

Regular measurements of ²²²Rn content in near-ground atmosphere were performed with the 12 aid of Radon Monitor. The instrument was developed at the Institute of Environmental 13 Physics, University of Heidelberg, Germany (Levin et al., 2002), and made available for this 14 study. The instrument measures specific activity of ²²²Rn in air through its daughter products. 15 16 The air is pumped with the constant flow rate through a glass filter (Whatman QM-A, 2.2 µm) which is placed directly over the surface-barrier detector measuring alpha particles emitted by 17 ²²²Rn and ²²⁰Rn daughter products deposited on the filter. The instrument records alpha decay 18 19 energy spectra accumulating in 30-min counting intervals. The spectra contain peaks representing decay products of 222 Rn (214 Po, $E_{\alpha} = 7.7$ MeV and 218 Po, $E_{\alpha} = 6$ MeV) as well as 20 220 Rn (212 Po, E_a = 8.8 MeV and 212 Bi, E_a=6.1 MeV). By energy discrimination and a 21 dedicated data evaluation protocol taking into account disequilibrium between daughter 22 products of ²²²Rn at the end of each counting interval, as well as empirically determined 23 disequilibrium between ²²²Rn gas and its daughter products in the atmosphere, the specific 24 activity of the ²²²Rn gas in air can be calculated. The mean disequilibrium has been 25 determined by parallel measurements of atmospheric ²²²Rn activity with an absolutely 26 27 calibrated slow-pulse ionization chamber and the Radon Monitor to 1/1.367, for the elevation of 20 meters above the local ground in Heidelberg (Levin et al., 2002). 28

1 3.2. Measurements of surface ²²²Rn fluxes in Krakow

Two different approaches were used to quantify the magnitude and temporal variability of surface fluxes of ²²²Rn into the local atmosphere: (i) night-time ²²²Rn fluxes were derived from measurements of atmospheric ²²²Rn content near the ground, combined with quasicontinuous measurements of the mixing layer height within the Planetary Boundary Layer (PBL) and modelling of vertical ²²²Rn profiles in the atmosphere using regional transport model, and (ii) point measurements of soil ²²²Rn fluxes were performed using specially designed exhalation chamber system connected to AlphaGUARD radon detector.

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10 **3.2.1** Sodar-assisted estimates of night-time ²²²Rn fluxes

11 **3.2.1.1 Theory**

During the day, with active vertical convection of air in the lower atmosphere, radon emitted from the soil is diluted in a large volume of mixing layer within PBL, leading to relatively low ²²²Rn concentrations observed close to the ground. During late afternoon, when the vertical gradient of air temperature changes the sign, drastic reduction of vertical mixing occurs. This process leads to accumulation of ²²²Rn in near-ground atmosphere during the night. The rate of nocturnal increase of ²²²Rn concentration is controlled by the mixing layer height according to the mass balance equation:

$$19 \qquad H\frac{dC_m}{dt} = F_{in} - F_{out} \tag{1}$$

20 where:

21 H - height of the mixing layer,

22 C_m - mean concentration of ²²²Rn within the mixing layer,

23 F_{in} - surface flux ²²²Rn,

24 F_{out} - flux of ²²²Rn associated with removal processes (horizontal and vertical transport, 25 radioactive decay). For nights with low wind speed (< 1 m s⁻¹) and the adopted 26 frequency of measurements, this term can be neglected.

During stable atmospheric conditions with low wind speeds, a distinct vertical gradient of 222 Rn concentration is established within the PBL. As the measurements of 222 Rn content are performed close to the surface, at the height of approximately 20 m, a correction factor *k* 1 relating the increase of the mean ²²²Rn concentration within the mixing layer (dC_m/dt) to the 2 increase of ²²²Rn concentration observed close to the ground (dC_{surf}/dt) should be introduced 3 to eq.(1):

$$4 F_{in} = \frac{H}{k} \frac{dC_{surf}}{dt} (2)$$

5 where:

 $6 \quad H$ - mixing layer height

7 k – correction factor

- 8 C_{surf} concentration of ²²²Rn at the adopted measurement height.
- 9

10 **3.2.1.2 Modelling**

The correction factor k was quantified using vertical profiles of 222 Rn simulated by EMEP 11 model. The Unified EMEP model (http://www.emep.int/) was developed at the Norwegian 12 13 Meteorological Institute under the EMEP programme. In this work, the Unified EMEP model version rv2_6_1 was used. The model is fully documented in Simpson et al. (2012). It 14 15 simulates the atmospheric transport and deposition of various trace compounds, as well as 16 photo-oxidants and particulate matter over Europe. The Unified EMEP model uses 3-hourly 17 meteorological data from PARallel Limited Area Model with the Polar Stereographic map 18 projection (PARLAM-PS), which is a dedicated version of the HIgh Resolution Limited Area 19 Model (HIRLAM) model for the use within EMEP. The model has been extensively validated 20 against measurements (e.g. Jeričević et al., 2010). In the framework of the presented study the model was adopted to calculate regional transport of ²²²Rn. The model domain (Fig. 1S in the 21 Supplement) covers Europe and the Atlantic Ocean with the grid size of 50×50 km and 20 22 23 layers in vertical, reaching up to 100 hPa.

The Unified EMEP model was used to simulate vertical profiles of ²²²Rn concentrations in the atmosphere for the location of Krakow for one full year (2005) with hourly time resolution, assuming spatially constant exhalation rate of 1 atom cm⁻² s⁻¹ over the continent. Model results corresponding to the periods starting at sunset and ending 6 hours later for each night were used. The ²²²Rn concentration at the measurement height (20 m above the ground.) was evaluated from the simulated profiles using exponential fit to the model data. The mean radon concentration in the mixing layer was calculated as the mean ²²²Rn content in the 1 lowest five layers of the model representing approximately the first 600 m of the troposphere,

2 weighted by the thickness of the corresponding layers:

3
$$c_m = \frac{\sum_{i=1}^{5} c_i h_i}{\sum_{i=1}^{5} h_i}$$
 (3)

4 where:

5 C_m – modelled mean concentration of ²²²Rn within the mixing layer,

6 C_i – mean ²²²Rn concentration in the layer *i* (model data),

7 h_i – thickness of the layer *i*.

8 In the next step, the growth rates of the mean ²²²Rn concentration within the mixing layer 9 (ΔC_m) and at 20 m above the ground (ΔC_{surf}) were calculated for each considered period using 10 linear regression fit to the data. Finally, *k* factor was calculated for each night:

11
$$k = \frac{\Delta c_{surf}}{\Delta c_m}$$
(4)

12 The monthly mean k values were calculated using the values of this parameter assigned for 13 each night, after a two-step selection procedure. In the first step, only the periods characterized by well-defined growth rates ΔC_m and ΔC_{surf} (R^2 >0.8) were selected. In the 14 second step, periods characterized by high wind speeds within the first layer of the model (v >15 3 m s⁻¹) were removed. The monthly mean k values calculated using the above-outlined 16 procedure are presented together with their uncertainties in Fig. 2S in the Supplement. The k17 18 value for January 2005 is missing because no single night in this month fulfilled the adopted 19 selection criteria. Since model results were available only for 2005, the monthly mean k20 values presented in Fig. 2S were further smoothed using CCGvu 4.40 routine (Thoning et al., 21 1989). The smoothed curve (heavy line in Fig. 2S) was then used in calculations of surface fluxes of 222 Rn (see section 3.2.1.4). 22

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24 **3.2.1.3** Measurements of mixing height using sodar

The mixing layer height, *H*, was monitored using VDS sodar (Version 3) built by the Krakow Branch of the Institute of Meteorology and Water Management. Detailed description of the sodar system can be found in Netzel et al. (1995) and Zimnoch et al. (2010). Sodar records were analyzed manually. Stability of the surface layer was identified through unique features of sodar echoes and its range was defined by determining the height of the observed structures

at the upper level where mixing processes still exist (Piringer and Joffre, 2005). The sodar 1 system was located inside a park complex, between the city centre and the industrial district, 2 at the distance of ca. 6 km east from the location where atmospheric ²²²Rn measurements 3 were performed. To examine possible influence of the distance between the sodar and ²²²Rn 4 measurement sites on the calculated ²²²Rn fluxes, two dedicated measurement campaigns 5 were performed. During the first campaign lasting one month, ²²²Rn measurements were 6 performed directly at the sodar site. Then, the sodar system was moved to the permanent 7 location of the ²²²Rn monitoring system. During both campaigns neither significant change in 8 9 the mixing layer height variability nor the range of calculated fluxes were observed (Zimnoch 10 et al., 2010).

11

12 **3.2.1.4 Calculations of night-time ²²²Rn fluxes**

Each period considered in the calculations of night-time ²²²Rn fluxes started typically at 13 14 sunset and ended 6 hours later. Average height of the mixing layer was calculated from hourly sodar data for each analyzed night. In building monthly mean night-time ²²²Rn fluxes, 15 individual fluxes calculated for each night using eq. (2) were subject to two-step data 16 selection algorithm. In the first step, only the nights for which the growth rate of the measured 17 ²²²Rn concentration was well-defined ($R^2 > 0.8$), were selected. In the second step, the nights 18 with the standard uncertainty of the mean mixing-layer height u(H) larger than 30 m, 19 20 calculated from hourly values of this parameter derived from the sodar measurements, were 21 removed from the remaining dataset. For some months, the adopted data selection procedure resulted in significant reduction of the number of available data, leading to relatively large 22 uncertainty of the corresponding monthly mean values of the calculated night-time ²²²Rn 23 24 fluxes. The number of data used for calculation of the mean for each month varied between 5 25 and 23 with the mean value of 13. The first step of data selection reduced the amount of data 26 by 34% and the second step by additional 11%. Sensitivity analysis showed that modification of R^2 and u(H) values used in the data selection by 10% results in reduction of the pool of 27 28 data suitable for calculation of monthly means by 8% and 6%, respectively.

1 **3.2.2** Measurements of surface ²²²Rn fluxes using chamber method

The sodar-assisted assessment of surface nigh-time ²²²Rn fluxes in Krakow agglomeration 2 3 was supplemented by one-year long measurements of soil radon fluxes using in-growth 4 chamber method. The fluxes were measured using specially designed, automatic exhalation 5 chamber, connected with AlphaGUARD radon monitor (Mazur, 2008). The parameters of the chamber were as follows: (i) flow rate - 0.3 liter min⁻¹, (ii) insertion depth - 6 cm, (iii) 6 7 diameter -21.6 cm. The air trapped inside the chamber was circulated in a closed circuit for 8 about 90 minutes and the concentration of radon accumulated in the chamber was recorded 9 every 10 minutes. The radon flux was determined from the slope of the straight line fitted to 10 individual readings (Mazur, 2008; Vaupotič et al., 2010). Special device was constructed which enabled automatic movement of the exhalation chamber, down to the ground for radon 11 flux measurement and up for ventilation of the system (Mazur, 2008). The ²²²Rn fluxes were 12 measured up to eight times per day. Only night-time measurements (up to 4 in total) were 13 used for comparison with sodar-assisted estimates of surface ²²²Rn flux representing night-14 time periods. The available raw data are presented on Fig. 3S in the Supplement. 15

16

17 **4.** Results and discussion

Time series of hourly mean concentrations of atmospheric ²²²Rn obtained at Heidelberg and 18 Krakow between January 2005 and December 2009 are shown in Fig. 1. The gaps in time 19 series of ²²²Rn content from June till December 2007 in Krakow and in June-July 2006 in 20 Heidelberg were caused by technical problems with Radon Monitors. It is apparent from Fig. 21 1 that 222 Rn content varies at both sites in a relatively wide range, from less than 2 Bg m⁻³ to 22 approximately 40 Bq m⁻³ in Krakow and 35 Bq m⁻³ in Heidelberg. The mean ²²²Rn 23 concentration in Krakow averaged over entire observation period (5.86 ± 0.09 Bg m⁻³), is 30% 24 higher when compared to Heidelberg (4.50 ± 0.07 Bq m⁻³ - cf. Table 1). 25

26

27 **4.1** Diurnal changes of ²²²Rn content in near-ground atmosphere

Diurnal changes of ²²²Rn concentration in near-ground atmosphere over Krakow and Heidelberg, averaged separately for each hour of the day over entire observation period (January 2005 - December 2009) and for each season (spring, summer, autumn, winter), are summarized in Fig. 2a and b. Irrespectively of season, the measured ²²²Rn contents at both sites reveal characteristic behaviour, with elevated concentrations during night hours and reduced concentrations during mid-day. However, the shape and amplitude of daily changes of ²²²Rn content vary significantly with the season and the observation site.

During winter months (December-February), daily variations of ²²²Rn are remarkably 5 similar at both locations. The average 222 Rn content for that period is equal 5.65±0.17 Bq m⁻³ 6 at Krakow and 5.19±0.19 Bq m⁻³ at Heidelberg, with peak-to-peak amplitude of diurnal 7 changes reaching 1.5 Bq m⁻³ at both sites (cf. Fig. 2 and Table 1). Daily minima are shallow 8 9 and reduced in duration. During spring and summer months (March-May and June-August, respectively) the peak-to-peak amplitude of daily changes of ²²²Rn concentration increases 10 significantly. This increase is particularly well-pronounced in Krakow (the amplitudes reach 11 4.7 and 7.0 Bq m^{-3} , respectively) when compared to Heidelberg (2.5 and 3.5 Bq m^{-3} . 12 13 respectively). At the same time, broader daily minima are observed at both sites, reflecting 14 growing role of vertical mixing within the PBL, driven by longer exposure of the surface to 15 sunlight. Most pronounced differences between both sites are observed during autumn months (September-November). In Krakow, the maxima of atmospheric ²²²Rn content occur usually 16 in the early-morning hours (ca. 4-6 a.m. UTC) and reach 12 Bq m⁻³, followed by a distinct 17 minimum of 5 Bq m⁻³ recorded usually between 1 and 2 p.m. UTC. In Heidelberg, the early 18 morning maxima reach only 8 Bq m⁻³, while the minima are maintained at approximately the 19 same level as in Krakow (ca. 5 Bg m^{-3}). 20

Diurnal variations of ²²²Rn content at both monitoring sites, averaged over entire 21 observation period, are shown in Fig. 2c. It is apparent that higher peak-to-peak amplitude of 22 daily ²²²Rn variations in Krakow is primarily due to higher build-up of ²²²Rn during night 23 hours at this site when compared to Heidelberg. The nocturnal build-up of radon in Krakow 24 25 starts and ends up approximately 2 hours earlier when compared to Heidelberg (at ca. 3 p.m. 26 and 4 a.m. UTC, respectively). This difference may partly stem from the fact that while both 27 stations are located in the same time zone, the true solar time is shifted by approximately one 28 hour.

The source strength of ²²²Rn in the soil and the upward transport of this gas into the atmosphere does not vary significantly on hourly time scale, except of frontal situations with fast changes of atmospheric pressure or during prolonged rainfall events. Therefore, diurnal variations of the concentration of this gas in near-ground atmosphere presented in Fig. 2 1 primarily reflect the changes in stability of the lower troposphere and the resulting intensity of

- 2 vertical mixing.
- 3

4 4.2 Synoptic time-scale changes of ²²²Rn content in near-ground atmosphere

5 While the amplitude of diurnal changes of ²²²Rn content is controlled primarily by the 6 intensity of vertical mixing, its variations on synoptic time scale (days to weeks) should be 7 also a function of the origin of air masses passing through the given measurement site. One 8 example illustrating the influence of synoptic-scale phenomena on the atmospheric ²²²Rn 9 concentrations measured in Krakow is discussed below.

Temporal evolution of atmospheric ²²²Rn content in Krakow during the period from 10 March 1 to March 13, 2005, is shown in Fig. 3 in relation to wind speed, precipitation rates 11 12 and 96-hours backward trajectories of air masses passing the measurement location. Between March 1 and 4, the measured ²²²Rn content revealed strong diurnal variations (Fig. 3a), with 13 the maximum 222 Rn concentrations reaching 16 Bg m⁻³, superimposed on the growing trend of 14 mean daily concentration of this gas (Fig. 3b). Air masses passing the sampling site during 15 this period originated in north-central Europe (Fig. 3c). Very low wind speed facilitated 16 gradual build-up of night-time ²²²Rn in the local atmosphere during these days, as well as an 17 18 increase of day-time background level caused by the increase in "continentality" of the air masses. From March 5 onwards, the course of atmospheric ²²²Rn content has changed 19 radically; diurnal variations almost disappeared (Fig. 3a) while the average daily ²²²Rn 20 content dropped from 9.7 Bg m⁻³ for March 4 to 5.2 Bg m⁻³ for March 5 and 3.3 Bg m⁻³ for 21 22 March 6. This substantial change was clearly linked to change of circulation and sharp increase of wind speed (Fig. 3b,c). Between March 5 and 7 the sampling location was under 23 influence of air masses originating over Russia and Estonia, passing westward at relatively 24 25 high elevation over the Baltic Sea and then turning south-east in the direction of Poland. Between March 8 and 12 the sampling location was under the influence of maritime air 26 27 masses originating over the Arctic Sea and travelling southward at very high elevation with 28 high speed. The rainfall occurring in Krakow during this period (between 0.5 and 2.0 mm) was associated with frontal situation (drop of atmospheric pressure from 1000 hPa at March 8 29 to 965 hPa three days later). During the second part of this period, small increase of ²²²Rn 30 concentration correlated with significant reduction of the elevation of trajectories arriving in 31 32 Krakow was observed.

1 The data shown in Fig. 3 clearly demonstrate a strong link between the measured ²²²Rn 2 content in near-ground continental atmosphere and weather-related phenomena such as 3 history of air masses and wind speed. Concentrations of atmospheric ²²²Rn may change 4 significantly on synoptic time scale in response to these factors.

5

6 4.3 Seasonal variations of ²²²Rn content in near-ground atmosphere

Figure 4 shows monthly means of ²²²Rn content in near-ground atmosphere, as observed in Krakow and Heidelberg between January 2005 and December 2009. The medians reveal distinct seasonal trend, with a broad minimum of ²²²Rn content in spring and summer and maximum in autumn months. Monthly maxima are significantly higher in Krakow, reaching 25 Bq m⁻³, compared to 20 Bq m⁻³ in Heidelberg. Monthly minima are comparable at both locations.

13

14 **4.4 Surface fluxes of ²²²Rn in Krakow**

The monthly mean night-time fluxes of ²²²Rn in Krakow urban area, derived from 15 atmospheric ²²²Rn observations and sodar measurements of the mixing layer height using the 16 17 methodology outlined in section 3.2.1 are presented in Fig. 5. They cover the period: June 18 2004 - May 2009. Error bars indicate standard uncertainties of the monthly mean values. Blue 19 bars represent the number of nights used for calculation of the monthly means. As seen in Fig. 5, high night-time ²²²Rn fluxes were obtained for autumn months (September, October, 20 November) reaching 80-90 Bq $m^{-2} h^{-1}$. When averaged over entire observation period, the 21 mean 222 Rn night-time flux obtained for Krakow urban area is equal 46.8±2.4 Bg m⁻² h⁻¹. This 22 values coincides with approximately 46 Bq m⁻² h⁻¹ derived for the location of Krakow from 23 the map of 222 Rn flux in Europe obtained from terrestrial γ -dose rate data (Szegvary et al., 24 25 2009).

Direct measurements ²²²Rn exhalation rates from the soil using the methodology described in section 3.2.2 were performed over one-year period, from September 2005 till September 2006. To allow direct comparison with the night-time ²²²Rn fluxes derived from measurements of atmospheric ²²²Rn content and observations of the mixing layer height, only measurements performed during the days for which sodar-assisted ²²²Rn fluxes were available, were considered. The monthly mean ²²²Rn fluxes derived using both approaches

are shown in Fig. 6. The mean 222 Rn flux derived from chamber measurements (50.3±8.4 Bq 1 $m^{-2} h^{-1}$) agrees very well with sodar-assisted estimate of this flux (50.3±3.4 Bq m⁻² h⁻¹). A 2 large discrepancy is apparent for February and March 2006, for which the chamber 3 measurements differ from sodar-assisted estimates of the ²²²Rn flux by a factor of ten. The 4 reasons for such large difference remain unclear. It may stem from the fact that frozen soil 5 and snow cover observed at the measurement site in February and first half of March could 6 partly block the ²²²Rn flux from the soil, resulting in low readings of the chamber method, 7 whereas the sodar-assisted estimates of the ²²²Rn flux are spatial averages over the footprint 8 area of atmospheric ²²²Rn measurements which is in the order of several square kilometres. 9

10

4.5 Factors controlling seasonality of ²²²Rn content in near-ground atmosphere over central Europe

Distinct seasonality of atmospheric ²²²Rn concentrations apparent in the data presented in 13 Figs. 1 and 4 may have its roots in several processes: (i) it may reflect seasonal bias in the 14 15 origin of air masses arriving at the given measurement location, e.g. prevalence of maritime air masses with low ²²²Rn during summer and of continental air masses with higher average 16 17 ²²²Rn content during autumn and winter, (ii) it may reflect seasonal bias in the stability of the 18 lower atmosphere, with more frequent, prolonged inversion periods during autumn and winter months when compared to summer, and, finally (iii) it may reflect seasonality in the source 19 term i.e. seasonally varying ²²²Rn exhalation rates from the soil. All these three factors may 20 21 act together and they are examined in some detail below.

22 Backward trajectory analysis has been performed for Krakow and Heidelberg for the 23 period from January 2005 till December 2009 using HYSPLIT model (Draxler and Rolph, 24 2011). 96-hours backward trajectories were calculated for every hour within this time period. 25 Multiple trajectories were displayed by creating an arbitrary grid over the computational 26 domain, counting number of trajectories over each grid point (without multiple intersections of the same trajectory) and dividing it by the total number of trajectories. The plotting routine 27 28 was used (Rolph, 2011) to display the spatial distribution of trajectory densities over the 29 computational domain. The grid resolution has been set to one by one degree. The results 30 were averaged separately for each season and each site and are presented in Fig. 7. Yellow 31 colour represents grid points where more that 10% of all trajectories passed through. Blue, 32 green and cyan colours represent trajectory density range between 5 and 10%, 2 and 5% and 1

and 2%, respectively. The trajectory density plots shown in Fig. 7 are supplemented by statistical analysis of trajectory height. The distribution of the mean and the maximum trajectory height, separately for each season and each monitoring site, is presented in Fig. 4S. The characteristic feature of these distributions is generally lower height of trajectories arriving in Krakow when compared to Heidelberg. Also, the largest percentage of trajectories arriving in Krakow typically falls into the lowermost range of heights, whereas in Heidelberg a distinct maxima of the distributions are observed.

8 It is apparent from Fig. 7 that only minor seasonal variations occur in the spatial 9 extension and shape of the trajectory density distribution maps for 94-hours backward 10 trajectories arriving in Krakow and Heidelberg. The contours are generally skewed towards 11 W-E axis, reflecting dominance of westerly circulation. There are three distinct differences 12 between Heidelberg and Krakow: (i) Heidelberg receives significantly higher proportion of 13 maritime air masses originating over North Atlantic and arriving at the site within 96 hours. 14 when compared to Krakow, (ii) the surface area of the 1% contour map is larger for Heidelberg that for Krakow, indicating generally higher transport velocities and larger spatial 15 16 extensions of 96-hour backward trajectories for this site, and (iii) the maximum height of 17 trajectories is approximately three times higher for Heidelberg when compared to Krakow, indicating transport of air masses being less influenced by surface ²²²Rn sources. 18

19 In order to better understand the factors controlling the observed seasonality of atmospheric ²²²Rn levels at Krakow and Heidelberg, the available data averaged separately for 20 21 each month and for entire observation period were compared with other on-site parameters 22 which may influence this apparent seasonality. Figure 8a shows mean monthly distribution of ²²²Rn content in Krakow near-ground atmosphere, presented in the form of box-and-whisker 23 plot for the entire observation period from January 2005 till December 2009. The long-term 24 monthly mean ²²²Rn content shows a broad maximum in September-October-November. The 25 long-term monthly minima of ²²²Rn, as observed in Krakow and Heidelberg, are shown in 26 Fig. 8b. Within the quoted uncertainties of the mean values they are indistinguishable at both 27 sites in March, April and May. While the long-term monthly ²²²Rn minima are practically 28 constant from March till August in Heidelberg, in Krakow they start to increase already in 29 30 June. The long-term monthly minima peak in October and November, for Heidelberg and 31 Krakow, respectively.

32 Distinct seasonality observed in atmospheric 222 Rn concentration at both sites could be 33 also linked to the seasonality of 222 Rn emissions from the ground. Monthly mean night-time

²²²Rn fluxes, as estimated for Krakow and averaged over entire observation period (cf. Fig. 5), 1 2 are shown in Fig. 8c. Figure 8d shows the monthly means of ²²²Rn fluxes derived from chamber measurements performed in Krakow and averaged over the period from September 3 2005 till September 2006. The monthly night-time ²²²Rn fluxes (Fig. 8c) reveal a broad 4 5 maximum during summer and early autumn (June-October). Although the mean monthly ²²²Rn fluxes derived from chamber measurements are more variable, they also reveal a broad 6 7 maximum during summer and early autumn. Comparison of the amplitudes of seasonal 8 variation presented on Figs.8a and b with Figs.8c and d shows that the apparent seasonality in ²²²Rn emissions from the ground cannot fully account for the observed seasonality of the 9 ²²²Rn concentrations in near-ground atmosphere. Also other factors should be involved. 10

The surface ²²²Rn fluxes estimated in the framework of this study reveal distinct 11 seasonality, with a broad maximum during summer and early autumn and minimum during 12 13 winter. Theoretical considerations (e.g. Nazaroff, 1992, Sasaki et al., 2004) as well as large 14 body of experimental data (e.g. Rogers and Nielson, 1991; Greeman and Rose, 1996; Levin et al., 2002; Papachristodoulou et al., 2007; Sakoda et al., 2010) suggest that free pore space in 15 the soil available for diffusion-controlled transport of gaseous ²²²Rn exerts primary control 16 over ²²²Rn exhalation rates. This parameter can be approximated by the volumetric water 17 content in the soil profile. The maps of soil moisture available for the European continent 18 19 (http://edo.jrc.ec.europa.eu) reveal generally lower values of this parameter during summer and autumn when compared to winter and spring, for large parts of continental Europe. This 20 suggest that distinct seasonality of ²²²Rn exhalation rates, as observed in this study, may have 21 22 its origin in seasonal changes of moisture load in the upper soil level.

23 Monthly means of surface air temperature and the amounts of monthly precipitation, both 24 recorded in Krakow and averaged over the period from January 2005 till December 2009 are 25 shown in Fig. 8e. While the air temperature reveals distinct seasonal changes (the difference 26 between coldest and warmest month is larger than 20 °C), monthly precipitation data are more variable and the seasonality is less marked. Comparison of local air and soil temperatures 27 (four different depths) in the vicinity of ²²²Rn concentration measurements in Krakow is 28 shown in Fig. 8f. As seen in Fig. 8f, during autumn and winter the soil is warmer than air, 29 thus making the soil air unstable. This may facilitate transport of ²²²Rn from the soil to the 30 31 atmosphere (Schubert and Schulz, 2002).

The last two graphs in Fig. 8 (Fig. 8g and h) show seasonal variations of two parameters characterizing stability of the lower atmosphere in Krakow: (i) the wind speed, and (ii) the

1 duration of inversion episodes. In Fig. 8g monthly means of the percentage of calm periods 2 (wind speed below 1 m s⁻¹), averaged over entire observation period (January 2005 -December 2009), is shown. It is apparent that percentage of periods with wind speed below 3 one m s⁻¹ is highest in Krakow during autumn and winter. Also, the duration of the inversion 4 periods is highest during that time (October-February), contributing to enhanced stability of 5 the lower atmosphere. Although the data presented in Fig. 8h refer to another 5-year period 6 (January 1994 - December 1999), it is assumed that this specific feature of the local 7 8 atmosphere in Krakow, when averaged over 5-year period, is valid also for the period considered in this study (January 2005 - December 2009). 9

10

11 **4.6** Assessment of ²²²Rn build-up in the atmosphere over the European

12 continent

Fluxes of ²²²Rn over the ocean are two to three orders of magnitude smaller than those over 13 the continents (Schery and Huang, 2004). Consequently, maritime air masses entering the 14 continent will have very low ²²²Rn content and will be gradually laden with ²²²Rn until new 15 equilibrium is reached. Since both ²²²Rn observation sites discussed in this study are located 16 17 in the same latitudinal band (ca. 50°N) and are exposed to westerly circulation, with Heidelberg being situated ca. 600 km from the Atlantic coast and Krakow approximately 18 1000 km further inland, it was of interest to quantify the extent of ²²²Rn build-up in the air 19 masses on their way from Heidelberg to Krakow. The evolution of ²²²Rn content within the 20 21 PBL can be described by a simple mass balance equation:

$$22 \qquad \frac{dC_{Rn}}{dt} = S_{Rn} - (\lambda_d + \lambda_e)C_{Rn} \tag{5}$$

where: C_{Rn} - concentration of ²²²Rn in the convective mixed layer of PBL; S_{Rn} - source term 23 linked to the surface flux of ²²²Rn. $S_{Rn} = F_{Rn}/h$ where F_{Rn} is the surface flux of ²²²Rn and h is 24 the height of the convective mixed layer; λ_d - decay constant of ²²²Rn; λ_e - rate constant 25 associated with removal of ²²²Rn across the PBL boundary. Equation (5) implies perfect 26 mixing within the PBL and assumes that net exchange of ²²²Rn due to horizontal transport 27 28 perpendicular to the direction of air mass movement is equal zero. It has to be noted that 29 convective mixed layer referred to in eq. (5) extends to PBL boundary while the nocturnal mixing layer discussed in section 3.2.1 is generally less extensive (H < h) and occupies only 30

1 part of PBL (Stull, 1988). Analytical solution of eq. (5), with the initial condition $C_{Rn}(o) = 0$, 2 constant surface flux of ²²²Rn and constant height of the convective mixed layer, reads as 3 follows:

4
$$C_{Rn}(t) = \frac{S_{Rn}}{\lambda_d + \lambda_e} \left(1 - e^{-(\lambda_d + \lambda_e)t} \right)$$
(6)

5 Substituting $S_{Rn} = F_{Rn}/h$ and $T_{eff} = 0.693/(\lambda_d + \lambda_e)$ it follows:

6
$$C_{Rn}(t) = \frac{F_{Rn} \cdot T_{eff}}{h \cdot 0.693} \left(1 - e^{-\frac{0.693}{T_{eff}}t} \right)$$
 (7)

Equation (7) can be used to calculate the expected build-up of ²²²Rn content within PBL for maritime air masses travelling eastward from the Atlantic Ocean towards Heidelberg and Krakow. It is apparent from the preceding discussion that local effects are decisive in shaping up atmospheric ²²²Rn levels at both sites. Therefore, careful screening and selection of ²²²Rn data had to be performed before an attempt is made to quantify the "continental effect" between both sites.

13 In the first step, only trajectories arriving in Krakow, which also passed Heidelberg at the distance of less then 100 km were selected for further analysis (Fig. 9). Out of this population 14 15 of trajectories, only those which were travelling from the vicinity of Heidelberg to Krakow during the period shorter than 24 hours, were selected. Finally, only the data representing 16 minimum ²²²Rn concentrations measured at both sites during these periods, taking into 17 18 account the time lag due to atmospheric transport from Heidelberg to Krakow, were 19 considered. This step was aimed at minimizing the impact of local effects (inversion episodes) on the measured ²²²Rn concentrations at both sites. The selected minimum ²²²Rn 20 21 concentrations (94 pairs), averaged over entire observation period (January 2005 - December 2009) separately for each station are equal 1.12 ± 0.09 and 1.90 ± 0.11 Bg m⁻³, for Heidelberg 22 and Krakow, respectively. The difference (0.78±0.12 Bq m⁻³) represents mean build-up of 23 ²²²Rn content between Heidelberg and Krakow. 24

25 Sensitivity analysis of eq. (7) was made with an attempt to fit simultaneously three values: 26 (i) the measured mean minimum 222 Rn concentrations at Heidelberg and Krakow, and (ii) the 27 difference between them. Three parameters in eq. (7) were treated as adjustable parameters: 28 (i) the mean transport velocity (*V*) of the air masses within convective mixed layer of PBL, on 29 their route from the Atlantic coast to Heidelberg and Krakow, (ii) the mean surface flux of

²²²Rn (F_{Rn}) on the European continent from the Atlantic coast till Krakow, and (iii) the rate 1 constant of ²²²Rn removal across the top of PBL (λ_e). Since HYSPLIT routine returns also the 2 3 mixing height of the calculated backward trajectories, it was possible to derive the value of 4 this parameter directly from the model output. The mixing heights were calculated for all trajectories which satisfied the selection criteria outlined above and the maximum mixing 5 6 height for each trajectory was selected. Averaged over all trajectories considered in the calculation, this procedure yield the mean value of the convective mixed layer height equal 7 8 1083±31 m. In calculations 1100 m was adopted. The selection of 94-hours trajectories impose a lower limit on the velocity of air masses travelling from the Atlantic coast to 9 Heidelberg and Krakow, equal approximately 2.8 m s⁻¹. The values of transport velocities 10 used in sensitivity analysis varied between 3.0 and 4.0 m s⁻¹. It appeared that while the 11 calculated ²²²Rn concentrations at Heidelberg and Krakow depend on the assumed transport 12 13 velocity, the difference between them is relatively insensitive to the actual value of this parameter. The rate constant of radon removal across the top of PBL (λ_e) and the mean 14 surface ^{222}Rn flux varied in the calculations from $\lambda_e=0.5\lambda_d$ to $\lambda_e=1.5\lambda_d$ and from 30 to 60 15 Bq $m^{-2}h^{-1}$, respectively. 16

Goodness of the fitting procedure was quantified by calculating the sum of squared 17 differences $(\Sigma(C_{Rn(m)} - C_{Rn(c)})^2)$ between the measured $(C_{Rn(m)})$ and calculated $(C_{Rn(c)})$ mean 18 minimum ²²²Rn concentration at Heidelberg and Krakow, and the difference between them. 19 The best fit (Σ =9.9x10⁻⁴) was obtained for the following combination of the adjusted 20 parameters: $V = 3.5 \text{ m s}^{-1}$, $F_{Rn} = 36 \text{ Bg m}^{-2}\text{h}^{-1}$ and $\lambda_e = \lambda_d$. Similarity between the rate constant 21 of radon removal across the top of PBL and the decay constant of ²²²Rn was suggested also by 22 Lui et al. (1984). The mean continental ²²²Rn surface flux between the Atlantic coast and 23 Krakow obtained through the fitting procedure (36 Bq $m^{-2}h^{-1}$) is lower than the mean annual 24 222 Rn flux obtained for Krakow urban area in the framework of this study (ca. 47 Bq m⁻²h⁻¹) 25 and is significantly lower than the annual mean ²²²Rn flux (ca. 57 Bq m⁻²h⁻¹) estimated for 26 27 Heidelberg area from long-term flux measurements at five locations with different soil texture (Schmidt et al., 2003). If the mean surface ²²²Rn flux of 52 Bq m⁻²h⁻¹ is assumed (arithmetic 28 average of Heidelberg and Krakow best estimates of this parameter), an equally good fit of 29 the measured minimum ²²²Rn concentrations at both stations and the difference between them 30 31 is also possible, although with significantly larger height of the convective mixed layer (h =32 1600 m). This suggest that the modelled value of convective mixing layer is underestimated.

1 5 Conclusions

Systematic observations of ²²²Rn concentration in near-ground atmosphere at two continental sites in Europe, supplemented by measurements of surface ²²²Rn fluxes, allowed a deeper insight into factors controlling spatial and temporal variability of ²²²Rn in near-ground atmosphere over central Europe. The available data allowed to address the role of local and regional factors in controlling the observed atmospheric ²²²Rn levels and their variability.

Atmospheric concentrations of ²²²Rn at both observation sites vary on daily, synoptic and monthly time scales. Generally higher and more variable ²²²Rn concentrations recorded in Krakow are mainly due to specific characteristics of the local atmosphere, such as lower wind speed and more frequent inversion periods of prolonged duration when compared to Heidelberg, thus leading to enhanced stability of the lower atmosphere at this monitoring site.

12 The presented data reveal a distinct asymmetry in the shape of seasonal variations of surface ²²²Rn fluxes and ²²²Rn concentrations measured in the local atmosphere of Krakow. 13 While atmospheric ²²²Rn contents peak in November, the ²²²Rn exhalation rates reach their 14 maximum in September-October. This distinct phase shift stems most probably from 15 16 increased stability of the lower atmosphere during autumn months (higher percentage of still periods, longer duration of ground-based inversion episodes). These factors may collectively 17 lead to the observed ²²²Rn maximum in the local atmosphere in November, despite of already 18 19 weakening soil ²²²Rn flux at that time of the year.

Although the atmospheric ²²²Rn levels at Heidelberg and Krakow appeared to be 20 controlled primarily by local factors, it was nevertheless possible to evaluate the "continental 21 effect" in atmospheric ²²²Rn content between both sites, related to gradual build-up of ²²²Rn 22 load of maritime air masses travelling eastward over the European continent. Satisfactory 23 agreement obtained between the measured and modelled minimum ²²²Rn concentrations at 24 both sites and the difference between them, derived from a simple box model coupled with 25 26 HYSPLIT analysis of air mass trajectories, allowed to put some constraints on the parameters of atmospheric ²²²Rn transport over the European continent and its surface fluxes. 27

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Table 1. Indicators of daily variations of ²²²Rn activity in Krakow (KR) and Heidelberg (HD)
 during the period January 2005 – December 2009. The quoted uncertainties represent
 standard uncertainties of the mean values.

Period		Mean daily ²²² Rn activity (Bq m ⁻³)	Mean daily maximum ²²² Rn activity (Bq m ⁻³)	Mean daily minimum ²²² Rn activity (Bq m ⁻³)	Peak-to-peak amplitude (Bq m ⁻³)
Winter (Dec Febr.)	KR	5.65 ± 0.17	9.36 ± 0.27	2.99 ± 0.11	6.37 ± 0.23
	HD	5.19 ± 0.19	7.93 ± 0.26	3.10 ± 0.14	4.83 ± 0.17
Spring (March - May)	KR	4.00 ± 0.09	8.81 ± 0.23	1.52 ± 0.05	7.29 ± 0.22
	HD	3.14 ± 0.08	5.74±0.16	1.54 ± 0.05	4.21 ± 0.15
Summer (June August)	KR	5.50 ± 0.13	12.11 ± 0.30	2.02 ± 0.05	10.09 ± 0.27
	HD	3.55 ± 0.07	6.88 ± 0.17	1.61 ±0.04	5.27 ± 0.15
Autumn (SeptNov.)	KR	8.70 ± 0.23	15.59 ± 0.39	3.83 ± 0.13	11.76 ± 0.35
	HD	6.08 ± 0.17	10.13 ± 0.27	3.38 ± 0.13	6.75 ± 0.21
Mean 2005-2009	KR	5.86 ± 0.09	11.28 ± 0.16	2.55 ± 0.05	8.72 ± 0.14
	HD	4.50 ± 0.07	7.67 ± 0.12	2.42 ± 0.05	5.26 ± 0.09

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Fig. 1. Time series of hourly mean ²²²Rn content in near-ground atmosphere, recorded between
January 2005 and December 2009 in Krakow, southern Poland (a) and Heidelberg, south-west
Germany (b).



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Fig. 2. Diurnal variations of ²²²Rn concentration in near-ground atmosphere over Krakow (a) and Heidelberg (b) during the period January 2005 - December 2009, averaged separately for each hour and for four seasons: winter (DJF), spring (MAM), summer (JJA) and autumn (SON). Diurnal variation of ²²²Rn content at both sites averaged for entire observation period is also shown (c). Vertical marks accompanying the data points indicate standard uncertainties of the calculated mean values.



Fig. 3. (a) - Changes of atmospheric ²²²Rn concentration in Krakow between March 1 and 13, 2005.
Horizontal lines with arrows and letter symbols mark different air masses identified in (c). (b) - Daily
means of atmospheric ²²²Rn concentration, wind speed and rainfall amount in Krakow. (c) - 96-hours
backward trajectories of the air masses arriving in Krakow, starting at 25%, 50% and 75% of the time
period A, B and C marked in (a). (d) - changes of the elevation of trajectories shown in (c).



Fig. 4. Box-and-whisker plot of monthly ²²²Rn concentration recorded in near-ground atmosphere of
Krakow (a) and Heidelberg (b) between January 2005 and December 2009. Marked percentiles
represent 5%, 25%, 75% and 95% confidence interval, black squares represent medians and horizontal
lines represent mean values.



Fig. 5. Monthly mean night-time fluxes of ²²²Rn in Krakow urban area for the period June 2004 - May
2009, derived from atmospheric ²²²Rn observations and sodar measurements of the mixing layer height
(see text for details). Error bars indicate standard uncertainties of the mean values. Blue bars represent
the number of nights used for calculation of the monthly means.



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Fig. 6. Comparison of monthly means of night-time surface fluxes of ²²²Rn in Krakow for the period from September 2005 till September 2006, derived from direct measurements of soil ²²²Rn flux using chamber method (red bars) and from atmospheric ²²²Rn observations combined with sodar measurements of the mixing layer height (black bars). Error bars indicate standard uncertainty of the mean values.



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Fig. 7. Spatial distribution of 96-hours backward trajectories of the air masses arriving in Heidelberg, 3 southern Germany, and Krakow, southern Poland, during the period January 2005 - December 2009,

- shown for four seasons, calculated using HYSPLIT model (Draxler and Rolph, 2011; Rolph, 2011) see text for details. The maps are supplemented with statistics of trajectory height for each season and
 station. Marked percentiles represent 5%, 25%, 75% and 95% confidence interval, black squares
 represent medians and horizontal lines represent mean values.



Fig. 8. (a) - Box-and-whisker plot of monthly ²²²Rn concentration measured in near-ground 3 4 atmosphere of Krakow. Median is represented by horizontal line while arithmetic average is shown by full dot. (b) - Monthly minima of atmospheric ²²²Rn content, as observed in Krakow and Heidelberg, 5 calculated as arithmetic averages of daily ²²²Rn minima for the given month, averaged over entire 6 7 observation period from January 2005 till December 2009. (c) - Monthly means of night-time ²²²Rn 8 fluxes in Krakow, averaged over the period from June 2004 till May 2009. (d) - Monthly means of soil 9 ²²²Rn fluxes in Krakow derived from chamber measurements, averaged over the period from 10 September 2005 till September 2006. (e) Monthly means of surface air temperature and precipitation 11 in Krakow, averaged over the period from January 2005 till December 2009. (f) - Monthly means of 12 surface air and soil temperatures in Krakow. (g) - Percentage of periods with wind speed below 1 m s⁻ ¹ in Krakow, averaged over the period from January 2005 till December 2009. (h) - Monthly means of 13

- 1 number of hours with ground-based inversion per day, averaged over the period from January 1994 till
- 2 December 1999.
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Fig. 9. Spatial distribution of 96-hour backward trajectories of the air masses arriving in Krakow,
southern Poland, which passed Heidelberg, southern Germany, at the distance of less than 100 km,
calculated using HYSPLIT model (Draxler and Rolph, 2011; Rolph, 2011 - see text for details). The
analysis comprised the period from January 2005 till December 2009.