1	Isotopic signatures of production and uptake of H <sub>2</sub> by soil
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Abstract: Molecular hydrogen  $(H_2)$  is the second most abundant reduced trace 23 24 gas (after methane) in the atmosphere, but its biogeochemical cycle is not well understood. Our study focuses on the soil production and uptake of H<sub>2</sub> and the 25 associated isotope effects. Air samples from a grass field and a forest site in the 26 Netherlands were collected using soil chambers. The results show that uptake 27 and emission of H<sub>2</sub> occurred simultaneously at all sampling sites, with strongest 28 29 emission at the grassland sites where clover ( $N_2$  fixing legume) was present. The H<sub>2</sub> mole fraction and deuterium content were measured in the laboratory to 30 determine the isotopic fractionation factor during  $H_2$  soil uptake ( $\alpha_{soil}$ ) and the 31 isotopic signature of  $H_2$  that is simultaneously emitted from the soil ( $\delta D_{soil}$ ). By 32 considering all net-uptake experiments, an overall fractionation factor for 33 deposition of  $\alpha_{soil} = k_{HD} / k_{HH} = 0.945 \pm 0.004$  (95% CI) was obtained. The 34 difference in mean  $\alpha_{soil}$  between the forest soil 0.937±0.008 and the grassland 35 0.951±0.025 is not statistically significant. For two experiments, the removal of 36 soil cover increased the deposition velocity ( $v_d$ ) and  $\alpha_{soil}$  simultaneously, but a 37 general positive correlation between  $v_{\rm d}$  and  $\alpha_{\rm soil}$  was not found in this study. 38 When the data are evaluated with a model of simultaneous production and 39 uptake, the isotopic composition of H<sub>2</sub> that is emitted at the grassland site is 40 calculated as  $\delta D_{soil} = (-530 \pm 40)$  ‰. This is less deuterium-depleted than what is 41 expected from isotope equilibrium between H<sub>2</sub>O and H<sub>2</sub>. 42

## 44 **1. Introduction**

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46  $H_2$  is considered as alternative energy carrier to replace fossil fuels in the future. 47 However, the environmental and climate impact of a potential widespread use 48 of  $H_2$  is still under assessment. Several studies suggested that the atmospheric 49  $H_2$  mole fraction might increase substantially in the future due to the leakage 50 during production, storage, transportation and use of  $H_2$ , which could 51 significantly affect atmospheric chemistry (Schultz et al., 2003; Tromp et al., 52 2003; van Ruijven et al., 2011; Warwick et al., 2004).

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In the troposphere,  $H_2$  has a mole fraction of about 550 parts per billion (ppb = 54 nmol mol<sup>-1</sup>) and a lifetime of around 2 years (Novelli et al., 1999; Price et al., 55 2007; Xiao et al., 2007; Pieterse et al., 2011; 2013). H<sub>2</sub> can affect atmospheric 56 chemistry and composition in several ways. Firstly, it increases the lifetime of 57 the greenhouse gas methane  $(CH_4)$  via its competing reaction with the hydroxyl 58 59 radical (OH) (Schultz et al., 2003; Warwick et al., 2004). Additionally, H<sub>2</sub> affects air quality because it is an ozone  $(O_3)$  precursor and indirectly increases 60 the lifetime of the air pollutant carbon monoxide (CO) through competition for 61 62 OH. In the stratosphere,  $H_2O$  that is produced through the oxidation of  $H_2$ increases humidity, which can result in increased formation of polar 63

64 stratospheric clouds and  $O_3$  depletion (Tromp, et al., 2003), but this effect may 65 be weaker than estimated initially (Warwick et al. 2004; Vogel et al., 2012).

The main sources of tropospheric  $H_2$  are the oxidation of  $CH_4$  and non-methane hydrocarbons (NMHC) (48%), biomass burning (19%), fossil fuel combustion (22%) and biogenic  $N_2$  fixation in the ocean (6%) and on land (4%), while the main sinks are soil uptake (70%) and oxidation by OH (30%) (Pieterse et al, 2013).

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The biogenic soil sink of H<sub>2</sub> is the largest and most uncertain term in the global 73 atmospheric H<sub>2</sub> budget. Conrad and Seiler (1981) assumed that the soil uptake 74 of atmospheric H<sub>2</sub> is most likely due to consumption by abiotic enzymes, since 75 there were no soil microorganisms known to be able to fix  $\mathrm{H}_{2}$  at the low 76 atmospheric mole fraction at that time. This remained the basic hypothesis of 77 78 many further soil uptake studies (Conrad et al., 1983; Conrad and Seiler, 1985; Ehhalt and Rohrer, 2011; Guo and Conrad, 2008; Häring et al., 1994; Smith-79 Downey et al., 2006). However, Constant et al. (2008a) were first to identify an 80 aerobic microorganism (Streptomyces sp. PCB7) that can consume  $H_2$  at 81 tropospheric ambient mole fractions, and suggested that active metabolic cells 82 could be responsible for the soil uptake of  $H_2$  rather than extracellular enzymes. 83 Further studies showed that uptake activity at ambient H<sub>2</sub> level is widespread 84 85 among the streptomycetes (Constant et al., 2010) and it was postulated that high affinity  $H_2$ -oxidizing bacteria are the main biological agent responsible for the soil uptake of atmospheric  $H_2$  (Constant et al., 2011). Khdhiri et al. (2015) suggested that the relative abundance of high affinity  $H_2$ -oxidation bacteria and soil carbon content could be used as predictive parameters for the  $H_2$  oxidation rate. Determining the dominant mechanism of the  $H_2$  soil uptake activity is still an active area of research.

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It has been shown that soil uptake of H<sub>2</sub> can coexist with soil production 93 (Conrad, 1994).  $H_2$  is produced in the soil during  $N_2$  fixation (e.g. by bacteria 94 95 living symbiotically in the roots of legumes such as clover or beans) and dark fermentation. Although the H<sub>2</sub> produced in the soil by e.g. N<sub>2</sub> fixation can be 96 largely consumed within the soil, a significant amount of H<sub>2</sub> escapes to the 97 atmosphere (Conrad and Seiler, 1979; 1980). Conrad and Seiler (1980) 98 estimated that 2.4 to 4.9 Tg  $a^{-1}$  of  $H_2$  is emitted into the atmosphere through  $N_2$ 99 100 fixation on land.

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102 One approach to better understand the sources and sinks of  $H_2$  is to investigate 103 the isotopic fractionation processes involved, which act as a fingerprint for  $H_2$ 104 emitted from different sources or destroyed by different sinks. The isotopic 105 composition of  $H_2$  is expressed as:

$$\delta(\mathrm{D},\mathrm{H}_2) = \frac{R_{\mathrm{sa}}}{R_{\mathrm{VSMOW}}} - 1$$

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108 where  $R_{sa}$  is the D/H ratio of the sample H<sub>2</sub> and  $R_{VSMOW} = (155.76\pm0.8)$  parts per 109 million (ppm = mmol mol<sup>-1</sup>) is the same ratio of the standard material, Vienna 110 Standard Mean Ocean Water (VSMOW) (De Wit et al.,1980; Gonfiantini et al., 111 1993). For brevity, we will use the notation  $\delta D$  (= $\delta D(D, H_2)$ ) throughout the 112 rest of this paper. The  $\delta D$  values are usually given in per mill (‰). Recent 113 studies showed that the global mean  $\delta D$  value of atmospheric H<sub>2</sub> is about +130 ‰ 114 (Batenburg et al., 2011; Gerst et al., 2000, 2001; Rice et al., 2010).

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The HH molecule is consumed preferentially over HD during both OH 116 117 oxidation and soil uptake, with OH oxidation causing a much stronger isotope fractionation effect. Only a few studies have investigated the soil uptake of H<sub>2</sub> 118 119 with isotope techniques. Gerst and Quay (2001) carried out field experiments in Seattle, United States and found  $\alpha_{soil} (= k_{HD}/k_{HH})$  to be 0.943±0.024 (1 $\sigma$ ). Note 120 that  $k_{\rm HD}$  and  $k_{\rm HH}$  are removal rate constants for HD and HH respectively. Rahn 121 et al. (2002a) collected air samples from four forest sites in ecosystems of 122 123 different ages in Alaska, United States, in July 2001, and obtained a similar average value (0.94±0.01). They suggested that  $\alpha_{\rm soil}$  depends on the forest 124 maturity, with smaller fractionation for more mature forests. Since the more 125 mature forests showed larger deposition velocity  $(v_d)$  of H<sub>2</sub>, they further 126

suggested that lower uptake rates involve greater isotopic fractionation ( $\alpha_{soil}$ ) 127 128 further from 1) than fast uptake rates. Rice et al. (2011) performed deposition experiments in Seattle and found  $\alpha_{soil}$  varying from 0.891 to 0.976, with a mean 129 of 0.934. They found  $\alpha_{soil}$  to be correlated with  $v_d$ , with smaller isotope effects 130 ( $\alpha_{soil}$  closer to 1) occurring at higher  $v_d$ , which agreed with the suggestion by 131 132 Rahn et al. (2002a). In addition, unpublished experiments from Rahn et al. (2005) yielded  $\alpha_{soil} = 0.89 \pm 0.03$  in three upland ecosystems that were part of an 133 Alaskan fire chronosequence. The data suggest that variability in the 134 soil/ecosystem affects  $\alpha_{soil}$  but no significant variability of  $\alpha_{soil}$  with season was 135 detected. Hitherto, only  $\alpha_{soil}$  values from studies in Seattle and Alaska are 136 available, and values from other locations and ecosystems are needed to learn 137 more about the factors influencing  $\alpha_{soil}$ . 138

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The  $\delta D$  of H<sub>2</sub> from various surface sources has been reported as about -290 ‰ 140 141 for biomass burning (Gerst and Quay, 2001; Haumann et al., 2013) and between -200 ‰ and -360 ‰ for fossil fuels combustion (Rahn et al., 2002b; Vollmer et 142 al., 2012). So far no field studies have determined the isotopic composition of 143 144 the H<sub>2</sub> emitted from soil. Two laboratory studies examined the isotopic signature of H<sub>2</sub> produced from N<sub>2</sub> fixation. Luo et al. (1991) reported a 145 fractionation factor  $\alpha_{\text{H2/H2O}} = R(D/H, H_2)/R(D/H, H_2O) = 0.448\pm0.001$  between 146 the  $H_2$  produced from  $N_2$  fixation and the  $H_2O$  used to grow the  $N_2\mbox{-fixing}$ 147 148 bacteria for Synechococcus sp. and 0.401±0.002 for Anabaena sp., respectively.

Walter et al. (2012) reported  $\alpha_{H2/H2O} = 0.363 \pm 0.019$  for the N<sub>2</sub>-fixing 149 150 rhizobacterium Azospirillum brasiliensis. It has been proposed that microbiological H<sub>2</sub> consumption and production could modify the thermal 151 isotopic equilibrium between H<sub>2</sub> and H<sub>2</sub>O in low-temperature hydrothermal 152 fluids (Kawagucci et al., 2010). Compared to the surface sources, H<sub>2</sub> produced 153 from CH<sub>4</sub> and NMHC oxidation is isotopically strongly enriched in deuterium, 154 155 with  $\delta D$  beween +120 and +180 % (Rahn et al., 2003; Röckmann et al. 2003a, 156 Pieterse et al., 2011).

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Here we report measurements of the isotopic fractionation factors of  $H_2$  during soil deposition at two different sites in the Netherlands, a forest and a grassland site. For the grassland site we also determine the apparent isotopic composition of the  $H_2$  that was simultaneously emitted from the soil during the experiment.

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# 164 **2. Methods**

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## 166 **2.1 Sampling**

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168 Air samples were collected from a soil chamber at two locations in the
169 Netherlands (Fig. 1): a grass field around the Cabauw tall tower (51°58' N,

4°55' E) and a forest site near Speuld (52°13' N, 5°39' E). Two types of ground
cover (grass with and without clover) were sampled at Cabauw, while three
types of forest (Douglas fir, beech and spruce) were selected in Speuld. More
information about the soil and vegetation type can be found in Beljaars and
Bosveld (1997) for the Cabauw site, and in Heij and Erisman (1997) for the
Speuld site.

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Flask samples were filled with air from a soil chamber, using a closed-cycle air 177 178 sampler (Fig. 2). The soil chamber consisted of two parts: the chamber body 179 with a metal base at the bottom that was inserted about 2 cm into the soil, and a 180 removable transparent lid with two connections for air sampling. The chamber had a height of 40 cm, an area of 570 cm<sup>2</sup> and a volume of 22.8 L; the air inside 181 was mixed by a fan. The sampler could hold four flasks installed in series, 182 which could be bypassed independently; the flow and pressure in the flasks 183 were controlled. The air was dried using  $Mg(ClO_4)_2$ . After passing through the 184 flasks the air was returned to the soil chamber, which kept the pressure inside 185 the chamber approximately constant during sampling. 186

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Air samples were collected from the chamber in 1 L glass flasks at 0, 10, 20 and 30 minutes after closing the chamber (time interval changed to 5 minutes in Speuld because of the faster uptake). The gas flasks (Normag, Ilmenau, Germany) were made of borosilicate glass 3.3 with O-ring-sealed stopcocks

192 made of PCTFE (Kel-F) and covered with a dark hose. Thorough tests have 193 demonstrated that air samples with typical trace gas content are stable in these 194 flasks (Rothe et al., 2004). In the beginning, the whole sampling unit (all lines, connections and flasks) was flushed with ambient air for about 10 minutes at a 195 flow rate of 2 L min<sup>-1</sup> and a pressure of 100 kPa, with all flasks open and the 196 chamber lid open. This initial flushing process was designed to fill the flasks 197 198 with background air. The air pressure inside the flasks was increased to 200 kPa 199 (180 kPa for Speuld samples) by adjusting the flow control valve and the valves 200 on two pressure gauges (Fig. 2) before chamber closing and then maintained 201 constant during the whole sampling time. The flow rate was maintained at 2 L 202 min<sup>-1</sup> at ambient pressure and temperature with a rotameter and the pressure 203 inside the chamber was maintained at 100 kPa during the whole sampling time. 204 The temperature was not recorded during the sampling. After the initial flushing, 205 the first flask was closed and then the chamber was closed as well. Afterwards, 206 the air was flushed from the chamber through three flasks (the first flask was by-passed) and back to the chamber. After 10, 20 and 30 minutes, the second, 207 208 third and fourth flasks were closed.

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A total of 36 sets of air samples were collected in Cabauw during summer (June,
July and August) 2012 and 12 sets were collected in Speuld in September 2012.
Each set contains four air samples. In total, 186 valid samples were analyzed for
H<sub>2</sub> mole fraction and its deuterium content (6 were lost during sampling,

transportation and measurement). All the Speuld samples and about half of the Cabauw samples were further used for analysis in this study. The reason why 50% of the Cabauw experiments were not used is that these experiments showed neither strong  $H_2$  emission nor  $H_2$  uptake and the isotopic signals were weak. Most experiments were conducted with the 22.8 L volume soil chamber as described above, while 10 experiments were conducted with a larger automated soil chamber with a volume of 125 L and a height of 22.5 cm.

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# 223 2.2 Laboratory determination of H<sub>2</sub> mole fraction and deuterium 224 content of air samples

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The mole fraction and the  $\delta D$  of H<sub>2</sub> were measured with a gas chromatography 226 isotope ratio mass spectrometry (GC/IRMS) setup (Rhee et al., 2004). For H<sub>2</sub> 227 228 mole fractions, the laboratory working standards are linked to the MPI-2009 229 scale (Jordan and Steinberg, 2011). The  $\delta D$  values of the laboratory reference gases are indirectly linked to mixtures of synthetic air with H<sub>2</sub> of known 230 isotopic composition, certified by Messer Griesheim, Germany (Batenburg et al., 231 232 2011). Most of the samples collected from Cabauw were measured within two months after sampling, while the samples from Speuld were kept in a dark 233 234 storage room for around four months before measurement.

The operational principle of the GC/IRMS system is to separate  $H_2$  from the air matrix at low temperature (about 36 K) and measure the HH and HD content with a mass spectrometer. The measurement includes four main steps:

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(1) A glass sample volume (750 ml) is evacuated and subsequently filled with sample air to approximately 700 mbar. This volume is then exposed to a cold head (36 K) of a closed-cycle helium compressor for 9 minutes. During this stage, all gases except  $H_2$ , helium (He) and neon (Ne) condense.

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(2) The remainder in the headspace of the cold head and sample volume is then flushed with He carrier gas to a pre-concentration trap where  $H_2$  is collected on a 25 cm long, 1/8 inch OD (outside diameter) stainless steel tube filled with fine grains (0.2 to 0.5 mm) of 5 Å molecular sieve, for 20 minutes. The preconcentration trap is cooled down to the triple point of nitrogen (63 K) by keeping it in a liquid N<sub>2</sub> reservoir that is further cooled down by pumping on the gas phase.

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253 (3) After the collection of  $H_2$ , the pre-concentration trap is warmed up to release 254 the absorbed  $H_2$ , which is then cryo-focused for 4 minutes on a capillary (25 cm 255 long, 0.32 mm ID (inside diameter)) filled with 5 Å molecular sieve at 77 K. 256 After that, the cryo-focus trap is warmed up to ambient temperature and the  $H_2$  sample is flushed with He carrier gas onto the GC column (5 Å molecular sieve,  $\approx 323$  K) where H<sub>2</sub> is chromatographically purified from potential remaining interferences.

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261 (4) In the end, the purified  $H_2$  is carried by the He carrier gas via an open split 262 interface (Röckmann et al., 2003b) into the IRMS for D/H ratio determination. 263

264 More details about the GC/IRMS system and measurement steps can be found 265 in Rhee et al. (2004) and Röckmann et al. (2010). The data correction 266 procedures and isotope calibration are similar to those described in Batenburg et 267 al. (2011). Four reference gases were used to determine the  $\delta D$  values of the 268 samples. Two of them (Ref-1 and Ref-2) with  $\delta D$  values of (+207.0 ± 0.3) % 269 and  $(+198.2 \pm 0.5)$  ‰ were calibrated and used previously in Batenburg et al. 270 (2011). The other two new reference gases (Ref-3 and Ref-4) were calibrated 271 versus Ref-1 and Ref-2. The  $\delta D$  value of Ref-3 was (-183± 2.4) ‰. Ref-4 was a frequently measured reference gas that was measured usually about 5 times per 272 273 sequence of measurement, while other three reference gases were measured about 1 to 3 times per sequence of measurement. The  $\delta D$  value of Ref-4 274 dropped linearly with time from -115 % to -157 % between 1 Jun 2012 and 15 275 276 Feb 2013, while the other three reference gases were stable.

#### 278 **2.3 Non-linearity of the GC/IRMS system**

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Ideally, the  $\delta D$  of H<sub>2</sub> measured with the GC/IRMS should not depend on the total amount of H<sub>2</sub> used for analysis, but in practice a dependence of the isotopic composition on the amount of H<sub>2</sub> is observed for low mole fractions. This is called non-linear behavior, and it is a particularly severe limitation for soil uptake studies, since the mole fraction in such samples can decrease by more than an order of magnitude. For comparison, in ambient background air the H<sub>2</sub> mole fraction variations are usually no more than 20%.

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Experiments were carried out with different quantities of air from various 288 laboratory reference bottles with known  $\delta D$  to determine a suitable correction 289 290 for the non-linear behavior. The measured  $\delta D$  increases with the mass 2 sample peak area, which is proportional to the  $H_2$  quantity in the sample. In the peak 291 292 area range of 0.2 Vs to 1 Vs this relation can be parameterized by a logarithmic function  $\delta D = 54.6 \ln$  (peak area/Vs) %, which is used as correction function 293 for the measurements at low peak areas (Fig. 3). The linearity correction 294 introduces an additional uncertainty due to uncertainties in the logarithmic fit, 295 particularly at low peak areas. The total assigned uncertainty for each 296 measurement is calculated from the analytical and fitting uncertainty, as a 297 298 function of peak area (Fig. 4). It is 2 % for ln (peak area/Vs) of 1.5 or more

(equivalent to more than 600 ppb  $H_2$  in an air sample), but increases to 32 %299 when ln (peak area/Vs) drops to -1.6 ( $\approx 20$  ppb H<sub>2</sub> in air sample). In total, the 300 301  $\delta D$  results of 18 Speuld samples that were measured at these low peak areas were corrected with this linearity correction. Possible additional systematic 302 303 errors (a few %) may arise from uncertainties in the initially assigned  $\delta D$ values of the commercial calibration gases, changes of these values in the 304 process of creating calibration mixtures with near-ambient H<sub>2</sub> concentration, 305 and the calibration measurements themselves (Batenburg et al., 2011). 306

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#### 309 2.4 Data evaluation

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Assuming first order kinetics for  $H_2$  removal and a constant production rate *P* over the course of a deposition experiment, the time evolution of the mole fraction *c* of non-deuterated  $H_2$  (HH) inside the soil chamber can be expressed as:

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$$\frac{\mathrm{d}\,c}{\mathrm{d}t} = P - kc \tag{1}$$

316

where *k* is the first order uptake rate constant of HH. For well-mixed air in the chamber,  $k = v_d/h$ , where  $v_d$  is the gross deposition velocity of H<sub>2</sub> and *h* is the 319 chamber height. The gross deposition velocity is the deposition velocity 320 corrected for production, which is different from the net deposition velocity 321 reported in some studies in the past that showed the effective uptake of  $H_2$  from 322 the atmosphere. The solution of Eq. (1) is of the form:

323

$$c = (c_{\rm i} - c_{\rm e})e^{-kt} + c_{\rm e}$$
 (2)

324

where c,  $c_i$  and  $c_e$  (= P/k) are the mole fractions of HH at time t, initially and at equilibrium, respectively. Therefore, P and k can be obtained by fitting an exponential function to the time evolution of HH inside the chamber. Similarly, we can obtain P' and k' from the time evolution of HD.

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$$c' = (c'_{\rm i} - c'_{\rm e})e^{-k't} + c'_{\rm e}$$
(3)

330

where  $c', c_i', c'_e (= P'/k'), P'$  and k' are the corresponding parameters for HD.

Equations (2) and (3) constitute the mass balance model that we used to analyze our data. When k, k', P and P' have been determined,  $\alpha_{soil}$  and  $\delta D_{soil}$  can be calculated simply as:

$$\alpha_{\rm soil} = \frac{k'}{k} \tag{4}$$

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$$\delta D_{\rm soil} = \frac{P'/P}{2R_{\rm VSMOW}} - 1 \tag{5}$$

338

However, fitting an exponential curve to only four sample data yields relatively large errors for k, k', P and P', which propagate to large errors for  $\alpha_{soil}$  and  $\delta D_{soil}$ if they are determined directly from Eqs (4-5).

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In Rice et al. (2011), Equations (2) and (3) were combined to calculate  $\alpha_{soil}$  in the presence of both source and sink of H<sub>2</sub> using  $c_e$  and  $c_e'$  from the exponential fits:

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$$\ln \frac{c' - c_{\rm e}}{c_{\rm i}' - c_{\rm e}'} = \frac{k'}{k} \ln \frac{c - c_{\rm e}}{c_{\rm i} - c_{\rm e}}$$
(6)

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348  $\alpha_{soil} = k'/k$  can be obtained by plotting  $\ln \frac{c'-c'_e}{c'_i-c'_e}$  versus  $\ln \frac{c-c_e}{c_i-c_e}$  and fitting a linear 349 function. In the absence of soil emission ( $c_e = c'_e = 0$ ), Eq. (6) collapses to the 350 well-known Rayleigh fractionation equation that is used to quantify the isotope 351 fractionation during single stage removal processes in the absence of sources.

For the high emission measurements, where production overwhelms consumption, we use the relations  $c_e = P/k$  and  $c'_e = P'/k'$ , and obtain P'/P from the slope of  $c'_e \ln \frac{c'-c'_e}{c'_i-c'_e}$  against  $c_e \ln \frac{c-c_e}{c_i-c_e}$ . Then  $\delta D_{soil}$  is calculated from Eq. (5).

#### 357 2.5 Flask sampling model

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The advantage of sampling with the soil chamber system described in Section 359 2.1 was that the pressure in the soil chamber stayed constant even when several 360 large samples (2 L each) were taken. A disadvantage was that the volume of air 361 inside the flasks (8 L of air in total) was considerable compared to the volume 362 of air inside the soil chamber (22.8 L). This had two effects: (1) A significant 363 part of the air was at each time separated from the chamber and thus from the 364 365 soil production and uptake. (2) Because of the time lag to flush the samples, the air in a flask was not the same as the air in the chamber at the same time. 366

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We built a flask sampling model to derive correction factors that take into account the influence of the flask sampling system. For a given combination of uptake and production rates, the model simulates the evolution of the  $H_2$  mole fraction in two configurations: the soil chamber alone, and the soil chamber plus four flasks as in our experiments. The model is described in detail in Appendix A. An example of a simulation is shown in Fig. 5. Compared to the situation without flasks, there is a time lag in the decay of  $H_2$  for both the chamber and the flasks after introducing four flasks in the model. The time lag for the second flask is about 2.5 minutes. It increases to 5 minutes for the third flask and is even longer for the fourth flask.

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It is obvious that the sampling process strongly affects the uptake rate  $k_{app}$  and 379 production rate  $P_{app}$  obtained from the direct flask measurements, so we 380 corrected all  $k_{app}$  and  $P_{app}$  values with the correction coefficients derived from 381 this flask sampling model (Appendix A). For a fixed chamber volume, sample 382 pressure, flow rate and time interval of the flask collection that are all recorded 383 384 for each experiment, the relationship between the actual uptake rate constant  $k_{\text{true}}$  and apparent uptake rate constant  $k_{\text{app}}$  can be obtained (see Appendix A). 385 Under the same sampling conditions for a fixed value of  $P_{app}$ , the relationship 386 between actual production rate  $P_{true}$  and apparent production rate  $P_{app}$  depends 387 on  $k_{\text{true}}$  (Fig 10b). 388

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To evaluate the data, we first applied an exponential fit as in Eq. (2) to the measured HH mole fractions for the four flasks in each experiment and obtained *apparent* values  $k_{app}$ ,  $P_{app}$  and  $c_{e,app}$  from the fit parameters. Then we used the correction factors derived from the flask sampling model to retrieve true values 394  $k_{\text{true}}$  and  $P_{\text{true}}$  from the apparent values  $k_{\text{app}}$  and  $P_{\text{app}}$ . One can obtain  $k'_{\text{true}}$  and  $P'_{\text{true}}$ 395 by applying the same method to HD mole fractions inside four flasks.

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397 To determine 
$$\alpha_{soil}$$
, we plotted  $\ln \frac{c'-c'_{e,app}}{c'_1-c'_{e,app}}$  versus  $\ln \frac{c-c_{e,app}}{c_1-c_{e,app}}$  (Eq.6, Fig. 7) and  
398 obtained  $\alpha_{soil,app}$  from the slope of the linear regression. Here, *c* and *c*' are HH

and HD mole fractions in each of the four flasks;  $c_1$  and  $c'_1$  are HH and HD mole fractions of the first flask;  $c_{e,app}$  and  $c'_{e,app}$  are apparent HH and HD equilibrium mole fractions obtained from the exponential fits of HH and HD mole fractions inside the four flasks. We determined the relationship (Fig. 10c) between  $\alpha_{soil,true}$  and  $\alpha_{soil,app}$  obtained from  $\ln \frac{c'-c'_{e,app}}{c'_1-c'_{e,app}}$  versus  $\ln \frac{c-c_{e,app}}{c_1-c_{e,app}}$  using the flask sampling model (see Appendix A1.3). The correction coefficients for each experiment are given in Table 3.

407 Similarly, we obtained 
$$P'_{app}/P_{app}$$
 by plotting  $c'_{e,app} \ln \frac{c'-c'_{e,app}}{c'_1-c'_{e,app}}$  versus  
408  $c_{e,app} \ln \frac{c-c_{e,app}}{c_1-c_{e,app}}$  (Fig. 9), and calculated  $\delta D_{soil,app}$  by use of Eq. (5). Then we  
409 retrieved  $\delta D_{soil,true}$  by use of the flask sampling model (Fig. 10d). The  
410 corresponding correction coefficients for  $\delta D_{soil,app}$  for each net-emission  
411 experiment are shown in Table 3. More information about the retrievals of  
412  $\alpha_{soil,true}$  and  $\delta D_{soil,true}$  can be found in Appendix A.

414	Overall, the sampling effect on $\delta D_{soil}$ is small (less than 22‰). This means that
415	the flask sampling system strongly affects the temporal evolution of HH and
416	HD individually (Fig. 5), and the uptake and production rates derived from flask
417	measurements, but the effects on the computed isotopic signature of the source
418	and sink are relatively small. More details and discussion of the flask sampling
419	model corrections are provided in Appendix A.
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422	3. Results
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424	3.1 Temporal evolution of $H_2$ , HD and $\delta D$
425	
426	Fig. 6 shows examples for the temporal evolution of $H_2$ , HD and $\delta D$ in Cabauw
427	and Speuld, with error estimates included. The errors for $\rm H_2$ and HD are about $4\%$
428	of the respective mole fraction. The error for $\delta D$ ranges from 2 ‰ to 17 ‰.
429	
430	Some of our Cabauw experiments show net soil emission of $H_2$ (upper panels)
431	and some show net soil uptake (middle panels), while all Speuld experiments
432	show net uptake of $H_2$ (lower panels). In the Cabauw net emission experiments,
433	the increase in $H_2$ mole fractions is associated with a strong decrease in $\delta D$ ,
434	showing a strongly depleted $H_2$ source. However, the net uptake experiments at

Cabauw show also a decrease in  $\delta D$ , albeit smaller. In the Speuld experiments, the uptake of H<sub>2</sub> is much faster; the  $\delta D$  increases in the beginning but then decreases again towards the end of the sampling, when the H<sub>2</sub> mole fractions are low.

439

As mentioned in the introduction, soil uptake tends to increase  $\delta D$  while soil emission tends to decrease  $\delta D$  of H<sub>2</sub>. The continuous decrease of  $\delta D$  with time in all Cabauw experiments and the eventual decrease of  $\delta D$  in all Speuld experiments clearly show that there is concurrent soil emission even with net uptake. Thus, the equilibrium H<sub>2</sub> concentration in our experiments is not just a threshold concentration where microbial uptake stops, but the isotopic evolution shows that there is an active overlapping emission (Conrad, 1994).

447

448

#### 449 **3.2 Emission and uptake strength of H**<sub>2</sub>

450

The production rate  $P = P_{true}$  and uptake rate constant  $k = k_{true}$  were obtained by applying exponential fits to the temporal evolution of H<sub>2</sub>, and applying the corrections derived from the flask sampling model (appendix A) to the  $P_{app}$  and  $k_{app}$  obtained from the exponential fits (Fig. 6). The deposition velocity ( $v_d$ ), 455 production flux  $(F_p)$ , initial uptake flux  $(F_u)$  and net flux at the beginning of the 456 experiment  $(F_n)$  were then calculated as follows:

457

 $v_d = kh \tag{7}$ 

458

$$F_p = \frac{Ph}{V_{\rm M}} \tag{8}$$

459

$$F_u = \frac{kc_1h}{V_M} \tag{9}$$

460

$$F_n = F_p - F_u \tag{10}$$

461

where h,  $V_M$  and  $c_1$  are the chamber height, standard molar volume (=22.4 L mol<sup>-1</sup>) and H<sub>2</sub> mole fraction of the first flask, respectively. We note that with our method we derive  $v_d$  as deposition velocity for the gross uptake, unlike most of the results reported in the literature that just measured net uptake.

466

The strongest soil uptake occurs in the Speuld experiments (Table 1a), with a mean  $v_d$  of (0.17±0.02) (2 SE, n=12) cm s<sup>-1</sup> (SE represents standard error). On average, the Cabauw experiments show weaker soil uptake, with a mean  $v_d$  of (0.13±0.06) (2 SE, n=8) cm s<sup>-1</sup> for the net-uptake experiments (Table 1b) and 471 (0.06±0.03) (2 SE, n=9) cm s<sup>-1</sup> for the net-emission experiments (Table 2). In 472 terms of the net H<sub>2</sub> flux  $F_n$ , this is (-26.5±4.8) (2 SE, n=12) nmol m<sup>-2</sup> s<sup>-1</sup> for 473 Speuld experiments (Table 1a), (-13.6±8.6) (2 SE, n=8) nmol m<sup>-2</sup> s<sup>-1</sup> for Cabauw 474 net-uptake experiments (Table 1b) and (49.5±29.8) (2 SE, n=9) nmol m<sup>-2</sup> s<sup>-1</sup> for 475 Cabauw net-emission experiments (Table 2), indicating strong uptake, weaker 476 uptake and strong emission of H<sub>2</sub>, respectively.

477

478

#### 479 **3.3 Fractionation during soil uptake**

480

Soil uptake and soil emission have opposite effects on the isotopic composition of  $H_2$  and can partly cancel each other. This will lead to additional uncertainty and we expect to obtain the most robust fractionation factor for soil uptake when the soil uptake is larger than the soil emission (Table 1a&b).

485

The resulting  $\alpha_{soil}$  for Speuld (Table 1a) varies from 0.913 to 0.955, with a mean value of 0.937±0.008 (2 SE, n=12). Error estimates for HH and HD mole fraction at time t and at equilibrium are considered for the final error estimates of  $\alpha_{soil}$  for each experiment.

Table 1b shows  $\alpha_{soil}$  of the Cabauw net-uptake experiments. It should be noted 491 that the soil emitted H<sub>2</sub> interferes much more with the fractionation during 492 uptake in these Cabauw net-uptake experiments than for the Speuld experiments, 493 which is illustrated by the consistent decrease in  $\delta D$  in the middle panel of Fig. 494 6. The derived values for  $\alpha_{\rm soil}$  vary from 0.911 to 1.019 with a mean value of 495 0.951±0.026 (2 SE, n=8) for these 8 selected Cabauw net-uptake experiments. 496 497 Both the mean and the standard error are higher than for the Speuld experiments  $(0.937 \pm 0.008)$ , but the difference is not significant at the 0.1 confidence level. 498

499

To graphically illustrate the calculation of  $\alpha_{soil}$  with the mass balance model, we 500 plot  $\ln \frac{c'-c'_{e,app}}{c'_1-c'_{e,app}}$  versus  $\ln \frac{c-c_{e,app}}{c_1-c_{e,app}}$  for all Speuld and Cabauw net-uptake 501 experiments in Fig. 7. A linear fit is applied to all the data and the overall  $\alpha_{soil,app}$ 502 is found to be 0.947±0.004 (95% CI). Applying a correction factor is not 503 504 straightforward now because this analysis combines the results from different experiments. If we use the average of  $\alpha_{\text{soil,true}}/\alpha_{\text{soil,app}}$  ratios (0.998) for all net-505 506 uptake experiments in Table 3 as the correction coefficient for this overall  $\alpha_{\text{soil,app}}$ , the overall  $\alpha_{\text{soil}}$  is 0.945±0.004 (95% CI). 507

508

Fig. 8 shows  $\alpha_{soil}$  as a function of  $v_d$  for all Speuld experiments and Cabauw netuptake experiments. The R<sup>2</sup> value is nearly zero and the p-value is 0.53 for the linear regression of all experiments, so no significant correlation between  $\alpha_{soil}$  512 and  $v_d$  is found. Also, no significant correlation is found when considering the 513 Speuld and Cabauw net-uptake experiments separately.

514

515

## 516 **3.4 Isotopic signature of H<sub>2</sub> emitted from soil**

517

As discussed in Section 2.4, the isotopic signature of  $H_2$  emitted from the soil 518  $(\delta D_{soil})$  can be obtained from the mass balance model. In order to minimize the 519 influence of soil uptake on the computed  $\delta D_{soil}$  and obtain the most robust result, 520 521 we only consider the Cabauw experiments with strong soil emission and weak soil uptake ( $c_{e,app} > 1500$  ppb). In total, 9 Cabauw experiments are selected 522 (Table 2) and a linear fit is applied to the plot of  $c'_{e,app} \ln \frac{c'-c'_{e,app}}{c'_1-c'_{e,app}}$  versus 523  $c_{e,app} \ln \frac{c - c_{e,app}}{c_1 - c_{e,app}}$  for each experiment (Fig. 9). It can be seen that the linear 524 function fits the data very well for each experiment. The slope of the linear fit 525 yields  $P'_{app}/P_{app}$ . This  $P'_{app}/P_{app}$  ratio is used to calculate  $\delta D_{soil,app}$  (Eq. (5)). After 526 correcting for the flask sampling effects (see Appendix A), the corresponding 527  $\delta D_{soil}$  values are shown in Table 2. The  $\delta D_{soil}$  value ranges from -629 % to -528 451 ‰, with a mean value of (-530±40) ‰ (2 SE, n=9), which is very D-529 depleted, but still considerably enriched relative to the value around -700 % 530 expected for thermodynamic equilibrium between H<sub>2</sub> and H<sub>2</sub>O (Bottinga, 1969). 531

532

533

# 534 **4. Discussion**

535

## 536 4.1 Emission and uptake strength of H<sub>2</sub>

537

The deposition velocity  $v_d$  is a measure of the strength of soil uptake. Both microbial removal and diffusion can affect  $v_d$ , and they can both be influenced by the temperature and moisture content of the soil (Ehhalt and Rohrer, 2013a; 2013b). On average, the  $v_d$  obtained in this study is larger in the forest region (Table 1a) than in the grass/clover region (Table 1b and 2), in agreement with the conclusion from Ehhalt and Rohrer (2009).

544

The  $v_d$  of (0.06±0.03) cm s<sup>-1</sup> found in our Cabauw net-emission experiments 545 (Table 2) is similar to those reported in Conrad and Seiler (1980) (0.07 cm s<sup>-1</sup>, 546 both grass and clover) and Gerst and Quay (2001) (0.04 cm s<sup>-1</sup>, grass), while the 547  $v_{\rm d}$  of (0.13±0.06) cm s<sup>-1</sup> in Cabauw net-uptake experiments (Table 1b) is larger 548 than those studies with similar soil cover but close to values of 0.12 to 0.14 cm 549 s<sup>-1</sup> found in savanna soil (Conrad and Seiler, 1985). The stronger soil uptake in 550 Speuld forest ( $(0.17\pm0.02)$  cm s<sup>-1</sup>) agrees well with the beech forest results (0.06) 551 to 0.22 cm s<sup>-1</sup>) in Förstel (1988) and Förstel and Führ (1992). However, other 552

studies at forest sites cited in Ehhalt and Rohrer (2009) showed lower  $v_d$  than our Speuld results. We note here that the  $v_d$  values reported in Conrad and Seiler (1980; 1985) were gross deposition velocities while those reported in Gerst and Quay (2001) were net deposition velocities. The specific method used to obtain  $v_d$  was not documented in the other studies.  $v_d$  obtained from our experiments are gross deposition velocities.

559

The net uptake flux  $F_n$  in our Speuld experiments and Cabauw net-uptake 560 experiments is much larger than those found in Smith-Downey et al. (2008). 561 They found a  $F_n$  of about -8 nmol m<sup>-2</sup> s<sup>-1</sup> for the forest, desert, and marsh, which 562 was similar to that for loess loamy soil in Schmitt et al. (2009). Our results are 563 within the  $F_n$  range found in the mixed wood plains by Constant et al. (2008b) 564 and the Harvard forest by Meredith (2012). Previously at our Cabauw site, Popa 565 et al. (2011) obtained a  $F_n$  of only -3 nmol m<sup>-2</sup> s<sup>-1</sup> by using the radon tracer 566 567 method. However, the Cabauw net-uptake experiments used for this evaluation were from selected places where uptake was strong, while the results in Popa et 568 569 al. (2011) represented the overall uptake in the footprint of the Cabauw site, which is a much larger area (tens of  $km^2$ ). 570

571

572 Khdhiri et al. (2015) performed microbiological analyses on soil samples from 573 the Cabauw and Speuld sites, in order to find the drivers of soil  $H_2$  uptake. They 574 observed that the  $H_2$  uptake rate under standard incubation conditions was

significantly lower for the Cabauw soil samples than for the Speuld ones, which is consistent with our findings. The main factors that explained the differences were the relative abundance of high affinity  $H_2$ -oxydizing bacteria and the soil carbon content, both lower on average for the Cabauw site.

579

580 The emission of  $H_2$  from the soil is large for the Cabauw net-emission experiments, with  $F_n$  ranging from 13.7 to 150.2 nmol m<sup>-2</sup> s<sup>-1</sup> and a median 581 value of 41.0 nmol m<sup>-2</sup> s<sup>-1</sup> (Table 2). One experiment, "CBW-28", shows 582 unusually high emission, with H<sub>2</sub> increasing to 3010 ppb within 30 minutes. In 583 comparison, Conrad and Seiler (1980) found a  $F_n$  of 23-32 nmol m<sup>-2</sup> s<sup>-1</sup> for a 584 clover field. Except for the experiments "CBW-28" and "CBW-31", our 585 Cabauw net-emission experiments are close to the  $F_n$  found by them. The 586 variability in  $F_n$  could be attributed to different N<sub>2</sub> fixation flux in our 587 experiments, which could be affected by both spatial density of  $N_2$  fixation 588 organisms and their N<sub>2</sub> fixation activities. The N<sub>2</sub> fixation activity could be 589 regulated by various factors including temperature, moisture, light availability 590 591 and carbon storage etc. (Belnap, 2001), which were not measured are therefore not discussed here. 592

593

594

#### 595 4.2 Fractionation during soil uptake

597 Fractionation during soil uptake of H<sub>2</sub> can happen during the diffusion into the soil and due to microbial removal within the soil. To further investigate the 598 factors determining  $\alpha_{soil}$ , information about the soil cover is provided in Table 599 1a&b. It is evident that no large differences exist between the Douglas fir, 600 601 spruce and beech sites, i.e. the variability between sites is similar to the 602 variability within sites. The small number of experiments impedes examining the possible small differences between sites. In order to investigate the diffusion 603 effect, we removed the soil cover in experiments "SPU-8" and "SPU-12" at the 604 605 same place of experiments "SPU-7" and "SPU-11". The removal of leaves ("SPU-8") and needles ("SPU-12") increased  $\alpha_{soil}$  by  $\approx$  0.014, thus towards 606 smaller fractionation, which indicates that diffusion contributes to the 607 fractionation. As  $v_d$  also increases when the soil cover is removed, faster 608 deposition is associated with smaller fractionations in these experiments, which 609 610 is similar to the results from Rice et al. (2011).

611

The  $\alpha_{soil}$  for the Cabauw net-uptake experiments is higher and more scattered than that for the Speuld experiments (0.951±0.026 vs. 0.937±0.008). This could be caused by the interference of D-depleted H<sub>2</sub> from the strong soil emission in Cabauw, which may not be perfectly captured via the mathematical models applied. As can be seen from the strong decline of  $\delta D$  with time in the middle panel of Fig. 6, though soil uptake of H<sub>2</sub> dominates for the Cabauw net-uptake experiments, soil production is still considerable. If part of the source signature is not taken into account properly and appears in  $\alpha_{soil}$ , then  $\alpha_{soil}$  will be larger, because soil production tends to decrease  $\delta D$  of H<sub>2</sub>. This could explain why  $\alpha_{soil}$ is even larger than 1 in "CBW-7".

622

The overall  $\alpha_{soil}$  (0.945) obtained by plotting  $\ln \frac{c' - c'_{e,app}}{c'_1 - c'_{e,app}}$  versus  $\ln \frac{c - c_{e,app}}{c_1 - c_{e,app}}$  and 623 applying the average correction factor for all the Speuld and Cabauw net-uptake 624 experiments is similar to the results of 0.943±0.024 from Gerst and Quay (2001) 625 and 0.94±0.01 from Rahn et al. (2002a). They suggested that the overall  $\alpha_{soil}$  is 626 627 more accurate as it is less susceptible to outliers. We argue here that the average  $\alpha_{soil}$  of all individual experiments in Speuld (0.937) and Cabauw (0.951) is 628 representative for a spatially averaged fractionation factor for those sites and is 629 useful for e.g. characterizing the phenomenon and comparing with other 630 631 fractionation results. If all experiments are included in one fit, their weight for determining the slopes depends on how much H<sub>2</sub> has been removed, so 632 experiments with a lower  $c_{\rm e,app}$  have a larger weight than experiments with a 633 higher  $c_{e,app}$  (i.e. experiments with a higher  $v_d$  have a larger weight than 634 experiments with a lower  $v_d$ ). The fractionation factor obtained by fitting all 635 data together is therefore representative for a flux weighted average, which is 636 637 the relevant number for the global atmospheric isotope budget.

639 **4.3 Relationship between**  $\alpha_{soil}$  and  $v_d$ 

640

Rice et al. (2011) proposed a significant positive correlation between  $\alpha$  and 641 deposition velocity  $v_d$  in their soil uptake experiments. Fig. 8 shows that no 642 significant correlation between  $\alpha_{soil}$  and  $v_{d}$  is found when considering all Speuld 643 and Cabauw net-uptake experiments. The uptake rate is much stronger in the 644 Speuld experiments ( $v_d \approx 0.17 \text{ cm s}^{-1}$ ) than in the study of Rice et al. (2011) ( $v_d$ 645  $\approx$  0.04 cm s<sup>-1</sup>), but the  $\alpha_{soil}$  is virtually identical (0.937 versus 0.934). Therefore, 646 when the results from both studies are combined, the correlation reported in 647 Rice et al. (2011) between  $\alpha_{soil}$  and  $v_d$  disappears. We suggest that a positive 648 correlation between  $\alpha_{soil}$  and  $v_{d}$  may exist for a specific site where microbial 649 species are similar. This was suggested from the simultaneous increase of both 650  $\alpha_{soil}$  and  $v_d$  in two experiments ("SPU-8" and "SPU-12"), when soil cover was 651 652 removed at the same sampling location, as mentioned in Section 4.2.

653

We conclude that there is certainly not one single correlation between  $\alpha_{soil}$  and  $v_d$  that holds globally and the soil type might play an important role. Measurements at more sites may be needed to positively confirm whether local positive correlations between  $\alpha_{soil}$  and  $v_d$  are common.

658

#### 660 **4.4** $\delta$ **D** of H<sub>2</sub> emitted from the soil

661

The present study is the first field study to report  $\delta D$  of H<sub>2</sub> emitted from soils. 662 The  $\delta D_{soil}$  values (-629 % to -451 %) shown in Table 2 are less depleted than 663 the H<sub>2</sub> in isotopic equilibrium with water ( $\approx$ -700 %). Previous observations 664 665 from environmental H<sub>2</sub> production yielded a  $\delta D$  of -628 ‰ for two seawater samples (Rice et al., 2010), -778 % for a termite headspace sample and -690 % 666 667 for two headspace samples from a eutrophic water pond (Rahn et al., 2002b). Kawagucci et al. (2015) proposed that microbiological H<sub>2</sub> consumption and 668 production could destroy the thermal isotopic equilibrium between  $H_2$  and  $H_2O$ 669 670 in low-temperature hydrothermal fluids. Luo et al. (1991) and Walter et al. (2012) found fractionation factors of 0.448, 0.401 and 0.363 for  $\rm H_2$  generated 671 672 from water by different  $N_2$ -fixing bacteria in the laboratory.

673

In order to compare our  $\delta D_{soil}$  with the fractionation factors between H<sub>2</sub> and H<sub>2</sub>O found by Luo et al. (1991) and Walter et al. (2012), we converted their fractionation factors to  $\delta D(H_2)$  by assuming the  $\delta D(H_2O)$  to be the same as that of global rainwater (-37.8 ‰, Hoffmann et al., 1998). This results in  $\delta D(H_2)$ values of -651 ‰ to -569 ‰ for their N<sub>2</sub>-fixing bacteria. Although the ranges are considerable, it appears that the mean  $\delta D_{soil}$  (-530 ‰) obtained in our field 680 study is even higher than what was found for nitrogenase-derived  $H_2$  in 681 laboratory experiments.

682

It is known that H<sub>2</sub> produced by biogenic N<sub>2</sub> fixation can be largely recycled 683 within the soil before entering the atmosphere (Evans et al., 1987; Conrad and 684 Seiler, 1979; 1980). If this uptake process within the soil tends to increase the 685  $\delta D$  of the remaining H<sub>2</sub>, as the soil uptake process for atmospheric H<sub>2</sub> does, then 686 the  $H_2$  entering the atmosphere will be less D-depleted than pure biogenic  $H_2$ . 687 688 However, if the fractionation factor of removal in the soil is similar to that determined from the net-uptake experiments ( $\approx 0.94$ ), a large fraction of H<sub>2</sub> 689 needs to be removed in the soil before release to explain the D-enriched  $\delta D_{soil}$ 690 691 compared to the values reported in the literature.

692

The deuterium enrichment in the emitted H<sub>2</sub>, compared to the value expected in 693 694 isotopic equilibrium with water, could also be caused by different fractionations induced by different enzymes and/or a potentially enriched deuterium content of 695 the substrate water available for H<sub>2</sub> production in Cabauw. H<sub>2</sub> is generated from 696 the reduction of hydrogen ions (H<sup>+</sup> or D<sup>+</sup>) in intracellular water (Yang et al., 697 698 2012). It was found that the isotopic composition of intracellular water can be 699 different from that of extracellular water due to metabolic processing (Kreuzer-700 Martin et al., 2006). Due to the differences in H-bonding and hydrogen ion transport, the fractionation may be different for different microbe species, which could result in different isotopic signatures of the produced  $H_2$ . Measurements of the isotopic composition of produced  $H_2$  may be a tool to investigate such effects.

705

Finally, we note that if our Cabauw net-emission experiments are analyzed with a simple Keeling plot approach (i.e. without considering uptake), the y axis intercept is -703 ‰. We know from the temporal evolution of H<sub>2</sub>, HD and  $\delta$ D that this model is not adequate and that uptake was significant in our experiments, so a simple Keeling plot analysis can be misleading if uptake is not considered.

712

713

# 714 **5.** Conclusions

715

This study investigated the isotope effects associated with the production and uptake of atmospheric  $H_2$  by soil. Our aim was to quantify the fractionation factor  $\alpha_{soil}$  for  $H_2$  deposition and the isotopic signature of  $H_2$  emitted from the soil ( $\delta D_{soil}$ ) from experiments carried out at Speuld and Cabauw.

The experiments covered a wide range of conditions from situations with very strong net  $H_2$  uptake to situations with very strong net  $H_2$  emission. The superposition of deposition and production made the analysis with simple models like Rayleigh plot and Keeling plot impossible. Therefore, the mass balance model suggested by Rice et al. (2011) was used for evaluation.

726

The deposition velocity  $v_d$  was largest in the Speuld experiments ((0.17±0.02) cm s<sup>-1</sup>) where also the strongest net soil uptake occurred, followed by the Cabauw net-uptake experiments ((0.13±0.06) cm s<sup>-1</sup>) and Cabauw net-emission experiments ((0.06±0.03) cm s<sup>-1</sup>). The net H<sub>2</sub> flux  $F_n$  was (-26.5±4.8) nmol m<sup>-2</sup> s<sup>-1</sup> for Speuld experiments, (-13.6±8.6) nmol m<sup>-2</sup> s<sup>-1</sup> for Cabauw net-uptake experiments and (49.5±29.8) nmol m<sup>-2</sup> s<sup>-1</sup> for Cabauw net-emission experiments.

734

The mean fractionation factors  $\alpha_{soil}$  are 0.937±0.008 for the Speuld forest soil 735 736 experiments and 0.951±0.026 for the Cabauw grassland experiments, which are representative for a spatial average and useful for comparisons with other 737 fractionation studies. The Cabauw results may be affected by the relatively 738 strong concomitant soil emissions. The overall  $\alpha_{soil}$  by considering all net-739 740 uptake experiments is  $0.945\pm0.004$ , which is representative for a flux weighted average and useful for global isotope budget estimates. The fractionation factors 741 found in this work are in good agreement with previous studies. 742
No significant correlation between  $\alpha_{soil}$  and deposition velocity  $v_d$  was found 744 while considering all of our experiments. The  $v_d$  were overall much larger in our 745 study than those in Rice et al. (2011) and we obtained similar values for  $\alpha_{soil}$ . 746 This demonstrates that the positive correlation that was found previously does 747 not hold globally. From two of our Speuld experiments,  $\alpha_{soil}$  increased after the 748 749 removal of leaves or needles above the soil. This indicates that there may be a 750 fractionation associated with diffusion through the surface layer of leaves or 751 needles during soil uptake, but more experiments are required to confirm this.

752

753 The isotopic analysis clearly showed that the net uptake was always a 754 superposition of a larger gross uptake and a gross emission flux. In Cabauw, the 755 emission strength was very large at locations where clover was present. Using a simple mass balance approach, the isotopic composition of the emitted H<sub>2</sub> was 756 757 determined to be  $(-530\pm40)$  ‰, which is significantly higher than the value expected for H<sub>2</sub>O - H<sub>2</sub> isotope equilibrium. Although limited, other published 758 data on H<sub>2</sub> produced biologically via nitrogenase show also a tendency to more 759 enriched values. An additional isotope enrichment in our field soil study could 760 originate from fractionation during the recycling of H<sub>2</sub> within the soil before it 761 762 enters the atmosphere.

## 764 Appendix A

765

# 766 A1. Flask sampling model

767

A mathematical model is used to simulate the sampling and to correct for the 768 effects of the flask sampling method on the values of uptake rate constant (k), 769 production rate (P), fractionation factor ( $\alpha_{soil}$ ) and isotopic signature of H<sub>2</sub> 770 771 produced from soil ( $\delta D_{soil}$ ). We start with a pair of known (*true*) uptake and production rates and simulate the evolution of the mole fractions of H<sub>2</sub> and HD 772 in the flasks and chamber. From the modeled mole fractions we calculate the 773 apparent uptake and production rates and derive the correction needed to obtain 774 the *true* uptake and production rates from measurement of the *apparent* rates in 775 776 actual experiments.

777

### 778 A1.1 Mathematical description of the flask sampling model

779

The sampling setup is shown in Fig. 2 of the main paper. After 10 minutes of flushing, the chamber and the flasks contain ambient air with the prevailing H<sub>2</sub> and HD mole fractions. In the following we denote  $c_1(t)$ ,  $c_2(t)$ ,  $c_3(t)$ ,  $c_4(t)$  and  $c_0(t)$  the H<sub>2</sub> mole fractions for the first, second, third, forth flask and the chamber, respectively. The moment when the first flask and the chamber lid are closed is considered the starting time of the experiment (t=0). From this point on, only the chamber, the second, third and fourth flask are connected, and the initial H<sub>2</sub> mole fraction inside them is  $c_0(0) = c_2(0) = c_3(0) = c_4(0) = c_1$ . We start a simulation with an input uptake rate constant ( $k_{true}$ ) and an input production rate ( $P_{true}$ ). The simulation of the flask sampling is based on Eqs. (A1)-(A4) shown below.

791

Assuming that the air in each flask and in the chamber is well-mixed during the entire sampling process, the time evolution for the second flask  $c_2(t)$ , the third flask  $c_3(t)$ , the forth flask  $c_4(t)$  and the chamber  $c_0(t)$  in the first 10 minutes after starting the experiment can be expressed as:

796

$$\frac{\mathrm{d}c_2(t)}{\mathrm{d}t} = \frac{f}{V}c_0(t) - \frac{f}{V}c_2(t) \tag{A1}$$

797

$$\frac{dc_3(t)}{dt} = \frac{f}{V}c_2(t) - \frac{f}{V}c_3(t)$$
 (A2)

798

$$\frac{dc_4(t)}{dt} = \frac{f}{V}c_3(t) - \frac{f}{V}c_4(t)$$
(A3)

799

$$\frac{\mathrm{d}c_0(t)}{\mathrm{d}t} = \frac{f}{V'}c_4(t) - \frac{f}{V'}c_0(t) + (P_{\mathrm{true}} - k_{\mathrm{true}}c_0(t)) \tag{A4}$$

where V and V' are the air volumes of the flask and chamber, and f is the flow rate. These differential equations are solved using the Matlab ODE solvers at time steps of 0.01 min. The input parameters are  $c_0(0)$ ,  $P_{true}$ ,  $k_{true}$ , V, V' and f. For each time step the solvers calculate the hydrogen flux into and out of the chamber and each flask, as well as the new mole fractions there.

806

After 10 minutes, the second flask is closed and now contains air with mole fraction  $c_2 = c_2(10 \text{ min})$ . From this point on, only the chamber, the third and the fourth flask are connected, and the time evolution of the mole fractions can be expressed as:

811

$$\frac{dc_3(t)}{dt} = \frac{f}{V}c_0(t) - \frac{f}{V}c_3(t)$$
(A5)

812

$$\frac{\mathrm{d}c_4(t)}{\mathrm{d}t} = \frac{f}{V}c_3(t) - \frac{f}{V}c_4(t) \tag{A6}$$

813

$$\frac{dc_0(t)}{dt} = \frac{f}{V'}c_4(t) - \frac{f}{V'}c_0(t) + (P_{\text{true}} - k_{\text{true}}c_0(t))$$
(A7)

814

After another 10 minutes of sampling, the third flask is closed  $c_3 = c_3(20 \text{ min})$ , and only the chamber and the fourth flask are connected. Then, the time evolution for the fourth flask and the chamber can be expressed as:

$$\frac{dc_4(t)}{dt} = \frac{f}{V}c_0(t) - \frac{f}{V}c_4(t)$$
(A8)

819

$$\frac{dc_0(t)}{dt} = \frac{f}{V'}c_4(t) - \frac{f}{V'}c_0(t) + (P_{\text{true}} - k_{\text{true}}c_0(t))$$
(A9)

820

The H<sub>2</sub> mole fraction inside the chamber and the fourth flask at time t=30 minutes is  $c_0(30)$  and  $c_4(30)$ .

823

In the end, a set of four flasks with mole fractions  $c_1(0)$ ,  $c_2(10 \text{ min})$ ,  $c_3(20 \text{ min})$ 824 and  $c_4(30 \text{ min})$  is obtained. By fitting this set of four data points with an 825 exponential function  $c = ae^{-k_{app}t} + c_{e,app}$  (see Eq. (2) in the main paper), we 826 can obtain the *apparent* soil uptake rate constant  $(k_{app})$  and equilibrium 827 concentration  $(c_{e,app})$  and further calculate *apparent* production rate 828  $(P_{app}=k_{app}c_{e,app})$ . These apparent rates  $k_{app}$  and  $P_{app}$  are different from the assumed 829 true rates  $k_{true}$  and  $P_{true}$ . The flask sampling model enables us to establish a 830 relation between  $k_{app}$  and  $P_{app}$  and  $k_{true}$  and  $P_{true}$ , so that  $k_{true}$  and  $P_{true}$  can be 831 derived from  $k_{\rm app}$  and  $P_{\rm app}$  in actual experiments, where the true values are 832 unknown. To accomplish this, simulations are carried out with a wide range of 833 values for  $k_{\text{true}}$  and  $P_{\text{true}}$ , and a corresponding dataset of  $k_{\text{app}}$  and  $P_{\text{app}}$  is generated. 834

Similarly, we use a new set of input uptake rate constant  $k'_{true}$  and production rate  $P'_{true}$  for HD, and generate a corresponding dataset of  $k'_{app}$  and  $P'_{app}$ .

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#### 838 A1.2 The correction coefficients for *k* and *P*

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Here we discuss an example of the relationship between  $k_{\text{true}}$  and  $k_{\text{app}}$  for the 840 setup used in some Cabauw experiments (V'=22.8 L, f=2 L min<sup>-1</sup> and  $\Delta t=10$ 841 min). The pressure inside the flasks is 200 kPa and the pressure inside the 842 chamber is 100 kPa. The relationship between  $k_{true}/k_{app}$  and  $k_{app}$  is shown in Fig. 843 10a. The ratio  $k_{true}/k_{app}$  varies between 1.45 to 1.61 for our  $k_{app}$  range of 0.04 to 844 0.30 min<sup>-1</sup>. This relationship does not depend on  $P_{\text{true}}$  (with  $P_{\text{true}}$  varying from 50 845 to 650 ppb min<sup>-1</sup>). An additional uncertainty can arise from incorrect timing of 846 the flask sampling, but sampling times should be correct within few seconds, 847 which may lead to an additional uncertainty of below 1%. The uncertainty of 848 849 the flow rate obtained from the rotameter due to variations in ambient pressure 850 and temperature that were not recorded is less than 4%, and the effect on the ratio  $k_{true}/k_{app}$  ratio is below 1%. We can retrieve  $k_{true}$  by multiplying  $k_{app}$  with the 851 modeled value of  $k_{\rm true}/k_{\rm app}$  for each experiment. The ratio  $k_{\rm true}/k_{\rm app}$  for each 852 experiment is shown in Table 3. It depends on experimental setup and  $k_{\rm app}$  of 853 854 each experiment, with a range of 1.177 to 1.589.

After retrieving  $k_{true}$  from  $k_{app}$ , we investigate the relationship between  $P_{true}/P_{app}$ 856 and  $P_{app}$  for a fixed value of  $k_{true}$  (Fig. 10b). The ratio  $P_{true}/P_{app}$  depends slightly 857 on  $P_{\rm app}$  and  $k_{\rm true}$ , ranging from 1.40 to 1.59 for a wide  $P_{\rm app}$  range of 30 to 450 858 ppb min<sup>-1</sup> and a wide  $k_{true}$  range of 0.05 to 0.45 min<sup>-1</sup>. As for the correction of k, 859 uncertainties arising from incorrect timing of the flask sampling and from 860 pressure and temperature variations and their effect on the flow rate lead to 861 additional uncertainties of  $P_{true}/P_{app}$  ratio below 1%, which are not considered. 862 We can retrieve  $P_{\text{true}}$  by multiplying  $P_{\text{app}}$  with  $P_{\text{true}}/P_{\text{app}}$  for each experiment after 863 having determined  $k_{\text{true}}$  from  $k_{\text{app}}$ . The ratio  $P_{\text{true}}/P_{\text{app}}$  for each experiment is 864 shown in Table 3 and depends on the experimental setup,  $P_{app}$  and  $k_{app}$  of each 865 experiment. It ranges from 1.152 to 2.759 for most experiments, with an 866 exception of 7.472 for experiment SPU-2 where a very small  $P_{app}$  of 0.67 ppb 867 min<sup>-1</sup> is found. Although the ratio  $P_{true}/P_{app}$  of experiment SPU-2 is high,  $P_{true}$  of 868 SPU-2 is still smaller than the rest of the experiments.  $P_{true}/P_{app}$  ratios for 869 870 experiments SPU-10 and SPU-11 are null because these two experiments show a  $P_{\rm app}$  of zero. 871

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#### 873 A1.3 The correction coefficients for $\alpha_{soil}$ and $\delta D_{soil}$

874

In our experiments, the uncertainties of  $k_{app}$  and  $k'_{app}$  derived from exponential fits to the time evolution of HH and HD are rather large, which results in a large scatter of  $\alpha_{soil,app}$  if  $\alpha_{soil,app}$  is calculated directly as  $k'_{app}/k_{app}$ . Thus, we obtained

878 
$$\alpha_{\text{soil,app}}$$
 by plotting  $\ln \frac{c'-c'_{\text{e,app}}}{c'_1-c'_{\text{e,app}}}$  versus  $\ln \frac{c-c_{\text{e,app}}}{c_1-c_{\text{e,app}}}$  (Fig. 7) for each experiment

879 which yields a smaller scatter for  $\alpha_{\text{soil,app}}$ .

880

Correction coefficients to convert  $\alpha_{\text{soil,app}}$  to  $\alpha_{\text{soil,true}}$  are obtained using the flask 881 sampling model by comparing  $\alpha_{\text{soil,true}}$  used as input for the model run to  $\alpha_{\text{soil,app}}$ 882 derived from the plot of  $\ln \frac{c'-c'_{e,app}}{c'_1-c'_{e,app}}$  versus  $\ln \frac{c-c_{e,app}}{c_1-c_{e,app}}$  of the output values, like 883 in the experiments. Fig. 10c shows  $\alpha_{\text{soil,true}}/\alpha_{\text{soil,app}}$  as a function of  $\alpha_{\text{soil,app}}$  for a 884 wide  $\delta D_{\text{soil,true}}$  range of -750% to -250% with the sampling setup described 885 above (V'=22.8 L, f=2 L min<sup>-1</sup> and  $\Delta t=10$  min) for  $k_{true}=0.25$  min<sup>-1</sup> and  $P_{true}=50$ 886 ppb min<sup>-1</sup>. In this case the correction factor  $\alpha_{\text{soil,true}}/\alpha_{\text{soil,app}}$  varies from 0.98 to 887 1.00 for a  $\alpha_{\text{soil,app}}$  range of 0.90 to 1.00, and it does not depend on  $\delta D_{\text{soil,true}}$ . Thus, 888 after retrieving  $k_{\text{true}}$  and  $P_{\text{true}}$  as described in Section A1.2, we can retrieve  $\alpha_{\text{soil,true}}$ 889 from  $\alpha_{\text{soil,app}}$  for each experiment. The correction factors range from 0.984 to 890 1.007, depending on the experimental setup and  $\alpha_{\text{soil,app}}$  of each experiment 891 892 (Table 3).

893

Similarly, in our experiments, the uncertainties of  $P_{app}$  and  $P'_{app}$  derived from exponential fits of time evolution of HH and HD are large, which results in a large scatter of  $\delta D_{soil,app}$  if  $\delta D_{soil,app}$  is calculated directly from these  $P'_{app}$  and  $P_{app}$ . We therefore obtained the ratio  $P'_{app}/P_{app}$  by plotting  $c'_{e,app} \ln \frac{c'-c'_{e,app}}{c'_1-c'_{e,app}}$  versus

898 
$$c_{e,app} \ln \frac{c-c_{e,app}}{c_1-c_{e,app}}$$
 (Fig. 9) and calculated  $\delta D_{soil,app}$  from Eq. (4). This yielded  
899 smaller scatter for  $\delta D_{soil,app}$ . After retrieving  $k_{true}$ ,  $P_{true}$  and  $\alpha_{soil,true}$  as described  
900 above, we used the flask sampling model again to derived correction factors by  
901 comparing  $\delta D_{soil,true}$  used as model input with  $\delta D_{soil,app}$  obtained from  
902  $c'_{e,app} \ln \frac{c'-c'_{e,app}}{c'_1-c'_{e,app}}$  versus  $c_{e,app} \ln \frac{c-c_{e,app}}{c_1-c_{e,app}}$  of the model output, and retrieve  
903  $\delta D_{soil,true}$  from  $\delta D_{soil,app}$  for each experiment. Fig. 10d shows  
904  $(\delta D_{soil,true}+1)/(\delta D_{soil,app}+1)$  as a function of  $(\delta D_{soil,app}+1)$  for a  $\alpha_{soil,true}$  range of 0.90  
905 to 1.00 with the sampling setup described above (V'=22.8 L, f=2 L min<sup>-1</sup> and  
906  $\Delta t=10$  min) for  $k_{true}=0.25$  min<sup>-1</sup> and  $P_{true}=50$  ppb min<sup>-1</sup>. The ratio  
907  $(\delta D_{soil,true}+1)/(\delta D_{soil,app}+1)$  changes from 0.99 to 1.05 for a wide  $(\delta D_{soil,app}+1)$  ratio  
908 depends slightly on  $\alpha_{soil,true}$  at a fixed  $(\delta D_{soil,app}+1)$ , with a maximum difference  
910 of about 1% for a  $\alpha_{soil,true}$  range of 0.90 to 1.00. The ratio  
911  $(\delta D_{soil,true}+1)/(\delta D_{soil,app}+1)$  for each net-emission experiment is shown in Table 3,  
912 ranging from 1.007 to 1.048. The largest difference between  $\delta D_{soil,true}$  and  
913  $\delta D_{soil,app}$  is 21% for CBW-8. The mean  $\delta D_{true}$  and  $\delta D_{app}$  for these net emission  
914 experiments are -530% and -538% c, respectively.

916 In conclusion, the effect of the flask sampling process is relatively small for  $\alpha_{soil}$ 917 and  $\delta D_{soil}$ , but considerable for the uptake rate constants k and k' and emission 918 rates P and P'. The flask sampling model allows to derive corresponding

919	corrections that have been applied to correct for the bias introduced by the flask
920	sampling system.
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922	
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924	
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1229 Tables

1231 Table 1. The deposition velocity  $(v_d)$ , fractionation factor  $(\alpha_{soil})$  as well as its error estimate, 1232 and soil cover information for each Speuld experiment (a) and Cabauw net-uptake 1233 experiment (b). The STDEV represents standard deviation and SE represents standard error. 1234 The errors of  $\alpha_{soil}$  represent the 95% confidence interval (CI) for  $\alpha_{soil,app}$  obtained from

1235	$\ln rac{c' - c'_{e,\mathrm{app}}}{c'_1 - c'_{e,\mathrm{app}}}$	versus ln-	$\frac{c-c_{e,app}}{c_1-c_{e,app}}$
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(a)	$F_{\rm n}$ (nmol m <sup>-2</sup> s <sup>-1</sup> )	$v_{\rm d}$ (cm s <sup>-1</sup> )	$lpha_{ m soil}$	Error $a_{ m soil}$	soil cover
SPU-1	-30.1	0.20	0.924	0.032	D. fir, moss
SPU-2	-35.3	0.22	0.948	0.028	D. fir, needles
SPU-3	-37.7	0.20	0.945	0.008	D. fir, moss
SPU-4	-26.1	0.16	0.913	0.004	D. fir, moss
SPU-5	-24.9	0.16	0.918	0.006	D. fir, moss
SPU-6	-13.2	0.12	0.951	0.031	D. fir, moss
SPU-7	-19.6	0.12	0.939	0.005	beech, leaves
SDI 8	28.4	0.16	0.955	0.008	Same subsite as SPU-7,
51 0-0	-20.4	0.10	0.955	0.000	leaves removed
SPU-9	-20.4	0.12	0.925	0.002	beech, leaves
SPU-10	-22.3	0.13	0.949	0.060	spruce, moss
SPU-11	-19.4	0.13	0.936	0.068	spruce, needles
SDU 12	40.5	0.28	0.047	0.004	Same subsite as SPU-11,
Sr U-12	-40.2	0.20	0.747	0.004	needles removed
MEAN	-26.5	0.17	0.937	/	/

STDEV	8.2	0.05	0.014	/	/
SE	2.4	0.01	0.004	/	/
(b)	${F}_{ m n}$	$v_{\rm d}$	a	Frror a	soil cover
(0)	$(nmol m^{-2} s^{-1})$	$(cm s^{-1})$	$\alpha_{\rm soil}$	Error $\alpha_{soil}$	son cover
CBW-5	-6.6	0.04	0.943	0.004	few clover, grass
CBW-7	-3.1	0.03	1.019	0.005	few clover, grass
CBW-16	-22.9	0.18	0.993	0.001	bare soil, few grass
CBW-18	-39.3	0.24	0.950	0.054	grass
CBW-19	-7.4	0.14	0.935	0.105	grass
CBW-20	-14.9	0.20	0.940	0.260	bare soil
CBW-25	-8.0	0.12	0.911	0.014	clover, grass
CBW-26	-6.1	0.09	0.916	0.038	grass
MEAN	-13.6	0.13	0.951	/	/
STDEV	12.2	0.08	0.037	/	/
SE	4.3	0.03	0.013	/	/

Net	$F_{n}$	$v_{ m d}$			
emission	$(nmol m^{-2} s^{-1})$	(cm s <sup>-1</sup> )	0D <sub>soil</sub> (‰)	EITOI $OD_{\text{soil}}$ (700)	
CBW-8	24.5	0.05	-535	53	
CBW-10	16.1	0.03	-460	17	
CBW-14	13.7	0.02	-629	21	
CBW-17	20.3	0.03	-542	1	
CBW-21	42.0	0.04	-574	3	
CBW-28	150.2	0.14	-488	83	
CBW-30	41.0	0.05	-580	7	
CBW-31	92.0	0.09	-509	7	
CBW-33	46.2	0.10	-451	52	
MEAN	49.5	0.06	-530	/	
STDEV	44.7	0.04	59	/	
SE	14.9	0.01	20	/	

1238 Table 2. Net flux, deposition velocity and  $\delta D_{soil}$  (including error) obtained from the mass 1239 balance model for the net H<sub>2</sub> emission experiments.

	Pressure	Flow rate	Size	$\Delta t$	$k_{\rm app}$	$P_{ m app}$				$(\delta D_{soil,true}+1)/$
Exp.	(kPa)	$(L \min^{-1})$		(min)	$(\min^{-1})$	(ppb	$k_{\rm true}/k_{\rm app}$	$P_{\rm true}/P_{\rm app}$	$\alpha_{\rm soil,true}/\alpha_{\rm soil,app}$	$(\delta D_{soil,app}+1)$
						$\min^{-1}$ )		11		· • • •
SPU-1	200	2	S	10	0.199	4.12	1.494	1.601	0.984	/
SPU-2	200	2.2	S	5	0.206	0.67	1.589	7.472	0.998	/
SPU-3	200	3.1	S	5	0.204	3.58	1.496	2.475	0.999	/
SPU-4	200	2.8	S	5	0.160	7.51	1.526	2.136	1.004	/
SPU-5	200	2.6	S	5	0.156	4.16	1.546	2.759	1.004	/
SPU-6	160	3.2	L	5	0.232	7.61	1.184	1.446	0.999	/
SPU-7	160	3.2	S	5	0.128	5.40	1.418	2.264	1.006	/
SPU-8	160	2.5	S	5	0.172	4.23	1.438	2.381	1.001	/
SPU-9	160	2.8	S	5	0.128	4.56	1.440	2.513	1.007	/
<b>SPU-10</b>	180	2.7	S	5	0.128	/	1.502	/	1.005	/
SPU-11	160	2.2	S	5	0.130	/	1.490	/	1.006	/
SPU-12	180	2.3	S	5	0.272	11.30	1.529	1.720	0.994	/
CBW-5	200	2	L	10	0.086	18.24	1.204	1.248	1.001	/
CBW-7	200	1.9	L	10	0.048	11.57	1.260	1.361	0.999	/
CBW-16	210	2.1	S	10	0.183	45.21	1.498	1.505	0.999	/
CBW-18	200	2	S	10	0.240	38.07	1.532	1.527	0.986	/
CBW-19	200	2	S	10	0.145	56.69	1.457	1.463	0.991	/
CBW-20	200	2	S	10	0.196	65.81	1.491	1.494	0.988	/
<b>CBW-25</b>	200	2	S	10	0.122	44.85	1.449	1.460	0.994	/
CBW-26	200	2	S	10	0.088	31.05	1.452	1.475	1.002	/
CBW-8	200	2	S	10	0.044	82.92	1.542	1.438	/	1.048
CBW-10	200	2.6	L	10	0.069	111.00	1.177	1.152	/	1.010
CBW-14	200	2.5	L	10	0.035	82.53	1.251	1.166	/	1.042
CBW-17	220	2.1	L	10	0.047	117.40	1.268	1.198	/	1.024

Table 3. Sampling information and the correction coefficients  $(k_{true}/k_{app}, P_{true}/P_{app}, \alpha_{soil,true}/\alpha_{soil,app} \text{ and } (\delta D_{soil,true}+1)/(\delta D_{soil,app}+1) \text{ used}$ 

2 for each experiments. Size S refers to small chamber and size L refers to large chamber.

CBW-21	220	2	L	10	0.078	232.20	1.209	1.179	/	1.008
CBW-28	175	1.8	S	10	0.146	440.90	1.412	1.402	/	1.018
CBW-30	200	2	L	10	0.090	237.70	1.202	1.180	/	1.008
CBW-31	200	2	S	10	0.098	275.10	1.451	1.422	/	1.007
CBW-33	200	2	S	10	0.107	166.50	1.449	1.430	/	1.007

# **Figures**



Fig. 1. The location of the two sampling sites (Cabauw and Speuld) in the Netherlands, as

- 28 well as the plant species there.



1269 Fig. 2. Scheme of the sampling setup using the closed-cycle air sampler. The volume of the

- soil chamber was 22.8 L and the volume of each flask was 1 L.

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Fig. 3. Difference of  $\delta D$  from the assigned value for different gases including reference gases (Ref1-3) and laboratory flask samples (S1-7). A linear function (y = 54.6x) was fit to the data with peak area between 0.2 and 1.0 V s (green solid line; the dashed lines represent the 95% confidence interval of the fit). This function was used to correct the soil experiment data that were measured at low peak areas.



1304 Fig. 4. Calculated total assigned uncertainty of  $\delta D$  (consisting of analytical uncertainty and 1305 uncertainty arising from the linearity correction) for air samples with ln(peak area) ranging 1306 from -1.6 to 1.5.



Fig. 5. Results of the flask sampling model with the following parameters:  $k=0.1 \text{ min}^{-1}$ , P=10ppb min<sup>-1</sup> and  $c_1(t=0)=530$  ppb. The figure shows the evolution of  $H_2$  mole fraction in the chamber (green curve), in flask 2 (blue curve), flask 3 (red curve) and flask 4 (magenta curve) as a function of time, and what would be expected for a chamber without flasks (black curve). Flask 1 was closed before closing the chamber (at time 0 when all volumes contained the same air).



Fig. 6. Time evolution of H<sub>2</sub>, HD and  $\delta$ D in Cabauw (upper and middle panels) and in Speuld (lower panel) for representative experiments. HD is calculated from H<sub>2</sub> and  $\delta$ D. The H<sub>2</sub> data are fitted with an exponential function of the form:  $c = (c_1 - c_{e,app})e^{-k_{app}t} + c_{e,app}$ , where  $c_1$  and  $c_{e,app}$  are the H<sub>2</sub> mole fractions initially and in equilibrium, and  $k_{app}$  is the apparent soil uptake rate constant for H<sub>2</sub>. A similar exponential function is applied to the HD data. Error estimates for H<sub>2</sub>, HD and  $\delta$ D are shown. The connecting lines for  $\delta$ D data are included to guide the eye.





Fig. 7. Plot of  $\ln \frac{c'-c'_{e,app}}{c'_1-c'_{e,app}}$  versus  $\ln \frac{c-c_{e,app}}{c_1-c_{e,app}}$  for all Speuld and Cabauw net-uptake experiments. The slope of the linear fit to the data returns the fractionation factor  $\alpha_{soil,app}=0.947\pm0.004$  (95% CI). Errors in x and y direction for each data point were considered. One outlier ("CBW-18") was not included in the fitting. The 95% confidence intervals of the fit line are included as dashed lines but largely overlap with the fit line.



1367 Fig. 8. Correlation between  $\alpha_{soil}$  and  $v_d$  for all Speuld experiments and Cabauw net-uptake

1368 experiments. The errors for  $\alpha_{soil}$  were taken from Table 1.




1384 Fig. 9. Plot of  $c'_{e,app} \ln \frac{c'-c'_{e,app}}{c'_1-c'_{e,app}}$  versus  $c_{e,app} \ln \frac{c-c_{e,app}}{c_1-c_{e,app}}$  for 9 Cabauw net-emission 1385 experiments. A linear function was fit to each individual dataset and the slope was used to 1386 calculate the  $\delta D_{soil,app}$  value for each experiment. Errors in x and y direction for each data 1387 point were considered.

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Fig. 10. (a) The relationship between  $k_{true}/k_{app}$  and  $k_{app}$  for  $P_{true}$  of 50, 200 and 650 ppb min<sup>-1</sup>; (b) between  $P_{true}/P_{app}$  and  $P_{app}$  for  $k_{true}$  of 0.05, 0.25 and 0.45 min<sup>-1</sup>; (c) between  $\alpha_{soil,true}/\alpha_{soil,app}$  and  $\alpha_{soil,app}$  for  $(\delta D_{soil,true}+1)$  of 0.25 to 0.65 for  $k_{true}=0.25$  min<sup>-1</sup> and  $P_{true}=50$  ppb min<sup>-1</sup>; (d) between  $(\delta D_{soil,true}+1)/(\delta D_{soil,app}+1)$  and  $(\delta D_{soil,app}+1)$  for  $\alpha_{soil,true}$  of 0.90 to 1.00 for  $k_{true}=0.25$  min<sup>-1</sup> and  $P_{true}=50$  ppb min<sup>-1</sup>. The parameters of the sampling setup are V' =22.8 L, f=2 L min<sup>-1</sup>,  $\Delta t=10$ min and the pressures inside the flasks and chamber are 200 kPa and 100 kPa respectively.

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