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# Molecular hydrogen (H<sub>2</sub>) emissions and their isotopic signatures (H/D) from a motor vehicle: implications on atmospheric H<sub>2</sub>

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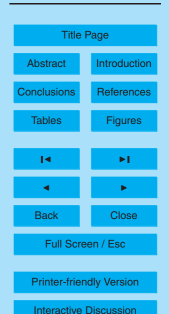
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#### Abstract

Molecular hydrogen ( $H_2$ ), its isotopic signature (deuterium/hydrogen,  $\delta D$ ), carbon monoxide (CO) and other compounds were studied in the exhaust of a passenger car engine fuelled with gasoline or methane and run under variable air-fuel ratios and 5 operating modes. H<sub>2</sub> and CO concentrations were largely reduced downstream of the three-way catalytic converter (TWC) compared to levels upstream, and showed a strong dependence on the air-fuel ratio (expressed as lambda,  $\lambda$ ). The isotopic composition of H<sub>2</sub> ranged from  $\delta D = -140\%$  to  $\delta D = -195\%$  upstream of the TWC but these values decreased to -270% to -370% after passing through the TWC. Post-TWC δD values for the fuel-rich range showed a strong dependence on TWC temperature with more negative  $\delta D$  for lower temperatures. These effects are attributed to a rapid temperature-dependent H-D isotope equilibration between H<sub>2</sub> and water (H<sub>2</sub>O). In addition, post TWC  $\delta D$  in H<sub>2</sub> showed a strong dependence on the fraction of removed H<sub>2</sub>, suggesting isotopic enrichment during catalytic removal of H<sub>2</sub> with enrichment factors  $(\varepsilon)$  ranging from -39.8% to -15.5% depending on the operating mode. Our results imply that there may be considerable variability in real-world  $\delta D$  emissions from vehicle exhaust, which may mainly depend on TWC technology and exhaust temperature regime. This variability is suggestive of a  $\delta D$  from traffic that varies over time, by season, and by geographical location. An earlier-derived integrated pure (end-member) δD from anthropogenic activities of -270% (Rahn et al., 2002) can be explained as a mixture of mainly vehicle emissions from cold starts and fully functional TWCs, but enhanced  $\delta D$  values by >50% are likely for regions where TWC technology is not fully implemented. Our results also suggest that a full hydrogen isotope analysis on fuel and exhaust gas may greatly aid at understanding process-level reactions in the exhaust gas, in particular in the TWC.

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#### 1 Introduction

The poorly understood budget of atmospheric molecular hydrogen  $(H_2)$  has received increased attention over the past years because of a potentially massive disturbance in the near future due to a potential shift towards a hydrogen energy economy. Recent studies have revealed potential negative impacts on stratospheric ozone and on earth's climate systems via the role of  $H_2$  in atmospheric chemistry; however, quantitative estimates are highly uncertain, largely due to unknown leakage rates of  $H_2$  to the atmosphere from future  $H_2$  energy systems. In the traffic sector, current  $H_2$  emissions from conventional combustion engine systems would disappear, but would likely be replaced by leakage and purged losses from  $H_2$ -powered vehicles.

 $H_2$  is abundant in the atmosphere at relatively high concentrations (hereafter expressed as dry air mol fraction) of 450 ppb–550 ppb (parts-per-billion,  $10^{-9}$ , molar). Its source and sink strengths are given here as ranges found in the literature (Novelli et al., 1999; Hauglustaine and Ehhalt, 2002; Sanderson et al., 2003; Rhee et al., 2006; Price et al., 2007; Xiao et al., 2007; Ehhalt and Rohrer, 2009): the major sources are in-situ photochemical production from methane ( $CH_4$ ) and other hydrocarbons (30–77  $Tg \, a^{-1}$ ), fossil fuel emissions (11–20  $Tg \, a^{-1}$ ), biomass burning (10–20  $Tg \, a^{-1}$ ), and minor emissions from microbial nitrogen ( $N_2$ ) fixation on land and in the oceans (6–10  $Tg \, a^{-1}$ ). The dominant sink of  $H_2$  is enzymatic destruction in soil (55–88  $Tg \, a^{-1}$ ), but there is also considerable removal through oxidation by the hydroxyl radical in the atmosphere (15–19  $Tg \, a^{-1}$ ). These fluxes result in a tropospheric lifetime of  $H_2$  of 1.4–2.2 a. Due to the dominant soil sink and the larger land coverage in the Northern Hemisphere, this hemisphere exhibits smaller tropospheric concentrations compared to the southern hemisphere, which is an unusual distribution for atmospheric trace gases.

Stable isotope measurements of  $H_2$  using the hydrogen/deuterium (H/D) ratio have greatly added to differentiating various sources and sinks (Brenninkmeijer et al., 2003; Ehhalt and Rohrer, 2009). This is particularly true because of largely differing isotopic signatures of the sources. For example,  $H_2$  from  $CH_4$  oxidation is particularly enriched

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in deuterium with  $\delta D$  values ~130% to 180% (Rahn et al., 2003; Rhee et al., 2006; Feilberg et al., 2007; Röckmann et al., 2003; Pieterse et al., 2009), biogenic H<sub>2</sub> is extraordinarily depleted ( $\delta D$ ~-700%, e.g. Rahn et al. (2002); Walter and et al. (2010)) and combustion-derived H<sub>2</sub> has  $\delta D$  values of approximately -170% to -270% (Gerst and Quay, 2001; Rahn et al., 2002).

Automobile traffic is believed to dominate anthropogenic  $H_2$  emissions, and has recently been estimated globally at  $4.2-5.4\,\mathrm{Tg\,a^{-1}}$  with a decreasing trend, based on a fleet-integrated tunnel emission study (Vollmer et al., 2007). The single available isotope study of regional traffic emissions using polluted air samples from the Los Angeles basin by Rahn et al. (2002) suggests an urban pollution isotopic signature of -270% for this region. Samples taken near the exhaust of individual vehicles in their study showed considerable variability in the isotopic signature, as did a few samples collected in a parking garage by Gerst and Quay (2001), which showed less depleted  $\delta D$ .

Engine and catalytic converter technology is evolving rapidly, and along with the increasing diversity of fuel types, presents a challenge for vehicle emissions characterization. Such studies can be conducted on various levels of detail, and range from regional scale atmospheric observations of polluted air masses to single unit process studies. Isotope investigations can greatly improve the distinction from other sources and sinks of pollutants. For vehicle exhaust, progress has recently been achieved through isotopic studies of e.g. carbon monoxide, CO (Tsunogai et al., 2003), methane, CH<sub>4</sub> (Chanton et al., 2000; Nakagawa et al., 2005), and nitrous oxide, N<sub>2</sub>O (Toyoda et al., 2008). Here we present results from a study on a passenger car engine for which we have measured pre- and post-catalytic H<sub>2</sub> concentrations and – to our knowledge for the first time – H/D signatures under variable engine and fuel settings.

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#### 2 Materials and methods

#### 2.1 Experimental setup

The engine experiment was conducted in 2008 on an engine test bench at the Empa Internal Combustion Engine Laboratory using a naturally-aspirated passenger car engine with four cylinders, 2-L displacement, and the capability to combust either gasoline or natural gas. The engine was equipped with a state-of-the art engine control unit, which was originally calibrated to achieve Euro-4 emission limits for a mid-size passenger car. On the test bench, the control parameters could be modified to study e.g. the influence of the air-to-fuel ratio on combustion and emissions. The combustion air fed to the engine was conditioned to a temperature of 295 K and a relative humidity of 50%. The exhaust system was equipped with a single three-way ceramic substrate catalytic converter (TWC) with a cell density of 600 cells per square inch (cpsi) and coated with palladium and rhodium as the catalytically active materials. Ceria, an oxide of the element cerium, was used in the TWC's washcoat to achieve the desired oxygen storage capacity. For the sake of the later temperature discussion, it is worthy to mention that a TWC is not actively heated but that it is initially cold (termed "cold start" in traffic studies) after engine start-up until sufficient hot exhaust gas passes through it to make it fully functional (light-off temperature). For air-to-fuel ratio feedback control, the engine was equipped with a linear lambda ( $\lambda$ ) sensor upstream of the catalyst and a switching-type  $\lambda$  sensor downstream of the TWC for bias control. Linear  $\lambda$  sensors are able to measure deviations of  $\lambda$  very precisely but they unfortunately show a relatively slow signal drift, which depends on many factors. To compensate for this drift, the bias of the linear  $\lambda$  sensor is corrected occasionally using the signal of a much simpler switching-type  $\lambda$  sensor mounted downstream of the TWC. Such switching-type sensors show a very stable and steep signal change at  $\lambda=1$  but no useful information of the actual  $\lambda$  value in fuel-rich or fuel-lean operating conditions. The engine control used a square signal " $\lambda$  – wobbling" function with an amplitude of 0.02 to the nominal λ setting. These rich-lean excursions are very often used in modern engine controls

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to enhance the TWC's efficiency, and to perform diagnosis of the TWC during engine operation using the TWC's dynamic response to this wobbling.

The experiment was conducted for 4 different operating modes (OMs) as is shown in Table 1, where OM-1 used the same engine settings as OM-2, but in the first case the TWC was cooled by active air ventilation of the exhaust manifold and the piping upstream of the TWC. All OMs reflect relatively low engine power, which are frequently used in normal operations of passenger cars. OM-2 is a standard setting, which is widely used in engine research for comparative purposes. OM-3 and OM-4 differ from OM-2 in torque and crankshaft speed and represent low rotation per minute (rpm) and high-rpm driving, both at double the engine power of OM-1 and OM-2. In order to study the air-to-fuel settings on the pollutant emissions before and after TWC, the  $\lambda$  for each OM was varied around  $\lambda=1.0$ . To prevent undesired fuelling changes caused by a  $\lambda$ bias control, bias control was turned off. The gasoline used for these experiments was market fuel with research octane number 95.5, a density of 736 kg m<sup>-3</sup>, a sulphur content of 22.6 mg kg<sup>-1</sup>, and an H/C ratio of 1.97. One of the purposes of this experiment was to study the interference of H<sub>2</sub> in the exhaust on the lambda probe. For this reason all operating modes were repeated using CH<sub>4</sub> as the engine fuel (commercial bottled CH<sub>4</sub> based on fossil-fuel natural gas). However, the focus of the H<sub>2</sub> stable isotope study was on the exhaust of the gasoline-powered engine, and we therefore discuss the CH<sub>4</sub> results only briefly in context of the H<sub>2</sub> isotopes. In addition to the chemical species discussed here, measurements in the exhaust were made of O2, nitrogen oxides (NO<sub>x</sub>, NO, NO<sub>2</sub>), CH<sub>4</sub>, short-chained hydrocarbons ( $C_2-C_4$ ), some aromatics, total hydrocarbons (HC), water (H<sub>2</sub>O), N<sub>2</sub>O, and sulphur dioxide (SO<sub>2</sub>).

Samples for D/H isotope analysis were collected at  $\sim$ 2 bar in 1-L glass flasks fitted with 2 stopcocks (NORMAG, Illmenau, Germany), and others at  $\sim$ 3 bar in  $\sim$ 2-L internally electro-polished stainless steel (ss) canisters using a small membrane pump (KNF model N86-KTE). The sampling stream was cooled in an ice bath before entering the pump and the condensed water was periodically removed. The samples were further dried using magnesium perchlorate (Mg(ClO<sub>4</sub>)<sub>2</sub>) before collection in the flasks.

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Analysis of the samples was performed within 3 months after their collection.

#### 2.2 Instrumentation

 $H_2$  concentrations were measured online using a commercially available electron impact ionization mass spectrometer (EIMS, H-Sense, V&F Analyse und Messtechnik, Absam, Austria). The instrument precision is ~1% (1 $\sigma$  of 1-min mean). However the measurements were affected by a blank of ~8 ppm (parts-per-million,  $10^{-6}$ , molar) likely caused by some  $H_2O$  interference in the system. This blank was determined and corrected for by comparing the EIMS results with the concentration results from the flask samples, which were available for the low-concentration range ( $\lambda$  >1). While this blank correction is of little relevance for the high-concentration samples, it resulted in considerable uncertainties of up to 50% for some of the low-concentration samples. Calibration of the instrument was achieved using commercial standards (30 ppm and 1000 ppm, Messer Schweiz, accuracies 0.5%) and during an independent validation study, the instrument was found to exhibit linear response within the range of our measured concentrations (Brühlmann, 2004). Overall accuracies of the  $H_2$  concentrations are estimated at 5 ppm or 3%, whichever is greater.

CO, CO<sub>2</sub>, and H<sub>2</sub>O were measured online using an automotive Fourier Transformation Infra-red (FTIR) instrument (AVL SESAM). This instrument was calibrated during commissioning using high-precision gases and did not need re-calibration during the experiment. Estimated uncertainties are 0.14% for H<sub>2</sub>O, 0.12% for CO and 0.33% for CO<sub>2</sub>. The CO and CO<sub>2</sub> concentrations were converted to dry air mole fractions using this instrument's H<sub>2</sub>O results. Earlier-mentioned compounds, which are not discussed here, were also measured on this FTIR and on a standard engine emission bench instrument (Horiba Mexa 9200).

Isotopic measurements of  $H_2$  from the flask samples were conducted in the isotope laboratory of the Institute for Marine and Atmospheric research Utrecht (IMAU), Utrecht University using isotope ratio mass spectrometry (Rhee et al., 2004). Isotope results are reported using the "delta" notation,  $\delta D = [(D/H)_s/(D/H)_{VSMOW} - 1] \times 1000\%$ , where s

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is the  $H_2$  of the sample and VSMOW is Vienna standard mean ocean water used as reference (Craig, 1961; Gonfiantini, 1978). The mean of the 1- $\sigma$  precisions from repeated analysis and duplicate samples is 11‰, which is poorer than for ambient air measurements and caused by the high concentrations employed in this study.

#### 2.3 Results and discussion

The results for our compounds of interest are listed in Table 2 and shown in Fig. 1 as a function of the air-fuel ratio. The air-fuel ratio is expressed here as a lambda  $(\lambda)$  value, whereby  $\lambda < 1$  is the range of fuel-rich combustion (typically during vehicle accelerations) and  $\lambda > 1$  corresponds to fuel-lean combustion (excess oxygen present). H<sub>2</sub> and CO are produced during incomplete combustion, and their abundance is related, amongst other chemical reactions, through the water-gas shift reaction H<sub>2</sub>+CO<sub>2</sub>  $\leftrightarrow$  H<sub>2</sub>O+CO. Pre-TWC results showed largest emissions under fuel-rich conditions, with concentrations up to  $\sim$ 14 000 ppm for H<sub>2</sub> and  $\sim$ 41 000 ppm for CO. These concentrations decreased with increasing  $\lambda$  to about 1000–1500 ppm for H<sub>2</sub> and 4000–7000 ppm for CO for the leanest samples. From the presence of large H<sub>2</sub> concentrations in all pre-TWC samples, we conclude that the H<sub>2</sub> concentration and isotope ratio in the intake air (which we measured in a nearby ambient sample with a concentration of 573 ppb and a  $\delta$ D of 83.3‰) is not relevant in the following discussion.

Post-TWC emissions of  $H_2$  and CO were lower compared to the pre-TWC emissions. Here the emissions were also larger in the fuel rich range, but nearly disappeared in the fuel-lean range. For  $H_2$ , most concentrations of the fuel-lean experiments were below typical ambient atmospheric values of ~0.5 ppm, implying that under these conditions,  $H_2$  is destroyed on the TWC. The post-TWC  $H_2$  concentrations in the fuel-rich range were highly dependent on the operating mode, with highest emissions for OM-4. In contrast, the post-TWC concentrations for CO showed remarkably similar  $\lambda$  dependence for all 4 operating modes.

The cooling of the TWC also had significant effects on post-TWC  $\rm H_2$  emissions. The reduction of the TWC temperatures in OM-1 by ~50 °C compared to OM-2 (Table 1)

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resulted in a reduction of H<sub>2</sub> emissions to less than half of those in the fuel-rich range, but had no effect on the CO emissions. These observations demonstrate that reactions other than the water-gas-shift play a role, because the latter alone would be driving the equilibrium to higher H<sub>2</sub> yield at reduced temperatures (Haryanto et al., 2009), and not the opposite we've observed. A possible, but likely not only explanation for these observations, is the TWC cocking with hydrocarbons under these conditions, thereby preventing adsorption of H<sub>2</sub>O required for the water-gas-shift reaction (e.g., Auckenthaler, 2005).

Our D/H isotope analysis shows largely differing isotope ratios between the individual settings, with the following three major observations. Firstly, for a given air-fuel ratio, the  $H_2$  in the post-TWC exhaust was more depleted in deuterium compared to the pre-TWC exhaust. For the range where pre- and post-TWC samples were available, the pre-TWC  $\delta$ D values ranged from -140% to -195% but these values decreased to between -270% and -370% after passing through the TWC. Secondly, the post-TWC D/H ratios showed a strong dependence on  $\lambda$  with lower  $\delta$ D in the fuel-rich range (lowest  $\delta$ D values are -370%) and higher  $\delta$ D in the fuel-lean range (up to  $\delta$ D=-160%). Thirdly, we observed systematic differences between the operating modes, with OM-1 showing the lowest  $\delta$ D values and OM-4 exhibiting the highest  $\delta$ D values.

## 2.4 H<sub>2</sub> Isotope equilibration with H<sub>2</sub>O

The finding of lower isotope  $\delta D$  values after vs. before the TWC is opposite to what would be expected if  $H_2$  was removed through a normal chemical (oxidation) process where the light isotopologues are removed preferentially. We hypothesize that our observed isotopic signatures are controlled by a temperature-dependent isotope equilibration process between  $H_2$  and  $H_2O$ . For such an equilibration process, the hotter engine exhaust ( $\sim 800~^{\circ}C-1000~^{\circ}C$ ) results in a higher isotope ratio compared to the relatively cooler exhaust downstream of the TWC ( $\sim 500~^{\circ}C-700~^{\circ}C$ ). A similar temperature-dependent isotope equilibration process was recently suggested by Affek and Eiler (2006) for the oxygen isotopes in  $CO_2$  and  $H_2O$  in car exhaust.

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To explore the above hypothesis we first discuss the post-TWC samples where the different TWC temperatures of the 4 operating modes are expected to show this equilibration effect. In Fig. 2 we compare the isotope ratios of the post-TWC fuel-rich samples with results from the theoretically derived H<sub>2</sub> – H<sub>2</sub>O isotope equilibrium as studied by Bottinga (1969). We plot the isotope results as a function of the exhaust temperature ranges as measured before and after the TWC. The comparison of our results to those of Bottinga (1969) are hampered by the fact that our isotope results are reported vs. the VSMOW reference where as the  $\delta$  notation in Bottinga (1969) denotes the H<sub>2</sub> isotope ratio relative to that of H<sub>2</sub>O with which it is in equilibrium. Although offset by ~100%, our results show a temperature-dependence that is very similar to that derived from theory (Fig. 2). To bring the calculated equilibrium values and our results to the same scale, information on the  $\delta D$  of the exhaust H<sub>2</sub>O is necessary. This can in principle be obtained by isotope analysis of the exhaust H<sub>2</sub>O, but unfortunately H<sub>2</sub>O was not collected. Alternatively, assuming that H<sub>2</sub>O is the dominant hydrogen pool in exhaust, its  $\delta D$  could be approximated by that of the gasoline, which is estimated at -80% to -110% vs. VSMOW (Schimmelmann et al., 2006). We therefore convert the theoretical results (Bottinga, 1969) to the VSMOW scale by shifting them by these offsets (Fig. 2). Our measured  $\delta D$  values of the fuel-rich post-TWC samples then agree well with the prediction at the corresponding temperatures. The  $\delta D$  of the sample collected when CH<sub>4</sub> was used as fuel, is offset by ~80% from the range for gasoline. This finding is consistent with our hypothesis because CH<sub>4</sub> in natural gas is even more depleted in D/H than gasoline, e.g.  $\delta D$  equals -190% to -220% vs. VSMOW for natural gas of Russian origin (Cramer et al., 1999), which accounts for a significant fraction of the European pipeline natural gas mixture from which the CH<sub>4</sub> in our experiment was extracted.

Our finding of an equilibration temperature similar to the temperature of the TWC is somewhat surprising. Exhaust flow is fast and so are its temperature changes. The  $H_2-H_2O$  isotopic equilibration rates at these high temperatures are assumed to be fast also, but likely to slow down as the exhaust cools. As a consequence it is to be ex-

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pected that the  $\delta D$  in H<sub>2</sub>, which is also changing rapidly in the cooling exhaust stream, leads to a final mean  $\delta D$ , which is derived from a composite (frequency distribution) of isotope ratios from a wider equilibration temperature range, rather than a  $\delta D$  from a single equilibration temperature. The width of this temperature distribution remains however unclear, but is likely related to the rate of exhaust cooling and to geometry and surface area effects in the exhaust stream. Rather than based on a broad temperature range, the mean  $\delta D$  can also be interpreted as a product of an equilibration at a single 'lock-in' temperature (of a very narrow-width temperature distribution). Although inconclusively, Affek and Eiler (2006) find evidence in their study that the oxygen isotope ratio in CO<sub>2</sub>–H<sub>2</sub>O corresponds to such a single lock-in temperature of ~200 °C, rather than a mixture of isotope ratios from a broad temperature spectrum (isotopic mean from e.g. 900 °C and 50 °C in their example). In our case of H<sub>2</sub>–H<sub>2</sub>O, a lock-in temperature of 200 °C would result in a theoretical  $H_2 - \delta D$  of approximately -600%, a value that is inconsistent with our laboratory observations or any field observations of combustion/industrial isotope signatures found in the past (Begemann and Friedman, 1959; Ehhalt et al., 1963; Gerst and Quay, 2001; Gonsior and Friedman, 1962; Rahn et al., 2002). As shown above, the lock-in temperatures for the post-TWC fuel-rich samples in our experiment are the TWC temperatures (~500 °C-700 °C). This suggests that, in the case of H<sub>2</sub>-H<sub>2</sub>O, the TWC catalytic activity facilitates a rapid isotope equilibration and that a potential isotopic re-adjustment at lower temperatures downstream of the TWC is insignificant. In contrast, the finding of Affek and Eiler (2006) suggests that for CO<sub>2</sub>-H<sub>2</sub>O, the lock-in of the isotope equilibration occurs well downstream of the TWC at much lower temperatures.

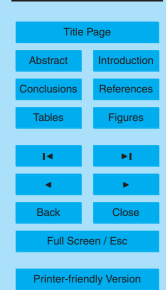
In Fig. 2, we also indicated the  $H_2$  –  $\delta D$  of our pre-TWC results by plotting these in the temperature range that is typical for exhaust gases leaving the engine's combustion chamber (800 °C–1000 °C). Qualitatively, the observation that these  $\delta D$  values are higher than those from the post-TWC samples again supports the temperature hypothesis, but quantitatively the  $\delta D$  values are more enriched for the respective temperatures than expected from theory. For the  $\delta D$  of these samples, the lock-in temperature would

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need to be much higher. The present study cannot resolve whether these signatures could be the remnants of  $H_2$ – $H_2$ O equilibration at the much higher temperatures (up to ~2300 °C) during the combustion process.

## 2.5 H<sub>2</sub> isotope fractionation during catalytic removal

The temperature-driven  $H_2$ – $H_2$ O isotope equilibration, as discussed in the previous section, can only explain some of the  $H_2$ – $\delta$ D variability we measured. For the individual operating modes, we observe a strong dependence of the isotope ratios on  $\lambda$ , particularly in the fuel-lean range (Fig. 1), even though there is only minor TWC temperature variability for each individual operating mode.  $\delta$ D of  $H_2$  increases with increasing  $\lambda$  (Fig. 1). To characterize the isotopic enrichment during  $H_2$  removal, we estimate the fractionation using the Rayleigh model where we relate the changes in  $\delta$ D to the fraction f of the post-TWC to the initial (pre-TWC)  $H_2$  concentrations (e.g. Kaiser et al., 2002):

$$ln(1 + \delta D_{res}) = ln(1 + \delta D_0) + \varepsilon \times ln(f)$$
(1)

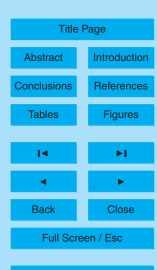
where  $\delta D_{res}$  and  $\delta D_0$  are the post (residual)- and pre-TWC  $\delta D$  values of H<sub>2</sub>, respectively, and  $\varepsilon$  is the enrichment factor. Note that a  $\delta D$  value of e.g. –300% has to be entered as –0.3 in Eq. (1). The  $\varepsilon$  is related to the fractionation factor  $\alpha$  (defined as D/H<sub>removed</sub>/D/H<sub>residual</sub>) through the definition  $\varepsilon = (\alpha - 1)$  and usually expressed in %, see (e.g. Friedman and O'Neil, 1977; Brenninkmeijer et al., 2003). The  $\varepsilon$  is calculated using least-square fitting techniques and is graphically shown in Fig. 3. We fit the data by linear regression, but in a first run we exclude the results for the very low H<sub>2</sub> concentration samples (ln(f)  $\approx$ –10) because due to their very low H<sub>2</sub> concentrations, any potential contamination with ambient air or isotope effects other than that discussed here, would have strong influences on the  $\delta D$  (the large uncertainty in these samples' H<sub>2</sub> concentrations is relatively small when plotted on a logarithmic scale). For the linear regressions of OM-2 and OM-3, we find  $R^2 \approx$ 0.6 each and large 95% confidence

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bands. The linear regressions for these operating modes and for OM-1 (dashed lines in Fig. 3) closely predict the results at the low  $\ln(f)$  end. This enhances the confidence that these samples may not have been altered significantly by any potential contamination, and we therefore include them in the linear regression (solid lines). This results in the following fractionation constants for the different operating points with 95% confidence bands: for  $\varepsilon_{\rm OM-1}$ =-39.8±3.1‰,  $\varepsilon_{\rm OM-2}$ =-15.5±3.0‰,  $\varepsilon_{\rm OM-3}$ =-18.8±5.2‰ ( $R^2$  >0.98 for all fits), and the slope of the line connecting the two OM-4 samples yields  $\varepsilon_{\rm OM-4}$ =-16.7‰. For the methane-fuelled experiment OM-4-CH<sub>4</sub>, we calculate a fractionation constant of  $\varepsilon_{\rm OM-4-CH4}$ =-20.2±69‰.

Our analysis thus suggests the presence of two separate isotope effects, a H<sub>2</sub>-H<sub>2</sub>O temperature-dependent isotope equilibration, and a kinetic isotope fractionation during H<sub>2</sub> removal. While our observations can be fully explained by these two mechanisms, it is the simultaneous appearance of both effects in our samples that remains puzzling. Since H<sub>2</sub> removal is largely bound to the TWC surface, this is the place where the kinetic isotope fractionation is believed to occur during the ~100 msec residence time in the TWC. The  $H_2-H_2O$  equilibration step must, however, also mainly occur in the TWC. This is obvious from the strong isotopic depletion when comparing post-TWC with corresponding pre-TWC isotope results. It is unlikely that the H<sub>2</sub>-H<sub>2</sub>O isotope equilibration would occur downstream of the TWC, as this would remove the clear isotopic signatures related to fractionation during removal. It is more likely that the equilibrium fractionation sets the general  $\delta D$  value at the TWC temperature, and subsequently there is a kinetic process that removes H<sub>2</sub> with the stated kinetic isotope effect. It should also be noted that the temperature of the exhaust gas increases by ~50 °C during passage through the TWC due to the exothermic reactions (the TWC is not actively heated). Given this, it is even more surprising to find the results of both effects downstream of the TWC. TWC chemistry is very complex also with regard to H<sub>2</sub>, and it is therefore also possible that some additional processes and fractionations may potentially influence our results. Hydrocarbon steam-reforming and partial oxidation, water-gas shift reaction, NO-reduction, and H<sub>2</sub> oxidation by O<sub>2</sub> are some of the

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processes involving hydrogen (D. Ferri personal communication, 2009; Salaün et al., 2009). In addition, increasing concentrations of  $H_2O$  along the TWC (with concentrations of ~15% compared to  $H_2$  concentrations of a factor ~10 less) may add to the fast and complex isotope chemistry of  $H_2$  in car exhaust.

## 5 2.6 Exhaust ratios of H<sub>2</sub>/CO, H<sub>2</sub>/CO<sub>2</sub> and CO/CO<sub>2</sub>

The ratios of H<sub>2</sub>/CO, H<sub>2</sub>/CO<sub>2</sub> and CO/CO<sub>2</sub> (in this paper reported on a molar basis) in traffic exhaust have recently received increased attention because of their potential to identify and characterize traffic pollution events in atmospheric observations, and to up-scale traffic emissions using the better constrained emission inventories of CO and CO<sub>2</sub>. Many studies over the past decades have revealed tight H<sub>2</sub> to CO relationships in raw engine exhaust (D'Alleva and Lovell, 1936; Leonard, 1961; Auckenthaler, 2005) and our pre-TWC H<sub>2</sub>/CO of 0.2-0.35 (Figs. 1 and 4) are in close agreement with these findings. Recent field studies of anthropogenically polluted air in Europe and North America have shown H<sub>2</sub>/CO ratios to vary in a relatively narrow range of ~0.3-0.6 (Novelli et al., 1999; Barnes et al., 2003; Steinbacher et al., 2007; Aalto et al., 2009), and when the traffic component was isolated from other H<sub>2</sub> and CO sources and sinks, yielded ratios of 0.48-0.51 (Vollmer et al., 2007; Hammer et al., 2009). Our post-TWC results show considerable variability (Figs. 1 and 4) for the four operating modes and the  $\lambda$  settings. In the fuel-rich range, H<sub>2</sub>/CO varied between 0.05 and 1.6 depending on the operating mode (with coolest TWC exhibiting the lowest ratios). Surprisingly the H<sub>2</sub>/CO reduced to about half for the actively-cooled TWC (OM-1 vs. OM-2) which is due to reduced H<sub>2</sub> emissions. H<sub>2</sub>/CO increases significantly near  $\lambda = 1$ . However, for  $\lambda \ge 1$ the H<sub>2</sub> and CO concentrations are tagged with high measurement uncertainties and consequently their ratios have large errors, and we are not further discussing these here. From an integrative aspect, because of their low concentrations, these ratios are also less important as they quantitatively add little to the overall traffic exhaust in a real-world case. In light of a comparison with real-world emissions, a more robust measure of  $H_2/CO$  is the cumulative  $H_2/CO$ . The overall cumulative  $H_2/CO$  over all

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gasoline post-TWC measurements is 0.25, about half of that observed in an integrative tunnel study (0.51 $\pm$ 0.11, Vollmer et al., 2007). This difference is likely caused by the somewhat arbitrary selection of  $\lambda$ -points which affects our cumulative estimate, and the generally large variability found throughout individual single-vehicle exhaust systems (Bond et al., 2010). Differences may also occur because our study uses more static settings compared to the dynamics found in real traffic. This is likely to cause some differences in the regime of  $O_2$  storage capacities and hence may influence  $H_2$  and CO levels too.

Our similarly-calculated overall ratio for  $H_2/CO_2$  is  $8.9\times10^{-3}$ , about double that found in the tunnel study  $(4.4\pm1.7\times10^{-3}, Vollmer et al., 2007)$ , and likely caused by the low-H<sub>2</sub> high-CO<sub>2</sub> diesel exhaust component in the tunnel. We also relate the H<sub>2</sub>/CO to the combustion efficiency, here defined as  $CO/CO_2$  (Fig. 4). Although the overall cumulative combustion efficiency in our experiment is significantly higher than in the tunnel study  $(36\times10^{-3} \text{ vs. } 8.4\pm2.8\times10^{-3})$ , these results demonstrate the generally more efficient combustion in vehicle engines compared to biomass burning (e.g., Röckmann et al., 2010). In conclusion, the simultaneous determination of H<sub>2</sub>, CO, and CO<sub>2</sub>, possibly along with isotopic measurements, may provide a tool to distinguish traffic exhaust from biomass burning in air pollution.

## 2.7 Implications for real-world emissions

Although here we present results from a single engine/TWC experiment, some results are likely applicable to other engine settings and to real-world driving conditions. For the investigated engine, H<sub>2</sub> removal by the TWC is large. This implies that single motor vehicle H<sub>2</sub> emissions from traffic have decreased strongly since the introduction of the catalytic converter in the 1970s. Due to increasingly stringent emission legislation around the world, improvements in catalytic converter technology, engine fueling precision, and the development of lambda control algorithms, global traffic H<sub>2</sub> and CO emissions are expected to decrease despite the increasing global car fleet (Vollmer et al., 2007).

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To place our isotope results in a real-world context, we compare these to earlier published D/H isotope measurements of air samples polluted with traffic exhaust. This comparison is shown in Fig. 5 as a "Keeling" plot, where the isotopic signature is plotted against the inverse concentration of the sample. Rahn et al. (2002) have measured traffic-polluted air samples in the Los Angeles basin in 2000, from which they derived a integrated pure anthropogenic isotopic signature (end-member) of -270%. This extrapolated value is bracketed by our pre-TWC and fuel-rich post-TWC results. Earlier results from Gerst and Quay (2001) from samples taken in 1999 in a parking garage in Seattle yielded higher concentration samples than those by Rahn et al. (2002), and were less depleted in D/H with results ranging from -173% to -183%. The insert of Fig. 5 also includes the samples with very low concentrations from the fuel-lean range. As these concentrations are sub-ambient, they cannot be quantitatively investigated in the Keeling diagram. Still, the isotope signatures are in the same range as the very high concentration samples. This suggests that in fact all of our samples are 'pure' end-member samples (not mixtures of produced H<sub>2</sub> with ambient background H<sub>2</sub>), but with largely differing source strengths.

On the one hand, all of our high-concentration post-TWC results show a  $\delta D$  equal or more negative than the end-member value predicted by Rahn et al. (2002), with the signature from the cooled TWC of OM-1 (and from the methane-powered engine) showing the largest depletion of about -370%. On the other hand, all of our pre-catalytic converter results show isotopic ratios that are much more enriched compared to the predicted end-member of -270%. These samples do, however, agree well with the results from Gerst and Quay (2001) from a parking garage in Seattle.

An interpretation of our results from a fleet-level perspective may provide some explanations for these apparent discrepancies. The isotopic signatures of exhaust at  $\lambda > 1$  are probably of little quantitative relevance because the  $H_2$  concentrations are very small. In contrast, emissions during cold starts and in the fuel-rich range ( $\lambda < 1$ ) are large, and hence their isotopic signatures are of importance. Emissions from single vehicle studies of standardized driving cycles show that cold start  $H_2$  emissions

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can contribute up to 40% of the cycles' total emissions (Bond et al., 2010). The extrapolated end-member of –270‰ by Rahn et al. (2002) based on their Los Angeles Basin study is likely a regional mixture of emissions from cold-starts (thus pre-TWC signatures) on the one hand (>–270‰) and fuel-rich TWC emissions (<–270‰) on the other. Because cold-start emissions are expected to continue contributing significantly to overall vehicle emissions, it is likely that the end-member value of –270‰ (Rahn et al., 2002) will be a useful average value for developed countries. For developing countries however, with parts of their fleets not equipped with TWC, it is likely that the high emissions from non-TWC vehicles will lead to a higher mean δD in H<sub>2</sub> traffic emissions.

It is tempting to attribute the relatively elevated  $\delta D$  found by Gerst and Quay (2001) to an apparent unrepresentative sampling location, since samples in a parking garage may be biased towards cold-start emissions (thus pre-TWC signatures). However Gerst and Quay (2001) state that their samples were taken in the morning when the majority of the vehicles arrive at the parking garage (presumably with fully-functional TWCs). Still the high emissions of a few potential cold-starts (with higher isotopic ratio) in the parking garage may have significantly influenced their results.

#### 3 Conclusions

If our results are representative of a large variety of vehicle engine/TWC systems, then our findings have several implications. Our results suggest that the isotopic signatures from vehicle emissions have 'lightened' over the last decades due to the introduction of catalytic converters in the 1970s. It is likely that this evolution has progressed in different manners in different regions of the world. Assuming that the majority of the third-world countries converted to TWC technology at a slower rate than industrialized countries, this would imply that there may have been (and still could be) an geographical difference in the isotopic signature of traffic exhaust. Our results also suggest that traffic sampling should be carried out with caution. Urban traffic with more frequent

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congested traffic used to be characterized by more fuel-rich conditions with high  $\rm H_2$  emissions hence probably lower isotope ratios. Current technological developments, however, avoid fuel-rich driving in urban traffic due to better lambda control systems, hence the fuel-rich driving (and thereby the low isotope ratios) may be limited to very high speed driving (e.g. German highways). The isotopic signature of cold-start emissions may resemble engine exhaust without TWC, hence parking garages may yield relatively enriched isotopic signatures.

Because the TWC is not actively heated, its temperature is partially a reflection of the exhaust gas temperature reaching the TWC. Therefore the driving mode will affect the TWC temperature, which in turn results in largely differing emissions of H<sub>2</sub> concentrations, their  $\delta D$ -H<sub>2</sub> and H<sub>2</sub>/CO ratio. However TWC and exhaust system geometry in combination with ambient temperatures will also significantly affect TWC temperatures and hence the emitted trace gases. Our results suggest that the exhaust  $\delta D-H_2$  for colder ambient air (e.g. winter) hence cooler TWC is significantly more negative compared to warmer air (e.g. summer). Consequently, temperature variations should also lead to seasonal and geographical variations in  $\delta D-H_2$ , with, e.g., the tropics showing different exhaust signatures compared to the high latitudes. Exhaust geometry, air flow and ambient temperatures may easily lead to temperature differences similar to the ~50 °C found for our OM-1 and OM-2 experiments, and could therefore lead to variability of  $\delta D-H_2$  of ~50%. Further studies are needed to fully confirm the hypothesis of a temperature-driven isotopic H<sub>2</sub>-H<sub>2</sub>O equilibration in motor vehicle exhaust and the potential variability caused by catalytic converters, exhaust geometry, and ambient temperature conditions. Studies on a fleet level (e.g. tunnel studies) in various seasons and parts of the world could greatly help to better characterize the H<sub>2</sub> traffic emissions and their isotopic signatures.

Except for the study by Affek and Eiler (2006), other isotope studies of car exhaust do not specifically address the question whether a significant thermal equilibration process between  $\rm H_2O$  and other exhaust compounds may occur. Combined pre- and post-TWC measurements were generally not conducted except for the  $\rm N_2O$  study by

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Toyoda et al. (2008). In their study, both isotopic enrichment and depletion were observed across the TWC, depending on the settings of the experimental variables, but in their case, the situation is complicated by both destruction and production of N<sub>2</sub>O in the exhaust stream. For CH<sub>4</sub> and CO stable isotope studies on cars, including one without catalytic converter (Nakagawa et al., 2005; Tsunogai et al., 2003), there is no clear evidence of thermal hydrogen or oxygen equilibration with H<sub>2</sub>O, possibly because of the overwhelming kinetic isotope fractionation during catalytic removal of CH<sub>4</sub> and CO. However, in the exhaust of the car without catalytic converter, (Tsunogai et al., 2003) find <sup>18</sup>O-depletion (but no <sup>13</sup>C depletion) in CO when the engine is still relatively cold compared to when it is hot. It remains unclear whether this could also be a thermal equilibration effect between CO and H<sub>2</sub>O, or, as the authors hypothesize, the cold-start is characterized by an O<sub>2</sub>-depleted combustion (hence the large CO emissions) possibly resulting in an enhanced kinetic isotope effect during the atmospheric oxygen destruction stage thereby producing <sup>18</sup>O depleted CO. It is possible that the biomass burning <sup>18</sup>O depletion in CO of colder smoldering combustion stages compared to hotter flaming stages (Tsunogai et al., 2003) could be partially caused by thermal equilibration processes with H<sub>2</sub>O or O<sub>2</sub>. Similarly, for biomass burning (Röckmann et al., 2010), the isotopic depletion in D/H with increasingly poor combustion efficiency ( $\Delta CO/\Delta CO_2$ ) may ultimately be partially related to temperature effects because poor combustion efficiency is generally related to lack of oxygen and hence to lower-temperature combustion.

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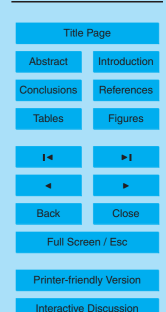
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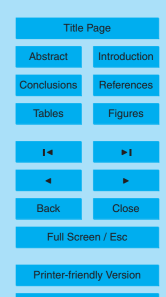


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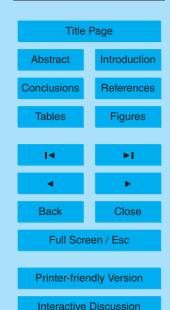
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**Table 1.** Operating modes (OM) for the experiments conducted on a passenger car engine. The three-way catalytic converter (TWC) was actively cooled in operating mode 1. OM-1 through OM-4 are gasoline-fuelled, OM-3-CH4 is methane-fuelled. Upstream and downstream TWC temperatures are lambda-dependent (low temperatures for low lambda). Although the TWC is not actively heated, some downstream temperatures are higher due to exothermic reactions occurring within the TWC.

Operating Mode (OM)	Crankshaft speed [r.p.m]	Torque [N m]	Power [kW]	downstream TWC cooling	upstream TWC temperature range [°C]	downstream TWC temperature range [ °C]
OM-1	2000	31.6	6.6	yes	460–490	456–534
OM-2	2000	31.6	6.6	no	530-556	512-586
OM-3	2000	63.2	13.2	no	598-629	566-642
OM-4	4000	31.6	13.2	no	690-721	665-753
OM-3-CH <sub>4</sub>	2000	63.2	13.2	no	566–590	558-620

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**Table 2.** Temperatures, trace gas concentrations and H<sub>2</sub> isotope ratio for pre-catalytic converter (pre-TWC) and post-TWC exhaust from a gasoline and methane-fuelled engine. The 5 different operating modes (OM) are specified in Table 1.

					Pre-TWC					Post-TWC	
Operating	Lambda	Temp.	H <sub>2</sub> conc.	$\delta$ D-H $_2$	CO conc.	CO <sub>2</sub> conc.	Temp.	H <sub>2</sub> conc	$\delta$ D-H $_2$	CO conc.	CO <sub>2</sub> conc
Mode	[]	[°C]	[10 <sup>3</sup> ppm]	[‰]	[10 <sup>3</sup> ppm]	[10 <sup>3</sup> ppm]	[℃]	[10 <sup>3</sup> ppm]	[‰]	[10 <sup>3</sup> ppm]	[10 <sup>3</sup> ppm]
1	0.892	466	13.91	-	38.86 18.93 11.63 9.60	122 139	456 483	9.37	_	37.93 17.03 8.05 5.27 4.59 3.97 3.17 2.51 1.92 1.23 0.60	124
1	0.942	481	6.52	-	18.93	139	483	0.79	-375	17.03	143
1	0.942 0.972 0.982	485 487	3.88 3.12	-	11.63	145 146	504 513	0.79 0.42 0.36		8.05 5.27	152 154
i	0.984	487	2.99	_	9.00	146	515	0.30		4 59	155
i	0.986	488	2.87	_	9.26 9.01 8.64 8.61	146	517 520 523	0.34 0.32	_	3.97	156
1	0.988	488	2.73	_	8.64	147	520	0.29	_	3.17	156
1	0.990	488	2.65		8.61	147	523	0.26	-	2.51	156
1	0.992	490	2.57	-141	8.41 8.17	147	525 528	0.22	-367	1.92	157
1	0.994	489	2.40	_	8.17	147	528	0.17	_	1.23	157
1	0.996	490 490	2.26	_	7.94	147	530	0.11	_	0.60	158 157
1	0.998 1.000	490	2.15 2.05	_	7.94 7.59 7.30	147	530 532 534	0.06 0.02	_	0.14 0.00	157
i	1.002	490	1.98	_ _ _	7.30	147	531	0.02	-162	0.00	157
i	1.012	491	1.50	-	7.13 5.71	145 146 146 146 147 147 147 147 147 147 147	531 521	0.00	-	0.00 0.00	155
2	0.892	530	13.92	_	39.18	122	512	9.58	_	37.80	123
2	0.942	545	6.50	_	18.89 11.78	139	537	2.44 2.42	-	16.48	143
2	0.972 0.982	551	3.81		11.78	145	556 563 566	2.42	-	8.06	152
5	0.982	553 554	3.03 2.88	- - - -	9.56 9.28	146	566	1.51 1.17	_	5.08 4.45 3.82 3.08 2.32 1.73 1.02	155 155
2	0.986	553	2.78	_	9.04 8.87 8.55 8.35 8.14 7.89	146	567	0.87	_	3.82	156
2	0.988	555	2.68	_	8.87	147	567 569 571	0.69	-324	3.08	156
2	0.990 0.992	553	2.54		8.55	147	571	0.56	_	2.32	156
2	0.992	555	2.44	-173	8.35	147	576	0.48	-328	1.73	157
2	0.994 0.996	553 555	2.31 2.17	-	8.14	147	576 581	0.35 0.26	_ -317	1.02 0.56	157 157
2	0.998	555	2.17	-	7.89	147	581	0.26		0.56	157
222222222222222222222222222222222222222	1.000	556	1 99	- - - -	7.67	147	580 586 580	0.14	_	0.13	157
2	1.002	556 555	1.99 1.83	_	6.90	147	580	0.02 0.00	-228	0.00	156
2	1.012	555	1.43	-	7.67 7.45 6.90 5.71	139 145 146 146 146 147 147 147 147 147 147 147 147	570	0.00	=	0.13 0.00 0.00 0.00	156
3	0.894	598	13.25	_	38.96	123 140	568	9.89	_	37.49 16.69 7.73 4.20 3.53 2.81 2.15 1.53	124
3	0.944	616	5.86	-	18.45	140	596	2.02	_	16.69	143
3	0.974 0.984	620	3.25 2.58 2.45 2.33	-	10.80	146	616	1.23	-319	7.73	153 156
3	0.984	624 623	2.58	_	8.85 8.55	148	627 627	0.96 0.88 0.78		4.20 3.53	156
3	0.988	623	2.43	_	8 24	148	629	0.00	_	2.81	156
š	0.990	625	2.18	- - - -	8.85 8.55 8.24 7.96 7.71	148	632	0.69	-296	2.15	157
3	0.992	625	2.05	_	7.71	148	632 633	0.57	-297	1.53	157
3	0.994 0.996	625	1.96	-195	7.36	149	634 638	0.46	-297	0.95	157
3	0.996	625	1.83		7.36 7.02 6.70	149	638	0.35	_	0.95 0.50 0.08 0.00	158
3	0.998 1.000	627 625	1.72 1.59	_	6.70	149	640 638	0.13 0.00	-287	0.08	157 157
3	1.000	626	1.59	_	6.16 5.95	149	638	0.00	=	0.00	157
3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3	1.002	626 626	1.44	_	5.95 5.74	146 148 148 148 148 148 149 149 149 149 150	638 635	0.00	-178	0.00 0.00	157
3	1.014	629	0.98	_	4.22	149	626	0.00	_	0.00	155
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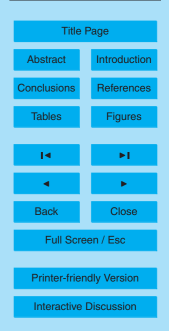




Table 2. Continued.

					Pre-TWC					Post-TWC	
Operating	Lambda	Temp.	H <sub>2</sub> conc.	$\delta D-H_2$	CO conc.	CO <sub>2</sub> conc.	Temp.	H <sub>2</sub> conc	$\delta D-H_2$	CO conc.	CO <sub>2</sub> conc
Mode	[]	[°C]	[10 <sup>3</sup> ppm]	[‰]	[10 <sup>3</sup> ppm]	[10 <sup>3</sup> ppm]	[°C]	[10 <sup>3</sup> ppm]	[‰]	[10 <sup>3</sup> ppm]	[10 <sup>3</sup> ppm]
4	0.896	690	13.93	-	41.39	120	668	14.00	-	38.46	123
4	0.946	710	6.57	_	20.73	139	701	5.32	-	17.29	143
4	0.976	715	4.04	-	13.50	143	725	3.02	-	7.98	152
4	0.986 0.988	717 719	3.25	-	11.32	145	737	2.04	_	4.31	155
4	0.988	719	3.13 2.99	-	11.08 10.67	145	739 741	1.78 1.50	_	3.61 2.88	156 156
4	0.990	717	2.99	_	10.67	145 145	741	1.50	_	2.88 2.19	157
4	0.992	719	2.70	_	9.89	146	743 745	0.92	_	1.52	157
4	0.994	719	2.62	-163	9.44	146	743 749	0.92	-273	0.91	157
4	0.998	720	2.44	-103	9.18	146	750	0.30	-273	0.43	157
7	1.000	719	2.35	_	8.97	146	753	0.04	_	0.06	157
4	1.002	720	2.18	_	8.63	146	751	0.00	_	0.00	156
4	1.004	721	2.07	_	8.43	146	748	0.00	_	0.00	156
<b>4</b>	1.006	720	1.98	_	8.28	147	745	0.00	-162	0.00	156
4	1.016	720	1.54	-	7.04	147	735	0.00		0.00	155
3-CH4	0.902	566	24.35	_	35.52	88	567	19.32	_	34.14	92
3-CH4	0.952	583	12.33	_	19.46	102	586	3.74	_	15.62	107
3-CH4 3-CH4	0.982	588 589	6.33	_	10.99	108	608 618	0.76	-377	4.11	117
3-CH4	0.992	589	4.92	_	9.20	110	618	0.21	_	0.24	119
3-CH4	0.994	590	4.64	_	8.97	111	617	0.13	_	0.06	119
3-CH4	0.996	588	4.41	-	8.62	110	620	0.09	_	0.02	119
3-CH4	0.998	589	3.96	-	8.25	111	620	0.15	-	0.06	119
3-CH4	1.000	588	3.70	_	7.79	110	615	0.00	-	0.00	118
3-CH4	1.002 1.004	588	3.50	-	7.39 7.19	110 111	609	0.00	-295	0.00	118 117
3-CH4 3-CH4	1.004	590 588	3.34 3.08	-		111	603	0.00	-	0.00 0.00	117 117
3-CH4 3-CH4	1.008	589	3.08	_	6.69 6.75	111	596	0.00	-255	0.00	116
3-CH4 3-CH4	1.008	589 586	2.80	_	6.20	110	589 590	0.00	-255	0.00	116
3-CH4	1.010	586	2.69	_	5.97	110	581	0.00	_	0.00	115
3-CH4	1.022	587	1.93	_	4.53	110	571	0.00		0.00	114

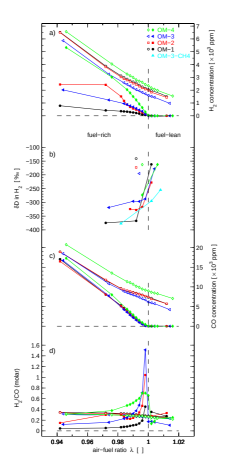
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**Fig. 1.** Engine exhaust concentrations for  $H_2$  (a) and its isotope ratio  $\delta D$  (b), CO (c), and the ratio  $H_2/CO$  (d) vs. the air-fuel ratio ( $\lambda$ ) upstream (open symbols) and downstream (filled symbols) of the three-way catalytic (TWC) converter. The colors differentiate the four main operating modes of the gasoline-fuelled engine and the isotope results for one methane-fuelled operating mode in (b). The data for the lowest  $\lambda$  are omitted to show the remaining results in closer detail. Vertical and horizontal dashed lines are visual aids at  $\lambda$ =1 and zero y-values, respectively.

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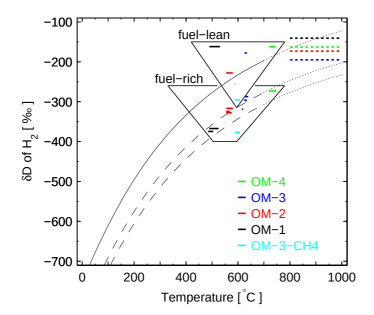
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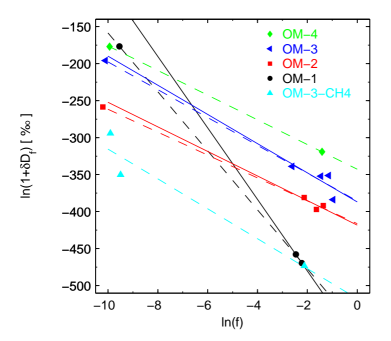
**Fig. 2.** Temperature-dependence of  $\delta D$  of H<sub>2</sub> produced in motor vehicle engine combustion. The short solid horizontal lines of variable size denote the  $\delta D$  in H<sub>2</sub> for various operating modes (see Fig. 1) on the VSMOW scale of the downstream three-way catalytic converter (post-TWC) exhaust samples. The horizontal extensions of the lines indicate the temperature ranges of the TWC. The solid curved line is a theoretical prediction (dotted lines is our extrapolation using a function  $a + b \times T + c \times T^{1/2}$ ) of  $\delta D$  in H<sub>2</sub> in isotopic equilibrium with H<sub>2</sub>O (Bottinga, 1969) and plotted relative to  $\delta D$  of H<sub>2</sub>O as a reference. This curved is shifted by the estimated  $\delta D$  of H<sub>2</sub>O in gasoline exhaust (–80‰ to –110‰ vs. VSMOW, (Schimmelmann et al., 2006), dashed curved lines) in order to match the scale of reference. The fuel-rich post-TWC gasoline  $\delta D$  agree well with the predicted values. The results for the pre-TWC samples are plotted as dashed lines in the upper right corner in the approximate temperature range of the exhaust gas when exiting the engine.

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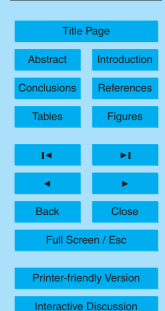




**Fig. 3.** Rayleigh fractionation diagram for the  $\delta D-H_2$  isotope fractionation of post-catalytic converter exhaust of a car engine. The solid lines are least-square linear fits through the data excluding those with low  $\ln(f)$  near -10, and the dashed lines are linear fits using all data points. The five colors denote different operating modes of the engine (see Fig. 1).

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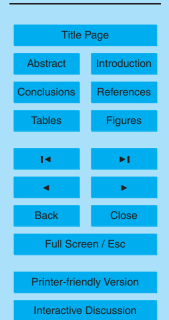
## approximate CO/CO<sub>2</sub> (molar) 0.05 0.1 0 OM-4 0.9 OM-3OM-2 0.8 OM-1 0.7 $H_2/CO$ (molar) 0.6 0.5 0.4 0.3 0.2 0.1 5 10 15 20 0 CO concentration [ $\times 10^3$ ppm]

Fig. 4. Molar  $H_2/CO$  ratio plotted vs. CO concentration. Open symbols denote pre-TWC samples and filled symbols denote post-TWC samples. The color-coding is the same as in Fig. 1.

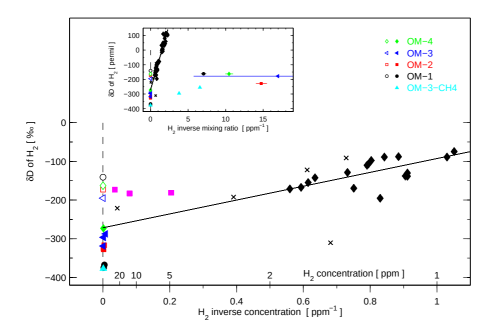
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**Fig. 5.** Isotope ratio  $\delta D-H_2$  vs. inverse concentration ("Keeling plot"). The open and filled symbols on the vertical dashed line denote pre- and post-catalytic converter samples, respectively. The color coding is the same as in Fig. 1. The black crosses (single vehicles) and diamonds (city air) are from a study in the Los Angeles Basin by Rahn et al. (2002) with a fit (solid line) through their data pointing to a -270% isotopic end-member. The magenta squares are from samples taken in a parking garage by Gerst and Quay (2001). The insert includes additionally the sub-ambient concentration samples of the post-catalytic lean engine combustion.

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