- 1 Deuterium excess in atmospheric water vapor of a
- 2 Mediterranean coastal wetland: regional versus local
- **3 signatures**

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## Abstract

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Stable isotopes of the water vapor represent a powerful tool for tracing atmospheric vapor 11 12 origin and mixing processes. Laser spectrometry recently allowed high time resolution measurements, but despite an increasing number of experimental studies, there is still a need 13 for a better understanding of the main drivers of isotopic signal variability at different time 14 scales. We present results of in situ measurements of  $\delta^{18}$ O and  $\delta D$  during 36 consecutive days 15 in summer 2011 in atmospheric vapor of a Mediterranean coastal wetland exposed to high 16 evapotranspiration (Camargue, Rhône River delta, France). A calibration protocol was tested 17 and instrument stability was analysed over the period. The mean composition of atmospheric 18 vapor during the campaign is  $\delta^{18}O=-14.66\%$  and  $\delta D=-95.4\%$ , with  $\delta_v$  data plotting clearly 19 above the local meteoric water line, and an average deuterium excess (d<sub>v</sub>) of 21.9%. At the 20 daily time step, we show a clear separation of isotopic characteristics with respect to the air 21 mass back trajectories, with the Northern air masses providing depleted compositions (8<sup>18</sup>O=-22 15.83%,  $\delta D$ =-103.5%) compared to Mediterranean air masses ( $\delta^{18}O$ =-13.13%,  $\delta D$ =-86.5%). 23 There is also a clear separation between d<sub>v</sub> corresponding to these different air mass origins, 24 25 with higher d<sub>v</sub> found for Northern air masses (23.2%) than for Mediterranean air masses (18.6%). However, since diurnal d<sub>v</sub> variations are more important than day-to-day 26 differences, an hourly time scale analysis is necessary to interpret the main drivers of dy 27 28 variability. Based on twenty-four average hourly data, we propose a depiction of typical daily 29 cycles, of water vapor isotopic composition under the two main regional meteorological 30 situations. Diurnal fluctuations are driven by 1) the influence of local evaporation, 31 culminating during day-time, and leading to an increase in absolute water vapor concentration 32 associated to a  $\delta_v$  enrichment and  $d_v$  increase; 2) vertical air mass redistribution when the PBL 33 vanishes in the evening, leading to a d<sub>v</sub> decrease, and 3) dew formation during the night, producing a  $\delta_v$  depletion with  $d_v$  remaining stable. Based on a two-component mixing model, 34 we calculate the average composition of the local vapor that produces the day-time increase in 35 d<sub>we</sub> and found higher d<sub>v</sub> for the local vapor composition under North Atlantic air mass 36 37 conditions, consistent with lower humidity conditions. In addition, we show that the stronger 38 advection rate that prevails under Northern conditions dilutes the contribution of local vapor to the ambient moisture, which indicates that regional vapor also carries a higher d<sub>v</sub> under 39 40 Northern conditions. This isotopic mass balance approach is proposed as a framework for 41 deciphering regional and local influences.

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# 1 Introduction

44 At the global scale, an acceleration of the hydrological cycle is expected in response to 45 climate change, with an intensification of both precipitation and evaporation. This effect has been recognized for the ocean (Dai et al., 2009), but less conclusive patterns appear for 46 continental surfaces, especially because of the complex behaviour of land evapotranspiration 47 (Ohmura and Wild, 2002; Roderick and Farquhar, 2002; Brutsaert, 2006; Roderick et al., 48 2007; Fu et al., 2009; Jung et al., 2010. The response of evapotranspiration to climate change 49 50 is controlled both by climatic and by hydrological parameters. Climate determines the 51 atmospheric evaporative demand (potential evapotranspiration), while hydrology controls surface water availability, thus limiting actual evapotranspiration. In addition, continental 52 53 vapor is involved in land-atmosphere feedbacks such as atmospheric water recycling: high 54 evapotranspiration may contribute to regional rainfall in contexts of tropical rainforest 55 (Brubaker et al., 1993; Eltahir and Bras, 1996; van der Ent et al., 2010) or large scale 56 irrigation (Boucher et al., 2004; Tuinenburg et al., 2012), or may even reduce potential 57 evapotranspiration by lowering temperature and increasing humidities near the earth surface (Destouni et al., 2010; Tuinenburg et al., 2012). A better understanding of the atmospheric 58 59 component of the water cycle and of the potential contribution of wetlands evaporation to the 60 regional water budget is thus important to anticipate the impact of global change, in particular in Mediterranean regions where more frequent droughts are expected. 61 62 Isotopic composition of atmospheric water vapor (subsequently called  $\delta_v$ ) represents a great

opportunity to explore land-atmosphere interactions, as the addition of moisture originating from evapotranspiration induces a modification of the isotopic content of the background vapor (Salati et al., 1979; Gat et Matsui, 1991; Gat et al., 1994; Gat, 2000; Worden et al., 2007; Yamanaka and Shimizu, 2007; Vallet-Coulomb et al., 2008; Risi et al., 2013). In addition, isotope tracers are now often incorporated into climate models (Sturm et al., 2005; Werner et al., 2011; Risi et al., 2012, 2013) in order to improve the representation of land-atmosphere interactions and feedbacks. A major challenge is thus to propose robust databases, and associated methodology for data acquisition and interpretation.

Laser spectrometers recently opened the door towards the high temporal resolution analysis of  $\delta_v$  variations, which was not possible with the traditional cold trap method. Whereas cryogenic trapping provides grab samples generally averaging a couple of hours, laser

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74 spectrometry performs continuous high frequency (up to  $\sim 1$  Hz) measurements of  $\delta_v$  (Kurita et al., 2012). This technology offers new insights into processes that affect the isotopic 75 76 composition of atmospheric water vapor and the number of studies based on continuous ground level isotope measurements over multi-week periods is continuously increasing, either 77 using Tunable Diode Laser (TDL) (Lee et al., 2006; Wen et al., 2010; Griffis et al., 2011; 78 79 Welp et al., 2012; Wen et al., 2012), Off-Axis Integrated Cavity Output Spectroscopy (OA-80 ICOS) (Sturm and Knohl, 2010; Sunmonu et al., 2012; Farlin et al., 2013; Steen-Larsen et al., 2013) or Wavelength-Scanned Cavity Ring-Down Spectroscopy (WS-CRDS) (Galewsky et 81 al., 2011; Noone et al., 2011; Tremoy et al., 2012; Steen-Larsen et al., 2013, 2014). Among 82 these experiments are some low latitude studies which have focused on the tracing of tropical 83 or sub-tropical convective activity in West Africa (Tremoy et al., 2012) or in South America 84 85 (Galewsky et al., 2011). In mid latitudes (Noone et al., 2011; Farlin et al., 2013), or high latitudes (Steen-Larsen et al., 2013), several studies have explored atmospheric mixing 86 processes at different time scales. Isotopes have also been used for partitioning 87 evapotranspiration into plant transpiration and direct evaporation, as the associated 88 89 fractionations are different (Yakir and Sternberg, 2000; Lai et al., 2006; Wang et al., 2010; 90 Griffis et al., 2011; Sun et al., 2014), for a better understanding of the role of ecosystems in 91 the hydrological cycle in the context of climate change (Wang et al., 2013). However, the number of studies remains limited because of technical difficulties associated 92 93 with field-deployed long-term measurements. Except for some work based on cold trap sampling (Jacob and Sonntag, 1991; Williams et al., 2004; Angert et al., 2008), there are to 94 95 our knowledge only two high time-resolution studies published from European sites (Iannone 96 et al., 2010; Aemisegger et al., 2014) and none at all treating Mediterranean areas. In addition, 97 there is a need for dual tracer databases, in order to use the deuterium excess signal (dexc = <u>δD-8×δ<sup>18</sup>O</u>; Dansgaard, 1964) as an additional indicator of atmospheric processes. Indeed, <u>d</u><sub>v</sub> 98 99 is often considered as a indicator of evaporation conditions, but its interpretation in 100 continental vapor remains complex since it is affected by the multiple vapor sources, combining the initial oceanic vapor and evapotranspiration from different continental water 101 102 sources (Gat et Matsui 1991; Vallet-Coulomb et al., 2008; Lai and Ehleringer, 2011; Welp et 103 al., 2012; Jouzel et al., 2013; Aemisseger et al, 2014).

In this paper, we provide results obtained in Camargue on 36 consecutive days during

summer 2011, using WS-CRDS technology. The Camargue region is emblematic of

Mediterranean wetlands, with important water inflow requirements and strong evaporation

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losses, making them highly sensitive to climate change and human pressures on water resource. Our experimental site is located in the center of Camargue, close to the main lagoon (Vaccarès lagoon,  $65 \text{ km}^2$ ). Considering the location of the site, and the important availability of open water in the surroundings, we expect a dominant contribution of evaporation upon transpiration fluxes. This study will provide the opportunity to investigate how local evaporation combines and interact with the other regional vapor sources. We will focus on identifying the main drivers of deuterium excess variability at different time scales, and explore the relevance of using either relative humidity, as an indicator of evaporation conditions, or specific humidity (mixing ratio) as a proxy of mixing between different vapor sources. Before analysing our results, we present technical aspects of measurement calibration and validation. We then analyse the day-to-day variations of  $\delta_v$  and  $d_v$  in relation to climatic parameters and the air mass back trajectories. Finally, average hourly variations are explored and we interpret the typical daily cycles according to the main regional meteorological conditions in order to depict their driving factors and the influence of local processes.

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# 2 Data acquisition

# 2.1 Protocols

Continuous in situ measurements of the isotopic composition of atmospheric water vapor ( $\delta^{18}O$  and  $\delta D$ ) were performed during summer 2011 between 20 July and 24 August, at 1.75m height and approximately 170m from the East border of the Vaccarès lagoon (Figure 1), using Wavelength-Scanned Cavity Ring-Down Spectroscopy (WS-CRDS). The instrument we use is the Picarro L1102-i isotopic liquid water and water vapor analyser (Picarro Inc., Sunnyvale, California, USA), which measures the isotopic composition of atmospheric water vapor every 5 to 7 seconds. Installed in an air-conditioned room, the analyser is connected to an outside air intake. As water vapor may be sticky on the walls of any tubing, a bypass configuration is used to bring the air in quickly, at a rate of 6L.min<sup>-1</sup> (with a Laboport vacuum pump N86 KN.18) through PVC tubing, in order to minimise wall effects that lead to fractionation in the inlet. The analyser then subsamples this air at a rate lower than 0.04L.min<sup>-1</sup>.

Calibration of laser measurements is performed according to <u>liquid</u> laboratory standards, and a three -way valve allows switching between the introduction of ambient air and vapor from the liquid sample vaporizer (vaporization module V1102-i adjusted to 110°C for flash vaporization to avoid fractionation). We used three laboratory standards whose isotopic spans

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a range of values including the composition expected in the atmosphere of the Camargue 139 140 (Table 1). 1.8µL of water standards are introduced is injected into the vaporizer using an 141 autosampler (CTC Analytics LEAP Technologies HTC PAL autosampler) with a SGE 5µL 142 syringe. We used synthetic air as the dry carrier gas (water content <45ppmv) delivered at a 143 pressure of 2.5±0.5psi. The analyser takes about nine minutes to perform approximately 40 144 measurements per injection, and these results are then averaged by the software. Between 145 each injection, the syringe is cleaned with 1-methyl-2-pyrrolidinone (NMP) solvent wash. 146 Our laboratory successfully participated in the IAEA 2011 proficiency test on routine analysis of  $\delta^{18}$ O and  $\delta$ D in liquid water (Wassenaar et al., 2012). 147 Previous studies have shown that optical spectrometric methods can induce a dependence of 148 149 isotopic measurements on water vapor concentration (Gupta et al., 2009; Schmidt et al., 2010; 150 Johnson et al., 2011; Tremoy et al., 2011). This does not affect liquid sample measurements, 151 since the water quantity introduced into the analyser through the vaporizer is nearly constant, 152 but important variations of vapor content do occur when analysing ambient atmosphere, 153 making it necessary to correct optical measurements. We have evaluated the concentration 154 dependence of our instrument for a large range of water concentrations, and for different 155 isotopic compositions using our three standards (Figure 2). In addition, for routine analysis, 156 an evaluation of the water concentration effect is performed at approximately 24-hour 157 intervals with the more depleted standard (Standard 1) whose isotopic composition is close to 158 the atmosphere of the Camargue. The autosampler is set to perform six injections of 0.9µL, 159 1.6μL and 2.0μL in the vaporization module to obtain three water vapor concentrations 160 ranging from about 8000 to 28000ppmy, comprising values expected in the study area. In 161 addition, calibration for isotopic composition is performed at least every 24 hours with our 162 three standards. For each standard, the autosampler is set to perform six injections of 1.8µL in 163 the vaporization module, corresponding to a water vapor concentration of approximately 20 164 000ppmv. The first two injections are disregarded to remove the memory effect, and the last four injections are averaged to obtain the  $\delta^{18}O$  and  $\delta D$  measurements of each standard. 165 166 Gaps in time series correspond to calibration periods, but may also reflect accidental power 167 cuts or data eliminated during episodes of condensation in the tubing. This latter problem was

detected thanks to the use of transparent air tubing, and was easily recognizable a posteriori

through the abnormally smooth shape of the water vapor concentration curve. A heating cable

was then used to avoid any condensation such as may occur when a negative temperature

gradient exists between the sampling point and the inlet of the analyser.

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# 2.2 Dependence on water vapor concentration

- 174 It has been shown that the dependence of WS-CRDS isotopic measurements on water vapor 175 concentration is instrument-dependent (Tremoy et al., 2011). The concentration dependence of our instrument, evaluated for a large range of concentrations (Figure 2), showed a linear 176 177 response in the range of water vapor concentration measured in the atmosphere of the Camargue (from 9000ppmv to 28000ppmv), with the relationship deviating slightly from the 178 linear trend at very low and high water vapor concentrations. For  $\delta^{18}O$  of standards 2 and 3 179 180 there is no concentration effect, but a small dependence did appear for the more depleted standard (-0.33%.10000ppmv<sup>-1</sup>; R<sup>2</sup>=0.75). For δD, we found significant regression 181 coefficients (R<sup>2</sup>=0.98, 0.99 and 1.00 for standards 1, 2 and 3 respectively), and the slope of 182 183 the dependence was only slightly different between standards (between 7.8 and 184 8.1‰.10000ppmv<sup>-1</sup>).
- 185 This justifies the use of only one standard (Standard 1) to perform the very time-consuming
- daily calibration of the water concentration effect (160 minutes for a 3-point calibration).
- 187 Measurements are corrected for concentration dependence before applying the isotopic
- 188 calibration in order to drive back atmospheric measurements to the water vapor concentration
- 189 w<sub>reference</sub> at which standards are measured, as follows (Schmidt et al., 2010):

- where  $\delta_{reference}$  (‰) is the reference  $\delta$  value at  $w_{reference}$  (20000 ppmv in our case);  $\delta_{observed}$  (‰)
- is the observed  $\delta$  value at  $w_{observed}$  (ppmv) the observed concentration; m (%.ppmv<sup>-1</sup>) is the
- slope of the concentration dependence (Table 2).

# 2.3 Instrument stability

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To evaluate the long-term stability of our instrument, statistics were calculated on raw delta values measured on our three standards during the entire measurement campaign (Table 1). Results show that the long-term variability is very low, but a regular calibration is nevertheless performed. Variability of both calibrations (for water vapor concentration and isotopic composition) is shown in Table 2. For  $\delta^{18}$ O, the calibration for concentration appears highly variable but often flat (no concentration effect), resulting in a linearity which is not always significant  $(0.26 \le R^2 \le 1.00)$ . Thus,  $\delta^{18}$ O was corrected only when the concentration

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dependence was significant. For  $\delta D$ , the regression is slightly variable in the long-term, and its linearity is always significant (0.89 $\leq$ R<sup>2</sup> $\leq$ 0.99). For daily calibration in isotopic composition (normalisation to the VSMOW2-SLAP2 scale), we used a three-point calibration, which allows to check for the linearity of the calibration, Linear regressions between raw measured values and absolute standard compositions are established as follows for  $\delta^{18}O$  and  $\delta D$ :

 $\delta_{calibrated} = slope \cdot \delta_{measured} + intercept$  (2)

Regressions are stable in the long-term (Table 2), and their linearity is always significant (R<sup>2</sup>=1.00). Accuracy and precision of our <u>instrument</u>, estimated by considering standard 2 as a sample which has been calibrated with standards 1 and 3, show good results (Table 3). <u>These performances are estimated on liquid measurements</u>, and are probably lower for vapor <u>measurements</u>.

3 Local atmospheric data,

Hourly air temperature and relative humidity were measured at the study site. Hourly wind speed and daily PET (Potential Evapotranspiration calculated from the Penman-Monteith

(www.meteothinker.com).

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speed and daily <u>PET (Potential Evapotranspiration calculated from the Penman-Monteith formula, Monteith, 1965)</u>, were obtained from a meteorological station located 400m far from the site of isotopic measurements. In addition, surface temperature was also measured in the <u>main pond of the Vaccarès lagoon\_system</u> at hourly time-step. <u>Backward three-dimensional trajectories were computed for each individual day with the Internet-based NOAA HYSPLIT Trajectory Model (<a href="http://ready.arl.noaa.gov/HYSPLIT.php">http://ready.arl.noaa.gov/HYSPLIT.php</a>), using GDAS (Global Data Assimilation System) meteorological data. <u>The HYSPLIT (HYbrid Single-Particle Lagrangian Integrated Trajectory) model is a complete system which allows computing simple air parcel trajectories.</u> Trajectories were parameterized to end up at the station location at 1200 UTC, at three heights of 50, 100 and 500 meters above ground level, with a total back run time long enough to get <u>back</u> to the oceanic source (maximum of 144 hours). Planetary boundary layer heights were obtained from GDAS meteorological data (3-hour time step). They were extracted and interpolated to our specific station using the MeteoInfo software</u>

In addition, we collected and analysed three rainfall samples corresponding to four small rainfall events (25, 26, 27 July, 7 August 2011). These samples account for a total of 7.6mm of the 9.6mm of cumulated rainfall during the measurement campaign; the remaining 2mm correspond to eight very small events between 14 and 23 August (not sampled).

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# 4 Results and discussion

The mean composition of atmospheric water vapor during the campaign is  $\delta^{18}O$ =-14.66‰,  $\delta D$ =-95.4‰ and  $d_v$ =21.9‰, with significant daily variations, from  $\delta^{18}O$ =-19.22 to -9.96‰ and  $\delta D$ =-125.2 to -61.7‰, while  $d_v$  varies between 7.8 and 31.2‰ (Figure 3). These deuterium excess values are higher than those of our three rainfall samples: 5.6‰, 4.5‰ and 7.0‰, (for  $\delta^{18}O$ =-3.89‰, -3.7‰, and -0.66‰ respectively), and also higher than the long-term average composition of regional precipitation in the neighbouring GNIP Station (Avignon, ≈50km far from the experimental site; d-exc = 9.2 ‰). The composition of vapor in equilibrium with rainfall is plotted for comparison (Figure 3). Values slightly enriched in  $\frac{18}{16}O$  compared to measured  $\delta_v$  suggest a probable evaporation of rainfall in the atmosphere, a classical feature of small summer rainfall events (Celle-Jeanton et al., 2001), while values slightly depleted in D compared to measured  $\delta_v$  points to the influence of an enriched vapor at ground level.

In measured atmospheric vapor, substantial day-to-day variations of  $\delta_v$  are observed, while  $d_v$  presents high hourly variations (Figure 3). In order to explore the relevant time scales for interpreting isotopic data in terms of synoptic and mesoscale meteorology versus local influences, we first analyse the relations of our data to daily climatic variables and air mass origins. Then, we focus on average diurnal variations for identifying the local influences.

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# 4.1 Correlations between isotope data and local climatic parameters

Quite good correlations are found between daily values of  $\delta_v$  and <u>local</u> climatic parameters (Table 4), the best fit being with the mixing ratio q, (R<sup>2</sup>=0.72 and 0.62 for  $\delta^{18}O$  and  $\delta D$  respectively). Such a correlation could result from the case of simple Rayleigh rainout processes, where condensation phases progressively remove water from the atmosphere and deplete the remaining atmospheric vapor in heavy isotopes (Dansgaard, 1964; Lee et al., 2006; Wen et al., 2010). The progressive rainout follows a temperature decrease, and should also result in a correlation between  $\delta_v$  and T, and subsequently between rainfall isotope content ( $\delta_p$ ) and T, which has led to the establishment of the isotopic thermometer (Dansgaard, 1964; Jouzel et al., 1997). Our data show a relation between  $\delta_v$  and T (Table 4, Figure 3) close to the isotopic thermometer:  $\Delta \delta^{18}O/\Delta T$ =0.53‰.°C<sup>-1</sup> (Jouzel et al., 1997), but the correlation is relatively poor (R<sup>2</sup>=0.30), indicating more complex processes, as was

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observed during a long-term German survey in which  $\delta_v$ -T correlation was degraded during summer because of the admixture of vapor originating from plant evapotranspiration (Jacob and Sonntag, 1991). Since the correlation between  $\delta_v$  and q is much stronger than the  $\delta_v$ -T correlation, it indicates a mixing between air masses characterized by different vapor concentrations as could result from different marine origins, and/or from the addition of continental vapor into the atmosphere.

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With respect to deuterium excess, we found that q is also the best predictor of d<sub>v</sub> variations at a daily time-step ( $R_v^2$ =0.54, Table 4). In addition, we observed a correlation between  $d_v$  and  $RH_a$  ( $R^2$ =0.51), consistent with Welp et al. (2012), but the correlation becomes very low (R<sup>2</sup>=0.19) when using relative humidity at surface temperature (RH<sub>8</sub>). The significance of deuterium excess in terms of a proxy for the conditions at the vapor sources comes from the fact that it is determined by the isotopic fractionation that occurs during evaporation, and more precisely by its variable kinetic component, which mainly depends on relative humidity (Craig and Gordon, 1965; Gonfiantini, 1986; Pfahl et Wernli, 2008; Uemura et al., 2008). However, relative humidity is only relevant for characterizing evaporation conditions if reported at surface temperature. In this case it represents the vapor concentration gradient between water and air, which controls the kinetic fractionation. Starting from a value around d<sub>v</sub>=0‰ in ocean water, the kinetic fractionation associated to initial seawater evaporation produces an increase in d<sub>v</sub> in oceanic vapor inversely related to humidity conditions (Merlivat and Jouzel, 1979; Armengaud et al., 1998; Pfahl et Wernli 2008; Uemura et al 2008; Jouzel et al., 2013; Steen-Larsen et al., 2013), During the air mass trajectory over land, dv may be further increased by the addition of vapor of continental origin (Gat and Matsui, 1991, Gat et al., 1994; Angert et al., 2008; Lai and Ehleringer, 2011; Welp et al., 2012). This "secondary" vapor is also expected to carry a  $d_v$  inversely related to humidity conditions at the vapor source (Aemisseger et al., 2014). However, the lack of correlation between d<sub>v</sub> and RH<sub>s</sub> in our data, whilst the dy-q correlation is stronger, suggests that mixing processes between air masses characterized by different vapor origins and concentrations have weakened the dv and RH<sub>s</sub> relation, as was the case for the  $\delta_v$ -q correlation.

Since it is hardly modified by equilibrium condensation,  $d_y$  can be considered as a conservative tracer of vapor sources during mixing processes. Nevertheless, the conservative behaviour of  $d_y$  during rainout is perturbed by the non-linearity in the definition of deuterium excess: the delta-notation approximation induces a small shift in  $d_y$  when there is a large

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decrease in  $\delta^{18}$ O and  $\delta$ D (Gat et al., 1996; Angert et al., 2008; Welp et al., 2012). In the observed range of  $\delta^{18}$ O (from -11 to -18‰), we have estimated this effect to induce a 2.5‰ increase in d<sub>v</sub>, while the substantial variations in d<sub>v</sub> observed in our daily data are significantly higher (from 15 to 26%), allowing the use of d<sub>v</sub> as a tracer of different vapor pools.

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# 4.2 Regional isotopic signatures.

In order to examine the link between the isotopic characteristics of vapor and the air mass trajectories since its oceanic origin, data were classified, on a daily basis, following the three main regional features schematically represented in Fig. 1. Corresponding hourly data are scattered in the  $\delta^{18}$ O- $\delta$ D plot distinctly above the local meteoric water line (LMWL, Figure 4), and show isotopically depleted compositions for air masses coming from the North Atlantic, while air masses coming from the Mediterranean Sea display isotopically enriched vapor (Table 5, Figure 4). For air masses coming from the Bay of Biscay, hourly data encompass the entire isotopic range, Local climatic data associated with the three meteorological situations reflect synoptic weather conditions related to origin of air masses, Air masses coming from the North Atlantic are associated with cold, dry and strong winds, while those from the Mediterranean are associated on the contrary with warm, wet and light winds (Table 5). The Bay of Biscay is an intermediate situation. It thus appears that the greater the distance over land (Figure 1), the more depleted is  $\delta_v$ . The northern trajectory corresponds to a "Mistral" situation: a typical cold, dry and strong north-northwest wind that affects the north of the occidental Mediterranean basin 130 days a year on average. The air mass is gradually dried out by rainout processes over land and accelerated in the Rhône River Valley and acquires a depleted isotopic signature. On the contrary, an air mass travelling over the Mediterranean Sea with the slight thermal wind or sea breeze coming from the South is moistened and acquires and maintains an enriched isotopic signature.

With respect to d<sub>v</sub>, North Atlantic air masses bring higher values (+23.2%) than Mediterranean air masses (+18.6%), but the main feature which appears on the  $\delta^{18}$ O- $\delta$ D plot (Figure 4) is the distinct separation between nocturnal and diurnal hourly data, with the latter plotting higher above the meteoric lines. This feature is observed for each of the three air mass origins.

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For the three classes of data, strong linear fits are evidenced (Figure 4), with slopes lower than eight, a value attributed to thermodynamic equilibrium, as it approximately represents the

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average ratio between D and <sup>18</sup>O liquid-vapor <u>equilibrium</u> fractionation. Such low slopes have also been observed in atmospheric vapor from Hawaii (7.02 (Bailey et al., 2013)), Greenland (between 6.47 and 7.44 (Steen-Larsen et al., 2013)), Western Siberia (between 5.6 and 7.7 (Bastrikov et al., 2014)), and in North America (7.5 (Berkelhammer et al., 2013)). The previously described shift in d<sub>v</sub> during Rayleigh-type rainout, due to the non-linearity in the deuterium excess definition, is not sufficient for explaining these low slopes, and non-equilibrium processes such as evaporation are also probably involved (Gibson et al., 2008).

# 4.3 Analysis of average daily cycles

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In order to observe the diurnal variations suggested by the night-day separation, 24-hour average values of isotopic and climatic data are plotted for the three air mass origins (Figure 5). A well-pronounced cyclicity appears for  $d_v$  in each meteorological condition. For Bay of Biscay origins, the average  $d_v$  cycle is very similar to the North Atlantic conditions, with larger standard deviations, reflecting a more variable climatic situation. For Mediterranean conditions, the  $d_v$  cycle is higher in amplitude than for North Atlantic conditions, and lower in absolute values, especially during the night. The high standard deviations observed during the last hours of the day correspond to calibration periods for which there are fewer data, but may also be due to variations in the timing of the PBL evening transition. The increase in  $d_v$  during the morning is mainly associated with a  $\delta D$  increase, while  $\delta^{18}O$  variations are smoother. Amplitudes of daily isotopic variations are higher for Mediterranean conditions (9.7%, 16.2% and 1.3 % for  $d_v$ ,  $\delta D$  and  $\delta^{18}O$  respectively) than for North Atlantic ones (7.0%, 13.1% and 1.1% for  $d_v$ ,  $\delta D$  and  $\delta^{18}O$  respectively).

From a climatic point of view, we observe high diurnal variations in RH<sub>a</sub>, mainly driven by the large amplitude of air temperature variations between night and day (not shown). As discussed above, the significance of relative humidity conditions in terms of evaporation conditions – and deuterium excess control – is only relevant at surface temperature. The variations are smoother for RH<sub>s</sub>, and it more or less follows the absolute amount of water (Figure 5). Under Mediterranean conditions, a distinct increase of q occurs in the morning, indicating a net addition of vapor in the atmosphere. Under North Atlantic conditions, the q increase is lower, but still detectable.

The daily cyclicity of the planetary boundary layer height (PBLH) is specifically related to the different atmospheric conditions. The PBLH attains around 1100m on average during the

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afternoon under North Atlantic and Bay of Biscay conditions, with a significant standard deviation. Under Mediterranean conditions, the PBLH is less variable and remains at almost 700m. Nocturnal values are also very low -less than 100m- in Mediterranean conditions, compared to 400-500m in the other weather situations. Low PBLH in Mediterranean conditions corresponds to weak turbulence and air stability resulting from light winds. while Northerly advection transports a relatively cold air mass over a warmer surface, leading to a weaker stratification.

# **Day-time processes: contribution of local evaporation**

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The diurnal increase in d<sub>v</sub> is a widely observed feature, which reflects the diurnal variation of the water and air mass balances of the planetary boundary layer. In many studies, it has been attributed to entrainment of free atmosphere into the boundary layer (Lai and Ehleringer, 2011; Zhang et al., 2011; Welp et al., 2012; Berkelhammer et al., 2013). However, it could also be due to the addition of locally evaporated vapor, as was observed by Welp et al. (2012). The direction of vapor concentration changes associated with these dv variations can help to identify which of these two processes dominates the PBL water mass budget. Entrainment brings a dryer air into the PBL, and thus causes a day-time decrease in vapor concentration, while an increase in mixing ratio points to the addition of an evaporation flux. Note that the transpiration component of evapotranspiration is not expected to carry high dw, as it has the same isotopic composition as the soil water at steady state (Yakir et Wang, 1996; Williams et al., 2004). In addition, the free atmospheric air is characterized by a isotopically lighter vapor (He and Smith, 1999; Bailey et al., 2013; Berkelhammer, et al 2013), and the dominance of day-time entrainment during day-time is generally associated with a  $\delta_V$  depletion (Lai C.-T. et al., 2006; Lai and Ehleringer, 2011; Tremoy et al., 2012; Bailey et al., 2013; Berkelhammer, et al 2013). Welp et al. (2012), compared six measurement locations and found that both local evapotranspiration and entrainment were involved: in some of their sites, the d<sub>v</sub> daily increase was associated with a  $\delta^{18}O$  decrease during the early morning due to entrainment of free atmosphere into the boundary layer during convective mixing, while the subsequent slight  $\delta^{18}$ O increase came from evapotranspiration. Our data display daytime increases in q,  $\delta^{18}$ O and  $\delta D$  (Figure 5), and thus indicates that the addition of surface vapor is the dominant process governing the d<sub>v</sub> cyclicity during the day.

**Nocturnal processes: dew formation** 

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Supprimé: The planetary boundary layer (PBL) mass budget over land is mainly controlled by the entrainment of free atmosphere, and PBL height has been shown to scale as the entrainment rate (Medeiros et al., 2005). This means that higher boundary layer heights would induce the "dilution" of surface mass fluxes by free atmosphere, when these surface fluxes (e.g. aerosols, atmospheric pollutants) do not directly depend on climatic factors (Quan et al., 2013). On the other hand, the main driver of the daily PBL growth is the buoyancy flux, which is mainly driven by the surface sensible flux and the latent heat flux. the latter being dominant in wetlands. In terms of water vapor mass budget, this suggests that higher PBL should correspond to higher contribution of both local evaporation and free atmosphere, but their respective contributions remain unknown.

The isotopic signature of atmospheric water vapor is a good means to decipher these two potential origins of water. Both processes influence  $d_v$  in the same direction, but not  $\delta_v$ : delta values are expected to decrease with altitude (Bailey et al., 2013; He and Smith, 1999), so that entrainment of free atmosphere would lead to a decrease of  $\delta_v$  during the development of PBL (Bailey et al., 2013; Tremoy et al., 2012), while evapotranspiration produces a more enriched vapor. Daily cyclicity in d<sub>v</sub> is thus a widely observed feature (Lai and Ehleringer, 2011; Welp et al., 2012; Zhang et al., 2011), but can be explained as a consequence of either the addition of vapor from evapotranspiration or of the entrainment of free atmosphere into the boundary layer.

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396 During the night, the PBL subsidence corresponds to the disappearance of both the surface 397 heat fluxes and entrainment. There is thus no further addition of water vapor to the 398 atmosphere. The large-scale motion becomes dominant and, without changing the total PBL 399 mass, redistributes it horizontally (Medeiros et al., 2005). We can thus interpret the nocturnal 400 isotopic values as representing an average signal, onto which are superimposed day-time 401 fluxes. The tendencies identified at daily time step which are interpreted in terms of regional 402 signature, that is to say, depleted (enriched)  $\delta_v$  and higher (smaller)  $d_v$  under North Atlantic 403 (Mediterranean) conditions, are recovered in night-time values (Table 5, Figure 6). However, the isotopic composition is not constant and follows a gradual depletion during the 20h-6h 404 405 UTC period, together with a decrease in water concentration and a stable dv value, which 406 indicates a progressive Rayleigh-condensation process, as was observed by Berkelhammer et 407 al. (2013). The averaged composition of the planetary boundary layer is thus represented by 408 the value measured just after the sunset (20h UTC), when the PBL vanishes and produces a 409 vertical mixing of the atmosphere, and before the influence of dew formation. The dv values 410 remains high during the night (on average 15,1 ±0,5% under Mediterranean conditions and 411 20.9 ±0.7% under North Atlantic conditions), and suggest that the influence of local vapor 412 still affects the average isotopic signal.

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# 4.4 Local isotopic signatures

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426 427 The concomitant increase of q and  $d_v$  during day-time in our average data indicates that the  $d_v$  cyclicity is mainly driven by local vapor fluxes. As stated above, the location of the experimental site close to the main lagoon, and the important availability of open water in the surroundings, point to a dominant contribution of evaporation upon transpiration. In order to calculate the local vapor isotopic composition ( $\delta_E$ ), we use a two-component mixing model based on the linear correlation between 1/q and  $\delta_v$ , derived from the "Keeling plot" method. The validity of this simple model and the possible contribution of regional advection will then be discussed.

The original Keeling approach (Keeling, 1958) was initially proposed to describe the addition of CO<sub>2</sub> into the atmosphere, and then further used to determine the isotopic composition of evapotranspired vapor flux (Yakir and Sternberg, 2000; Yepez et al., 2003; Williams et al., 2004; Wang et al., 2010; Zhang et al., 2010; Noone et al., 2011; Lee et al., 2012; Griffis et al., 2013; Noone et al., 2013). Assuming an isotopically stable background, with an initial mixing

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428 ratio q<sub>0</sub> (mmol/mol) and a vapor composition δ<sub>0</sub>, the addition of a water vapor amount W<sub>1</sub>
 429 (mmol) with a composition δ<sub>1</sub>, leads to the following mixing equations:

$$q = q_0 + W_1 \tag{3}$$

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where q is the mixing ratio (mmol/mol) of the resulting atmosphere. Attributing an isotopic composition to each of these vapor pools, the corresponding isotopic mass balance is:

$$\delta_{v} = (\delta_{0} - \delta_{1}) \cdot \frac{q_{0}}{q} + \delta_{1}$$

$$\tag{4}$$

where  $\delta_{v}$  is the vapor composition of the resulting atmosphere. In a data set characterising the progressive addition of vapor in a closed system, the  $\delta_{v}$  versus 1/q relation displays a linear shape, and the composition of the added vapor  $\delta_{1}$  is deduced from the intercept (for  $q \rightarrow \infty$ ). However, in an open atmosphere where water vapor is added through the mixing with a humid air mass, equation (3) becomes:

$$q = xq_0 + (1 - x)q_1 (5)$$

Where x is the fraction of the initial background air,  $q_1$  is the mixing ratio of the added air mass (mmol/mol), and the corresponding isotopic mass balance is:

$$\delta_{v} = (\delta_{0} - \delta_{1}) \cdot \frac{xq_{0}}{q} + \delta_{1} \tag{6}$$

In a  $\delta_{v}$  versus 1/q plot, the composition of the added vapor is then deduced for x=0, i.e. by extrapolating the linear trend until 1/ $q_1$ .

The application of this two-component mixing model thus requires the knowledge of the vapor content of the added air. If it comes from open water evaporation, we can use the mixing ratio which corresponds to saturation conditions (Noone, 2011, 2012). Indeed, as described by the diffusion model of Craig & Gordon (1965), evaporation results from the diffusion of vapor between a saturation concentration at the water-air interface and open air. We focus our discussion on the two most contrasted meteorological situations, i.e. Mediterranean and North Atlantic air mass origins. Strong correlations are found between average values of  $\delta_{v}$  and 1/q for the period of q increase, i.e 5h-13h UTC for North Atlantic situations and 5h-10h for Mediterranean situations ( $R^2$  values are between 0.85 and 0.96, see Figure 6). The local end-members are calculated for the average value of saturation mixing

(at 13h UTC or 10h UTC), Uncertainties are determined as resulting from error propagation

ratio at the water surface temperature (q<sub>s</sub>), during the daily maximum air water concentrations

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$$q_m = xq_0 + (1-x)q_{ET}$$

The mixing equation is as follows (Yakir and Sternberg, 2000):

$$\delta_m = (\delta_0 - \delta_{ET}) \cdot \frac{xq_0}{q_m} + \delta_{ET}$$

. (4) The mixing trend takes the shape y=ax+b in which  $\delta_{ET}=\delta_m$  when  $q_0=0$ , i.e. for  $q_m=q_{ET}$  the mixing ratio of the humid air end-member. Determination of the isotopic composition of water vapor source thus necessitates prior knowledge of the additional humid air mixing

water vapor source thus necessitates prior knowledge of the additional humid air mixing ratio (Noone, 2012; Noone et al., 2011), although there is some confusion concerning use of the mixing model to infer the iso ... [9]

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Our linear fits are robust for both isotopic species and under both climatic situations (Figure 6). Based on the

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457 of the linear model in the whole range of  $q_8$  standard deviation. The resulting compositions of local end-members ( $\delta_E$ ) are:  $\delta^{18}O=-12.5\%$  (-12.3% to -12.9%);  $\delta D=-48\%$  (-44% to -54%) 458 for North Atlantic conditions,  $\delta^{18}O=-11.8\%$  (-11.0% to -12.5%);  $\delta D=-58\%$  (-46% to -68%) 459 for Mediterranean conditions (Figure 7). These results show the high deuterium excess values 460 characterizing the surface vapor ( $d_v = 52\frac{\text{\%}}{\text{\%}}$  for North Atlantic and  $37\frac{\text{\%}}{\text{\%}}$  for 461 462 Mediterranean conditions). Such high values are consistent with the isotopic composition of 463 open water evaporation estimated from the Craig and Gordon model (Craig and Gordon, 1965; Gat et al, 1994). The high d<sub>v</sub> of evaporated moisture have made it possible to detect its 464 contribution to regional precipitation (Gat et Matsui, 1991, Gat et al, 1994) or to partition the 465 466 composition of a vapor flux into evaporation and transpiration (Williams et al., 2004). In our case, these high d<sub>v</sub> values confirm that evaporation is largely dominant upon transpiration, 467 and the higher dv under North Atlantic air mass conditions is consistent with lower RHs, 468 compared to Mediterranean conditions (Figure 5, Table 5). 469

a simple isotopic mass balance:

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$$\frac{W_E}{W} = \frac{\delta_{\nu(\text{max})} - \delta_{\nu(\text{min})}}{\delta_E - \delta_{\nu(\text{min})}}$$

Where W<sub>E</sub>/W represents the mass (or molar) ratio of local vapor to total atmospheric moisture,  $\delta_{v(min)}$  corresponds to the vapor background and  $\delta_{v(max)}$  is the maximum contribution of local vapor. We find 21% of local vapor in the ambient moisture under North Atlantic conditions, consistently when using  $\delta^{18}O$  and  $\delta D$  compositions, and 43 to 48% under Mediterranean conditions, when using  $\delta D$  and  $\delta^{18}O$  mass balances, respectively.

As a first attempt, we use this  $\delta_E$  composition for interpreting the diurnal amplitude of  $\delta_V$  from

**(7**)

The lower amplitude of diurnal variations observed under North Atlantic condition would thus indicate a lower proportion of local vapor, but, compared to the Mediterranean situation, North Atlantic conditions correspond to stronger average potential evapotranspiration rates: 5.8mm.day<sup>-1</sup> versus 3.8mm.day<sup>-1</sup> (Table 5). This apparent contradiction indicates that the isotopic imprint of local evaporation is diluted by advection, consistently with the high wind speed that prevails under North Atlantic conditions (2.0 m.s<sup>-1</sup> compared to 0.7 m.s<sup>-1</sup> for Mediterranean conditions). The local vapor is thus flushed by advection of dry northward air masses, and since this advected vapor may carry a different isotopic signature than the nocturnal background, a simple two-component model is not sufficient to describe the PBL water balance.

## Christine Vallet 7/5/15 08:46

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# Christine Vallet 1/5/15 12:19

**Supprimé:** Note that this saturation assumption corresponds to limit conditions at the water-atmosphere interface, such that our calculated humid end-member corresponds to a maximum value for the composition of the surface flux.

### Christine Vallet 6/6/15 13:58

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# Christine Vallet 4/6/15 14:41

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### Christine Vallet 5/6/15 13:16

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# Christine Vallet 9/5/15 18:43

Supprimé: The contribution of ET has thus a huge effect on regional  $d_{\nu}$ . In that sense, these results agree with both Welp et al. (2012) indicating that d<sub>v</sub> is not a conservative tracer of oceanic sources, and with Aemisegger et al. (2014), indicating that d<sub>v</sub> is a good proxy of continental recycling.

# Christine Vallet 7/5/15 15:03

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# Christine Vallet 7/5/15 15:03

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# Christine Vallet 7/5/15 15:03

Mis en forme: Police :Non Gras

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# Christine Vallet 3/6/15 15:38

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488 We have performed sensitivity analysis to evaluate the impact of a third component on the use of the  $\delta_v$ -1/q relationship for determining  $\delta_E$ , considering an advected air mass dryer than 489 490 local atmosphere (i.e. on the right hand of observed data in Figure 6). The occurrence of 491 regional advection remains compatible with a linear  $\delta_v$ -1/q trend, but slightly influences the 492 observed slope, and thus, the estimated humid end-member composition: if the advected 493 vapor composition is more enriched than the initial background, it leads to overestimate the  $\delta_E$ 494 value, and vice-versa. The composition of the advected vapor is isotopically heavier than the 495 5:00 UTC value, since the nocturnal dew formation induces a depletion of atmospheric vapor. 496 The use of a two-component mixing assumption would therefore lead to overestimate  $\delta_{\rm F}$ , and 497 the actual average composition of local vapor is expected to plot between this  $\delta_E$  estimate and 498 daily maximum measured  $\delta_v$  (Figure 7). 499 Nevertheless, this simple mass balance approach can be used as a framework for deciphering the complex regional and local influences affecting  $\delta_v$  and  $d_v$ . In addition, the processes 500 501 involved in the diurnal isotopic behaviour of atmospheric vapor can be summarized in see  $\delta^{18}$ O- $\delta$ D plot, which is a good mean to understand the drivers of  $d_v$  variations (Figure 7). The 502 503 two clusters of average vapor data are distributed along two linear trends defined by the q-504 increase period of the day, which also meet the calculated value of  $\delta_E$ . These trends result from the addition of local vapor into the ambient air, but, as discussed above, are also 505 506 influenced by the input (and flushing) of regional advection. The day-time vapor composition 507 then oscillates, but stays around the maximum  $\delta_v$  value, until the abrupt shift that occursent 508 20:00 UTC, during sunset, consistently for both meteorological situations. The  $d_v$  decrease 509 observed when the PBL vanishes corresponds to the vertical mixing of the PBL, as previously 510 stated, and is therefore driven by the mixing with a more regional vapor. The night-time 511 evolution of  $\delta_v$  is thus slightly shifted closer to the meteoric water line, compared to day-time 512 compositions. The progressive condensation that occurs between 20:00 and 5:00 UTC 513 maintains the  $\delta_v$  along a regression line roughly parallel to the LMWL. 514 The stronger advection rate which prevails under North-Atlantic conditions and which 515 smoothed the diurnal amplitude of variation, suggests that the higher average d<sub>v</sub> signature of 516 water vapor (Table 5) results from the combined influences of higher  $d_v$  in  $\delta_E$  and in the 517 regional vapor. Such high d<sub>w</sub> for North-Atlantic air masses can be explained by the longer 518 continental trajectory. A more detailed mass balance including a quantification of advection

### Christine Vallet 9/5/15 11:45

 $\label{eq:supprime:} \textbf{Supprime:} \text{ where } \delta_{ET} \text{ is the composition of local ET flux previously determined, } \delta_{v(min)} \text{ and } \delta_{v(max)} \text{ are, respectively, the minimum (night-time) and maximum (day-time) compositions of atmospheric vapor, and where $Q_{ET}/Q_{BG}$ is the relative contribution of daily vapor flux to the background. We found $Q_{ET}/Q_{BG}$ values of 0.21 for North Atlantic conditions, consistently when using $\delta^{18}O$ and $\delta D$ compositions, and 0.48 and 0.43 for Mediterranean conditions, when using, respectively, $\delta^{18}O$ and $\delta D$ mass balances.$ 

# Christine Vallet 9/5/15 18:33

Supprimé: where H is the maximum PBL height (m),  $T_p$  is the average potential temperature in the PBL, estimated from surface measurements (K), Patm is the atmospheric pressure (Pa), and M<sub>H2O</sub> is the molar mass of water (kg.mol<sup>-1</sup>). Compared to the Mæditerranean situation, North Atlantic contititions correstiond to stronger average potential evapotranspiration rates (5.8mm.day versus 3.8mm.day (see Table 5)), and to a lower daily variation in the total amount of water (8.6mm.day-1 versus 11.1mm.day-1 for W<sub>d</sub>-W<sub>n</sub>), mainly because the PBL remains high during the night. As a consequence, we found  $Q_{ET}/Q_{BG}$  values of 0.66 and 0.34 for North Atlantic and Mediterranean conditions respectively. We are aware that our water balance relies on rough assymptions relative to the average vertical profiles in the PBL. However, the calculation is only slightly sensitive to temperature and pressure variations, and is sufficient to allow comparison of the order of magnitude obtained from isotope and water mass balances, and thus, to evaluate the robustness of the assumptions on which the PBL water balance is based. For Mediterranean situations, the isotope mass balance indicates a higher proportion of local vapor than does the PBL water balance, which suggests a higher concentration of local vapor in the lower part of the PBL. On the contrary, under North Atlantic conditions, the isotopic contribution of the local flux is substantially lower than the water influx calculated from the PBL water balance, which indicates an addition water inflow from advection and/or entrainment. High wind speed under North Atlantic conditions ( $\hat{2}.0 \text{ m.s}^{-1}$  compared to  $0.7 \text{ m.s}^{-1}$  for Mediterranean conditions) suggests that the local atmosphere is affected by the contribution of dry northward air mas ... [10]

# Christine Vallet 9/5/15 18:38

**Supprimé:** by horizontal advection and by entrainment of free atmosphere. The contribution of surface evaporation into the PBL is thus modulated by the contribution of advection to the local water balance

# Christine Vallet 6/6/15 14:00

Mis en forme: Indice

fluxes would allow a more precise determination of the local evaporation composition, and would also help to determine the composition of regional vapor.

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# 5 Conclusions

Our results showed that daily averages of atmospheric vapor compositions are mainly controlled by synoptic and mesoscale weather conditions, related to air mass origin and trajectory. Cold, dry and strong winds coming from the North bring an isotopically depleted vapor. Inversely, warm, wet and light winds coming from the South bring an isotopically enriched vapor. In all situations, dv is higher than the global average of 10, and higher than deuterium excess measured in rainfall, both when considering the small rainfall events during the campaign, or the long-term average composition of regional precipitation. Higher dv is found for Northern air masses (23,2‰) than for Mediterranean air masses (18.6‰), but the amplitude of day-night differences is more important than day-to-day variations, and the drivers of these diurnal variations have to be understood for interpreting the isotopic signal with respect to its relevant spatiotemporal scales. Indeed, a day-time increase in dv can be caused either by entrainment of free atmospheric air, either by local evaporation, both processes having different spatial significances.

Unlike evaporation, the contribution of free atmospheric air would have induced a decrease in absolute vapor concentration, and our twenty-four hour average data clearly indicate that the diurnal  $\delta_v$  cycles are essentially driven by local evaporation. Based on the robust alignment of average data in a  $\delta_v$ -1/q plot for the q-increase periods of the day, we applied a two-component mixing model for estimating the composition of local evaporation ( $\delta_E$ ). This two-component mixing assumption does not account for the influence of regional advection, which modulates the amplitude of diurnal  $\delta_v$  variations. Sensitivity analysis showed that the error associated with the use of a two-component assumption leads to slightly overestimate the  $\delta_E$  values, especially under North-Atlantic conditions. Nevertheless, we found higher  $d_v$  for the local vapor composition under North Atlantic air mass conditions consistent with lower humidity conditions. In addition, the stronger advection rate that prevails under North-Atlantic conditions suggests that, in this situation, the higher average  $d_v$  of water vapor also reflects the signature of regional vapor.

Overall, our data showed that mixing ratio is a better predictor of deuterium excess variations than relative humidity, at both daily and sub-daily time scales, because mixing processes

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### Christine Vallet 25/2/15 17:29

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Christine Vallet 25/2/15 17:29
Supprimé: vapor with a lower d<sub>v</sub>
Christine Vallet 6/3/15 11:48
Supprimé: deuterium excess

### Christine Vallet 4/6/15 09:49

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Average hourly data clearly indicate that this high deuterium excess results from the contribution of local evapotranspiration. Whatever the synoptic situation, the strong diurnal cycles are essentially driven by evapotranspiration,

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# Christine Vallet 4/6/15 11:29

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# Christine Vallet 26/2/15 11:45

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**Supprimé:** very close results for the two main air mass trajectories

# Christine Vallet 26/2/15 11:47

Mis en forme: Indice

551 between different vapor sources have weakened the relevance of relative humidity as an 552 indicator of evaporation conditions. At the daily time step, mixing ratio is a proxy of the air 553 mass origin, since there is a huge contrast between water content of Northern and 554 Mediterranean air masses. At the sub-daily time step, the  $d_v$ -q correlation observed during the q-increase period of the day results from the addition of local evaporation. We have proposed 555 556 a depiction of typical daily cycles of water vapor isotopic composition under different 557 meteorological conditions, which could be used as a framework for further quantitative 558 analysis of vapor sources during specific periods.

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# Acknowledgements

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### Christine Vallet 4/6/15 09:20

Supprimé: In the  $\delta^{18}\text{O-}\delta\text{D}$  plot, average hourly  $\delta_{\nu}$  fluctuates between a maximum value corresponding to the maximum contribution of local vapor during daytime, and a minimum value, corresponding to the night-time background. The night-time background represents the averaged regional composition of the atmosphere, because the PBL mass is horizontally redistributed during the night, and its high deuterium excess indicates that it is still affected by the contribution of local vapor.

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### Christine Vallet 4/6/15 09:53

**Supprimé:** We thus show that d<sub>v</sub> measured in ground level atmospheric vapor carry the isotopic signature of local evapotranspiration and has lost the signature of its initial oceanic vapor source. Our measurements were performed during a summer high pressure period with high daily PET, and further measurements during low pressure situations would be useful in evaluating whether or not the influence of local surface processes remains dominant. This study provides a framework for depicting

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**Supprimé:** non-averaged data corresponding to particular situations, for a better understanding of land-atmosphere water exchanges.

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836 837 between 20 July to 24 August 2011).

Supprimé: calibrated with IAEA international standards by laser spectrometry.

Corresponding

Supprimé: average values and standard

deviations of the

Christine Vallet 2/4/15 15:03

Supprimé: measurements performed

Christine Vallet 3/6/15 13:51 Supprimé: from

Christine Vallet 1/4/15 17:32

Supprimé: Calibrated isotopic composition

Christine Vallet 23/4/15 17:54

Supprimé: value

Christine Vallet 2/4/15 15:04

Supprimé: raw measurements

Supprimé:

Standard deviation (‰)

Christine Vallet 2/4/15 15:05

Supprimé: ±

Christine Vallet 1/4/15 17:36

Tableau mis en forme

Christine Vallet 2/4/15 15:05

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		$\delta_{LS}$	Average and standar	d deviation, of
		(‰ vs VSMOW)	$\delta_{\text{raw}}$ (%0)	),
Standard 1	$\delta^{18}O$	-17.12	-13.89	0.19
Standard 1	$\delta D$	-133.3	-152.6	2.0
Standard 2	$\delta^{18}O$	-7.85	-4.85	0.11
Standard 2	$\delta D$	-53.5	-71.1	1.5
Standard 2	$\delta^{18}O$	0.68	3.41	0.08
Standard 3	$\delta D$	3.7	-12.8	1.8

Table 2. Evaluation of the stability of the calibrations in water vapor concentration and isotopic composition, with mean slope and mean intercept  $\pm$  1 standard deviation obtained for  $\delta^{18}O$  and  $\delta D$  with n calibrations performed from 20 July to 24 August 2011.

	$\delta^{18}{ m O}$	δD	
	Water vapor concentration		
Slope	$-0.16\ 10^{-4} \pm 0.13\ 10^{-4}$	$6.08\ 10^{-4} \pm 0.99\ 10^{-4}$	
Intercept	$-13.68 \pm 0.25$	$-165.4 \pm 2.0$	
$R^2$ (n=23)	$0.26 \le R^2 \le 1.00$	$0.89 \le R^2 \le 0.99$	
	Isotopic composition		
Slope	$1.03 \pm 0.01$	$0.98 \pm 0.01$	
Intercept $-2.83 \pm 0.07$		$16.3 \pm 1.7$	
$R^2$ (n=40) 1.00		1.00	

Table 3. Results of Standard 2 measurements ( $\delta_{std2-m}$ ) normalized to the VSMOW2-SLAP2 scale by using Standards 1 and 3 as working standards, following the IAEA Reference Sheet (IAEA, 2009). Statistics are performed from the set of 40 calibrations. Standard deviation provides the reproducibility, and the root mean square deviation from the known value (RMSE) provides the accuracy of liquid measurements.

	<u>δ</u> <sup>18</sup> O	<u>δD</u>	<u>d</u>
	(% VSMOW)	(% VSMOW)	<u>(‰)</u>
Average of $\delta_{\text{std2-m}}$	<u>-7.82</u>	<u>-53.4</u>	9.16
Standard deviation	<u>0.117</u>	0.89	1.29
<u>RMSE</u>	0.121	0.89	<u>1.30</u>

# Christine Vallet 2/4/15 15:06

Supprimé: Table 3. Accuracy and precision of isotopic liquid water measurement, estimated by considering Standard 2 as a sample which is calibrated with Standards 1 and 3, on a set of 42 calibrations. Accuracy is the deviation of the mean of Standard 2 isotopic composition from its known value, and precision corresponds to the standard deviation.

[12]

Unknown
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Table 4. Determination coefficients (R<sup>2</sup>) and slopes (S) of linear correlations between daily means of water vapor composition ( $\delta^{18}$ O,  $\delta$ D, d<sub>v</sub>) and air temperature (T<sub>a</sub>), mixing ratio (q), relative humidity at air temperature (RH<sub>a</sub>) and relative humidity at water surface temperature (RH<sub>a</sub>).

1	Christine Vallet 23/4/15 17:45					
	Mis en forme: Indice					
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		δ <sup>18</sup> O (‰)		δD (‰)		d <sub>v</sub> (‰)	
		$R^2$	S	$\mathbb{R}^2$	S	$R^2$	S
	T <sub>a</sub> (°C)	0.30	+0.54	0.27	+3.48	0.20	-0.88
ı	q (mmol.mol <sup>-1</sup> )	0.72	+0.38	0.62	+2.39	0.54	-0.66
	RH <sub>a</sub> (%)	0.64	+0.13	0.55	+0.80	0.51	-0.23
	<u>RH<sub>s</sub> (%)</u>	0.33	<u>+0.13</u>	0.31	+0.83	0.19	<u>-0.19</u>

Christine Vallet 23/4/15 17:45
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Automatique, Indice

Tableau mis en forme

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Table 5. Number of days (n) associated with the different origins of air masses (Figure 1). Corresponding mean values of  $\delta^{18}$ O,  $\delta$ D, deuterium excess (d<sub>v</sub>), air temperature (T), mixing ratio (q), relative humidity (RH), wind speed (V) and potential evapotranspiration (PET) measured in the experimental site,

Christine Vallet 6/3/15 12:13

Supprimé: and

### Christine Vallet 6/3/15 12:13

**Supprimé:** associated with the origin of air masses (Figure 1)

	North Atlantic	Mediterranean	Bay of Biscay
	[310°-360°]	[110°-220°]	[220°-310°]
N	10	13	13
$\delta^{18}$ O (‰)	-15.83	-13.13	-14.90
δD (‰)	-103.5	-86.5	-96.3
d <sub>v</sub> (‰)	23.2	18.6	22.9
T <sub>a</sub> (°C)	21.6	23.3	22.3
q (mmol.mol <sup>-1</sup> )	15.0	21.8	18.0
RH (%)	59.5	78.5	68.1
$V(m.s^{-1})$	2.0	0.7	1.6
PET (mm.day <sup>-1</sup> )	5.8	3.8	4.9

Christine Vallet 23/4/15 17:55

Mis en forme: Indice

- 864 Figure 1. Location of the experimental site (star) in Camargue (Rhône River Delta), 170m
- 865 from the Vaccarès lagoon and 12km from the Mediterranean Sea. Coloured arrows represent
- 866 the three main origins and trajectories of air masses affecting the study site, (1) North
- Atlantic, (2) Mediterranean and (3) Bay of Biscay.

- Figure 2. Raw  $\delta^{18}$ O and  $\delta$ D of our three laboratory liquid standards measured at various water
- 870 vapor concentrations. Error bars are smaller than the symbols. Linear regressions within the
- 871 concentration range encountered in Camargue during the field experiment and associated
- slopes and determination coefficients are shown.

873

- 874 Figure 3. Hourly mean values of air temperature (T), mixing ratio (q),  $\delta^{18}$ O,  $\delta$ D, deuterium
- 875 excess (d<sub>v</sub>) from 20 July to 24 August 2011 at the experimental site according to air mass
- origin (see Figure 1). Grey dots are  $\delta_v$  calculated in isotopic equilibrium with precipitation.

877

- 878 Figure 4. Hourly day-time and night-time isotopic composition of atmospheric water vapor in
- according to air mass origin. For reference, the GMWL and LMWL are also plotted.

880

- Figure 5. 24-hour average values of  $\delta^{18}$ O,  $\delta$ D, deuterium excess (d), relative humidity (RH),
- mixing ratio (q) and planetary boundary layer height (PBLH) calculated from 20 July to 24
- 883 August 2011 according to air mass origin. Grey shading indicates standard deviation of each
- 884 average.

885

- Figure 6. Relationships between 24-hour average values of  $\delta_v$  and 1/q according to the two
- 887 main air mass origins, and corresponding linear regressions for the period of q increase (see
- 888 text). Squares indicate the isotopic composition of the local vapor end-member calculated for
- a mixing ratio corresponding to saturation at the water surface temperature.

890

- 891 Figure 7. 24-hour average values of  $\delta_v$  according to the two main air mass origins,
- 892 corresponding linear regressions for the period of q increase (see text), and calculated local
- 893 vapor end-member (see Figure 6 for details). For reference, the LMWL is also plotted.

894