A portable infrared laser spectrometer for flux measurements of trace gases at the geosphere-atmosphere interface

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Abstract

A portable infrared laser absorption spectrometer named SPIRIT (SPectromètre Infra-Rouge In situ Troposphérique) has been set up for the simultaneous flux measurements of trace gases at the geosphere-atmosphere interface. It uses a Continuous Wave (CW) Distributed Feedback (DFB) Room Temperature (RT) Quantum Cascade Laser (QCL) and a patented new optical multi-pass cell. The aim of SPIRIT field studies is to get a better understanding of lands and water bodies to atmosphere exchange mechanisms of GreenHouse Gases (GHG). The analytical procedure to derive concentrations and fluxes are described, as well as the performances of the instrument under field conditions. The ability of SPIRIT to assess space and time dependence emissions of two GHG - nitrous oxide (N₂O) and methane (CH₄) - for different types of ecosystems is demonstrated through in situ measurements on a peatland, on a fertilized soil, and on water body systems. The objectives of these investigations and preliminary significant results are reported.
Keywords: infrared spectroscopy, Quantum Cascade Lasers, Infrared spectrometry, SPIRIT, CH₄, N₂O, flux measurements, chamber method, peatlands, water bodies, fertilized soils

1. **Introduction**

1.1. **Interest for the CH₄ and N₂O monitoring at ground level**

Global warming observed since the last century is mainly due to the anthropogenic increase of the three main long-lived GreenHouse Gases (GHG), namely carbon dioxide (CO₂), methane (CH₄) and nitrous oxide (N₂O). Atmospheric volume mixing ratios for CO₂, CH₄ and N₂O have raised from 275 to 379 ppm, from 715 to 1774 ppb and from 270 to 319 ppb, respectively, between 1750 (pre-industrial era) and 2005 (IPCC 4th assessment report, 2007a). Despite the smaller absolute concentration raises observed for CH₄ and N₂O relative to CO₂, CH₄ and N₂O emissions contribute significantly to current global warming and future trends for climate change because CH₄ and N₂O have a Global Warming Potential (GWP) averaged over 100 years of 25 and 298, respectively, relative to CO₂. Climate change magnitude in the next decades is mainly driven by the unknown future rate of atmospheric concentration increase of CO₂, CH₄ and N₂O and other minor trace gases.

Biogeochemical emissions represent a significant part of the atmospheric sources of these GHG (IPCC 4th assessment report, 2007b,c). Wetlands (such as peatlands) and cultivated soils contribute to a large part of the CH₄ and N₂O emission budget, respectively. CH₄ and N₂O emissions from these ecosystems exhibit strong changes within a short time scale in response to direct anthropogenic perturbations or to climate changes. Feedbacks in response to direct anthropogenic perturbations can be: sensitivity of peatland CH₄ emissions to drainage, to nutrient inputs from agriculture, or to peat extraction; sensitivity of cultivated soil N₂O emissions to mineral fertilizers or organic manure inputs, to soil labour (ploughing, compaction), to watering, or to crops and soils nature. Feedbacks in response to climate changes can be: net peatland carbon storage and CH₄ emissions changes, in response to temperature, soil humidity and biodiversity changes; sensitivity of N₂O emissions from soils to rainfall and temperature changes.

N₂O and CH₄ are mainly produced by the naturally occurring microbial processes of incomplete denitrification or nitrification, and by strictly anaerobic methanogenesis. However, their production processes in soils and the transport mechanisms at the biogeoosphere-atmosphere interface are poorly understood. As a consequence, gas fluxes must be understood in terms of (i) biological (microbial and vegetation) trace gas production/consumption including symbiosis processes and (ii) the physics of transport from soils and plants. Within an ecosystem, N₂O and CH₄ emissions are known to be strongly space (vegetation and soil properties) and time (diurnal and seasonal) dependent as it is the
case for crop fields (Laville et al., 1999; Flessa et al., 2002) and peatlands (Bortolluzi et al., 2006; Comas et al., 2008; Hendriks et al., 2010) with different efflux pathways for peatlands and water bodies (diffusive or sudden bubbling events). The implication is that fluxes of N₂O and CH₄ need to be measured at a local scale with a short integration time (even if chamber methods are used to measure fluxes).

An infrared (IR) laser portable absorption spectrometer called SPIRIT (SPectromètre Infra-Rouge In situ Troposphérique) has been developed for simultaneous volume mixing ratios and flux measurements of the main GHG and other trace gases of interest with significant impacts on the oxidation capacity of the atmosphere. SPIRIT mobility, rapidity for integrating atmospheric trace gases concentration, and sensitivity to cover a wide range of possible flux intensities are its major advantages to enable space and time dependence emissions investigations and detailed process-oriented studies of flux emissions (as a function of biota composition or abiotic parameters). SPIRIT instrument is designed to support three Quantum Cascade Lasers (QCL) operating at chosen wavelengths in the infrared region, depending on the type of the species measured. SPIRIT is currently equipped with one QCL operating for the detection of CH₄ and N₂O as a first step of concept validation, and because these two GHG gases (i) have a strong feedback in climate change in response to emissions changes from the geosphere and (ii) are challenging species to monitor. Their concentrations in ambient air and their emissions from ecosystems are indeed lower than those of CO₂, and many cost effective instruments easy to handle do exist for CO₂ monitoring.

1.2. Current IR spectrometers and main strengths of SPIRIT

IR absorption spectrometry provides the fastest online analysers for atmospheric trace gas concentration measurements. The very high resolution IR absorption spectrometry, possible thanks to laser technology, leads to larger sensitivity, selectivity and precision, compared to lower resolution IR spectrometry such as in Fourier Transform InfraRed (FTIR) spectrometry. Line strengths in the mid infrared region (2.5 - 25 µm) of absorbing atmospheric trace gases are known to be 2 or 3 orders of magnitude higher than in the Near Infra Red region (0.8 - 2.5 µm). Quantum Cascade Lasers (QCLs) are more and more used in the mid infrared region because (i) they provide high and reproducible output power (several tens of mW), (ii) they can now operate at or near Room Temperature (RT), with small Peltier thermoelectric coolers replacing large cryogenic liquid N₂ dewars (Beck et al., 2002), (iii) they present reproducible resonant modes, and (iv) they are tunable over large spectral windows (~10 cm⁻¹) giving versatility for the choice of the species to be measured. Moreover Distributed FeedBack (DFB) QCL, built with a grating inside the cavity, allow for exhibiting an excellent single-mode and a well-defined behaviour. In addition, Continuous Wave (CW) operating mode lasers provide better ease
and higher precision for concentration retrieval, larger selectivity and sensitivity than pulsed mode operating lasers because of a reduced linewidth (Herndon et al., 2007).

Sensitivity from high resolution IR spectrometers are also obtained from a multiple reflection cell made of high reflectivity mirrors, which can behave as (i) a non resonant cavity as it is the case for the patented SPIRIT cell (Robert, 2007) or modified Herriott type cells (Nelson et al., 2004; Kosterev et al., 2000) or (ii) a resonant cavity (Fabry-Pérot type) as it is the case in Cavity Ring Down Spectroscopy (CRDS) (e.g., Picarro Inc., CA, USA) and in Cavity Enhanced Absorption Spectroscopy (CEAS) (Romanini et al., 2006, for CH₄; Bertseva et al., 2002, for N₂O; Los Gatos Research Inc., CA, USA; Hendriks et al., 2008). By opposition to CRDS and CEAS, the non resonant multiple reflection cells provide shorter optical length paths and thus lower sensitivities but higher versatility as regard to the type and the concentration range of species to be measured in the atmosphere within the same cell because (i) the whole mid IR spectrum area can be used with the same type of mirrors (presenting an acceptable reflectivity over a larger band), and (ii) the optical path length can be varied, with a specific ease in the case of the patented SPIRIT cell.

SPIRIT is equipped with the best current laser technology, i.e. CW-DFB-RT-QCL, for measurement of atmospheric trace gases in general (present at or well below the ppb level) with high precision for concentration. For the more abundant species among them such as CH₄ and N₂O, most instruments are equipped with pulsed wave DFB-RT-QCL and a Herriott type Cell, such as the QCL-TILDAS-76 (Nelson et al., 2004; Kroon et al., 2007) model provided by Aerodyne Research Inc.® (ARI, USA) and another instrument developed by Kosterev et al., (2000) used for field studies (Wright, 2006). Some instruments have been recently modified to host CW-DFB-RT-QCLs, such as the ARI instrument (Neftel et al. 2010), because this QCL type is best to provide the required concentration precision at a 10 Hz sampling rate needed for flux measurements of N₂O and CH₄ using the Eddy Covariance method. MIR spectroscopy using CW-DFB-RT-QCLs and long path (>200m) multipass cells have now demonstrated to provide a precision better than 0.1% on the volume mixing ratio of ambient air trace gases. Such precision is needed for flux emission and isotopic ratio measurements (McManus et al. 2010 and 2011).

2. Instrument description and method for concentration retrieval

SPIRIT is a transport wheel instrument (~100 kg; 120 x 80 x 50 cm³). It is supplied in the field by an electricity gasoline generator and can operate within less than 20 min after deployment. The principle of SPIRIT has been explained through the description of a laboratory prototype in Joly et al. (2008). Only the optical cell and the laser component of this prototype have been kept for building the new portable field instrument with better performances described hereafter. In particular, two
measurement channels have been added, the electric current controller and temperature controller of
the lasers, the data acquisition system, the detection system and the method for concentration retrieval
have been redesigned, based on our experience on the balloon-borne instrument SPIRALE operating
with six lead-salt diode lasers simultaneously, which we previously built (Moreau et al., 2005). The
principle of the instrument is schematized in figure 1. The goal is to allow for SPIRIT future operation
with three QCLs (i.e. measurement channels) while keeping the use of one optical cell and two
detectors only – one for the measurement channel and the other one for the reference channel. Thanks
to the home-made QCL emission controller, the lasers will work sequentially, triggered and
synchronized by the data acquisition system. At present, only one DFB CW RT QCL laser (Alpes
Laser) is operating one third of the time, for simultaneous detection of N$_2$O and CH$_4$. This laser
produces a pure single mode radiation in the whole spectral range 1255-1265 cm$^{-1}$ by a cooling
between $+30^\circ$C and $-30^\circ$C with a Peltier thermoelectric cooler included in the QCL housing. A water-
glycol circuit regulated at a constant temperature of 15$^\circ$C is used for cooling the Peltier thermoelectric
cooler. The laser emission window has been designed to be thermally decoupled from the Peltier
thermoelectric cooler hosting the laser component by a 1-cm thick plastic material. Moreover the laser
housing is under vacuum, and therefore condensation is avoided on the emission window. Mode purity
has been checked by observing that null optical transmission from saturated lines of concentrated CH$_4$
and the laser switching-off give equal electric signals. This single mode property leads to a simpler
optical system, as shown in figure 2 and described below, without the need of another grating than the
DFB one. Temperature and current settings are provided by feedback controllers driven by a
numerical electronic module receiving commands from a laptop computer via a RS-232 interface. The
micro-window used for the present study is 1261.87-1262.30 cm$^{-1}$, selected by cooling the QCL at -9.0
$^\circ$C and injecting a current of 290 mA with a saw-tooth ramp of 34 mA repeated at 244 Hz, including a
turn-around of 100 $\mu$s. The laser output power varies between about 20 and 27 mW. The QCL housing
includes an off-axis parabolic mirror of 5 mm focal length, mounted on the same Peltier-cooled copper
mechanical support as the laser itself, in order to ensure optical alignment stability. The optical layout
is presented in figure 2, with a ray traced for the presently single laser. The beam goes out from this
housing through an anti-reflection coated ZnSe window, and is then divided in two parts by a beam
splitter (BaF$_2$). One minor part ($\sim$ 4%) is reflected toward a home-made Fabry-Pérot etalon used as
reference channel for relative spectral calibration. A location is anticipated for a small optical cell
containing gaseous species with well-established spectra to be added on the path of this reference
channel with the aim of absolute wavelength scale calibration. This reference beam is reflected on a
flat mirror and focused onto a 250-µm photovoltaic HgCdTe detector (Judson 519D10) by a spherical
mirror (140 mm radius). The other (major) part of the beam is directed and focused at the entrance of
the optical cell after one reflection over a flat mirror, one reflection over a focusing spherical mirror
(1000 mm radius) and two other ones by flat mirrors. The measurement output beam is collected by a
flat mirror and focused onto a similar detector by a spherical mirror similar as the reference beam. Both detectors are cooled by a Stirling cycle cooler (Thalès Cryogénie RM2-7i/02) mounted in a home-made Dewar. Liquid nitrogen is therefore no more used to cool the detectors.

The detected electric signal is amplified, first by a preamplifier with a fixed gain and secondly, by an automatically self-adjusted gain in the main electronic module. A 16-bit analog-to-digital converter allows for detecting a theoretical minimum absorption of $1/2^{16}$, i.e. $\sim 1.5 \times 10^{-5}$. One hundred and ten measurement spectra followed by eight reference spectra are acquired and averaged. With the 244 Hz frequency, this gives a measurement period of 1.5 s per laser, including the data processing and the transfer to the laptop computer via a RS-232 interface. Contribution from the detector dark current is measured during the two thirds of every period during which the laser is switched off. As shown in figure 3, each scan is composed of 1024 points as a function of time, including about 850 useful points and the other ones containing the dark current level and the transient signal during laser thermalization to the required temperature. This gives the maximum spectral resolution, namely $\sim 5 \times 10^{-4}$ cm$^{-1}$, depending on the micro-window ($\sim 0.43$ cm$^{-1}$) swept by the laser.

The optical cell is a patented multiple-reflection cell (French patent 05/08396 and International patent WO 2007/017570 A1) detailed in a previous paper (Robert, 2007). In brief, it behaves as multiplier of a Herriott cell from which it inherits the excellent opto-mechanical stability, but it is much more compact. Its other main advantages are that it is made of three standard spherical mirrors with great mechanical tolerance, it is easily set up and tuned for adjusting the path length from 20 to 400 m by the rotation of only one mirror, and it allows the use of higher numerical aperture beams. In the configuration used for the present experiments, the total optical path length $L$ is 14040 cm, thanks to 216 paths in a 65 cm long cell. One of the mirrors (the standing one represented on the right of both figures 1 and 2) is slowly and regularly moved by a piezoelectric actuator on a small range (< 100 μm) back and forth at 0.3 Hz frequency. By successive spectra averaging, this motion induces a reduction of unwanted fringes of interferences between the laser emission and the scattered light in the multi-reflection cell. The cell temperature is measured by a Pt100 resistance probe with a precision of 0.01 K. Sampled air is continuously drawn via PFA Teflon tubes (6 mm o.d., 10 m length) through this cell using a scroll pump (XDS10 Boc Edwards), with a flow rate of about 3.0 STP L min$^{-1}$ (Standard Temperature and Pressure: T = 273.15 K, P = 1013.25 hPa). This leads to an effective flow rate of more than 1.2 L s$^{-1}$ in the cell, maintained at a constant reduced pressure of about 30-40 hPa, precisely known ($\pm 0.01$ hPa) using a new gauge (0-130 hPa range, Scaime DL), by tuning a manually actuated dosing valve (EVN116 Pfeiffer Vacuum) located at the cell output, upstream of the pump. This pressure reduction enabled the 3.3 L cell content to be completely renewed in less than 3 s. The low pressure also allows for the trace gas species ro-vibrational lines broadening to be reduced, so that the
lines of different molecules are clearly distinguished and the baseline (i.e. the laser signal without absorption) is easily interpolated, which improves the measurement accuracy and the sensitivity.

The signal scans sampled as a function of time are converted into spectra as a function of wavenumber thanks to the Fabry-Perot etalon. This one generates interference fringes of $1.183 \times 10^{-2}$ cm$^{-1}$ free spectral range, whose optimum positions are detected and fitted by a numerical spline interpolation. The baseline of the signals is deduced from a polynomial interpolation over the full spectrum. The transmission $T(\tilde{\nu})$ as a function of the wavenumber $\tilde{\nu}$ is finally retrieved by dividing the experimental signal by the associated baseline. The N$_2$O and CH$_4$ ro-vibrational lines used in the present study are positioned at the central wavenumbers $\tilde{\nu}_0$ 1261.9874 and 1262.2285 cm$^{-1}$, with line intensities $S(T)$ of $1.044 \times 10^{-19}$ and $2.723 \times 10^{-20}$ cm molecule$^{-1}$ at $T = 296$ K, respectively (Rothman et al., 2009). The concentration retrieval is based on the Beer-Lambert law,

$$C = \frac{-\ln\left[T(\tilde{\nu})\right]}{S(T)g(\tilde{\nu}-\tilde{\nu}_0)L}$$

where $C$ is the molecular concentration (molecules cm$^{-3}$), $g(\tilde{\nu}-\tilde{\nu}_0)$ is the absorption profile (cm), and $L$ is the path length (cm). The principle of the retrieval is to fit the experimental optical depth, $-\ln[T(\tilde{\nu})]$, to the simulated one, $C.S(T)g(\tilde{\nu}-\tilde{\nu}_0)L$, by adjusting the concentration $C$. The fitting is performed by using a linear least-square algorithm. The intensity $S(T)$ includes the molecular partition functions and the lower-state energy of the transition $E''$. $g(\tilde{\nu}-\tilde{\nu}_0)$ is assumed to be a Voigt profile involving a Doppler (Gaussian) profile and a collisional (Lorentzian) profile parameterized by the air broadened half-width $\gamma_{\text{air}}$ and its temperature-dependence exponent $n_{\text{air}}$ calculated from the Hitran 2008 database (Rothman et al., 2009). The key advantage of the CW QCLs with respect to lead-salt diode lasers and pulsed QCLs is the better spectral quality, with a linewidth negligible (determined in the present study to have a HWHM $< 1.5 \times 10^{-4}$ cm$^{-1}$) with respect to the collisional and Doppler broadenings of the molecular line under the actual field pressure and temperature conditions. Thus, the laser linewidth does not need to be accounted for in the experimental absorption profile, which suppresses a systematic error source. Figures 4 (a) and (b) show an example of a transmission spectrum and the associated residual, i.e. the difference between the experimental and the simulated transmission spectra. Baseline signal variations of several origins (e.g., fringes of interferences between the laser emission and the scattered light from optical components with various free spectral ranges, fluctuations of the laser background emission signal, beam wandering on the detector surface) may complicate the retrievals with the direct absorption line fitting method in some cases, because the zero absorption signal is difficult to reconstruct. Another retrieval method consists of applying a numerical derivative that efficiently removes the continuous component of the signal, as already used.
and validated in the case of SPIRALE balloon-borne instrument (for more details see Moreau et al., 2005). The second derivative method is preferred, compared to the direct absorption method, because it transforms absorption lines into symmetrical signatures, which are easier to handle, and reduces the fringes efficiently, which is most useful for species with very low line strengths or in low abundances. Statistical noise is usually smaller in the second derivative due to the associated filtering. This method, illustrated in figures 4 (c)-(d), has been proven to be very useful, leading to accurate data for chemistry and dynamics studies of the lower stratosphere (see e.g., Berthet et al., 2006; Huret et al., 2006; Berthet et al., 2007; Müller et al., 2007; Pirre et al., 2008; Mébarki et al., 2010) as well as for validation of satellite instruments such as Envisat MIPAS and GOMOS, ACE-SCISAT FTS, Odin SMR and Aura-MLS (see e.g., Wang et al., 2007; Renard et al., 2008; Strong et al., 2008; Jégou et al., 2008; Mébarki et al., 2010). In the case of high absorptions such as for CH₄ or N₂O species, both methods lead to the same results and precisions (as shown in figure 4).

The overall uncertainties on the volume mixing ratios have been assessed by taking into account random errors and systematic errors. Sources of random error include the signal variations quoted above and other statistical noise. The theoretical detection limit of \( \sim 1.5 \times 10^{-5} \) in absorption is not achieved, due to these random errors of about \( 1 \times 10^{-4} \) magnitude, as observed as different frequency oscillations in figure 4 (b). For highly absorbing and/or abundant species such as N₂O or CH₄, i.e. absorbing by more than \( 5 \times 10^{-2} \) under ambient atmospheric conditions, a \( 10^{-4} \) precision in absorption leads to a precision of about 0.2% in volume mixing ratio, namely \( \sim 0.6 \text{ ppb} \) for N₂O and \( \sim 4 \text{ ppb} \) for CH₄. This is confirmed by the calculations of 1σ uncertainties for measurement series of at least 10 data (15 s), giving a precision of about 0.2%. This precision is considered as the sensitivity to concentration variations and also approximately as the detection limit for field measurement realistic times. In order to evaluate the SPIRIT performances in terms of time dependence precision and possible drift behaviour, the Allan variance has been calculated, as it is usually the case for on-line laser instruments (see e.g., Werle et al., 1993; Fried et al., 1998, Nelson et al., 2004; Joly et al., 2008; Skrinsky et al., 2009, Neftel et al., 2010, Tuzson et al., 2010). The log-log plot of Allan variance versus integration time shows a linear decay out to about 250 s, as shown for N₂O in figure 5 as an example. This time corresponds to a minimum standard deviation \( \sigma_{\text{Allan}} \) of 0.13 ppb for N₂O (0.040%) and 0.62 ppb for CH₄ (0.031%). This minimum describes the turn-over point where the white noise becomes dominated by additional drift noise, likely due to changes in optical alignment. This theoretical maximum precision could therefore be reached within this integration time (\( \sim 4 \) min) and lasts about 1 h. These results are of great improvement compared to those of our previous laboratory prototype (Joly et al., 2008) where the variance minimum was attained at 1280 s (\( \sim 21 \) min) for a few minutes and with a higher minimum \( \sigma_{\text{Allan}} \) of 0.5 ppb for N₂O and 3 ppb for CH₄ at ambient air concentrations. The main reasons are believed to be the new current and temperature controllers of the
lasers, the new data acquisition system and the new pressure gauge. These SPIRIT’s standard deviations $\sigma_{\text{Allan}}$ were obtained from laboratory room air non constant volume mixing ratios. Similar performances were obtained with ambient air in the field with shorter time integration ($< 30$ min). These performances can be considered as conservative estimates because Allan variance is best determined on samples from gas cylinders with constant mixing ratios.

SPIRIT exhibits performances similar to (i) the ARI QCL spectrometer for N$_2$O: $\sigma_{\text{Allan}}$ of 0.12 ppb (1 Hz), reached after 200 s averaging time (Neftel et al., 2010), and for CH$_4$: $\sigma_{\text{Allan}}$ of 0.23 ppb (1 Hz), reached after 500 s averaging time (Tuzson et al., 2010), and to (ii) the LGR Fast Methane (FMA) Analyser for CH$_4$: $\sigma_{\text{allan}}$ of 0.58 ppb (1 Hz), reached after 500 s averaging time (Tuzson et al., 2010).

Concerning potential systematic errors, they include the retrieval process, the spectroscopic parameters and the non-linearity of the detector response. The latter has been checked to be better than 0.5%. The most important potential systematic uncertainties originate from the spectroscopic data. The spectral micro-window can be chosen so that the Hitran spectroscopic parameters are more accurate than 5% (Rothman et al., 2009). A 5% uncertainty on the molecular line intensity $S(T)$ proportionally propagates in the concentration value uncertainty. The field temperature is close to the reference temperature of Hitran database (296 K) so that the uncertainties on the temperature dependencies of $S(T)$ (determined by the partition functions and $E''$) and the air-broadened coefficient $\gamma_{\text{air}}$ (determined by $n_{\text{air}}$) are negligible. Therefore, the only other important spectroscopic parameter to consider for the uncertainty evaluation is $\gamma_{\text{air}}$ at ground ambient temperature, primarily defining the line shape. The impact of its uncertainty has been evaluated in the concentration retrieval process by a sensitivity analysis using the fitting software. A 5% variation of $\gamma_{\text{air}}$ results in an uncertainty of 2.5% for the concentration. Finally, the precision on the retrieval, illustrated in figures 4 (b) and (d) for CH$_4$, is essentially limited by this $\gamma_{\text{air}}$ accuracy. In addition, concentration retrieval is found to be affected by line broadening effect linked to the air water vapour content as described by Neftel et al. (2010). Water vapour enrichment during field studies is significant enough to affect the gas matrix composition. Our current retrieval algorithm is parameterized with the HITRAN database “dry” air-broadened half-width $\gamma_{\text{dry air}}$ only, so without taking into account the species line broadening by water vapour. Instrument response to rapid changes in water vapour mixing ratio content has been quantified from chamber experiments, within the range of 0.5 to 3% representative from field conditions, leading to an apparent decrease of N$_2$O and CH$_4$ mixing ratios of 1% and 0.5 % at most, respectively, after correction from volumetric dilution by the water vapour. This decrease is observed only above 1% water vapour mixing ratio. One can note that by opposition to the EC (Eddy Covariance) method for flux measurement as used in Neftel et al. (2010), air line broadening correction do not affect significantly derived fluxes from chamber experiments and are usually not taken into account unless very small fluxes need to be quantified.
The sensitivity analysis has been also applied to the other essential parameters influencing the retrieval, i.e. pressure and temperature. Uncertainties of 0.2 hPa and 0.5 K have been found to influence the accuracy of the molecular concentration by 0.3% and 0.15%, respectively. Sensitivity to the outside temperature fluctuations on the retrieval process has also been quantified. Outside temperature fluctuations may induce a slight misalignment of the optical elements, which in turn induces a decrease of the signal intensity received on the detector. This does not significantly (< 1%) change the species volume mixing ratios measured in the ambient air, having in mind that this changes may also come from the natural ambient air variations. For instance, the same values of N₂O ambient air volume mixing ratio (within 1%) were measured within 25°C outside temperature variation in the La Guette peatland (see section 4.1.2. for more details on these field experiments) over a 20 h experiment duration. N₂O ambient air volume mixing ratio was considered to remain constant because La Guette peatland was not observed to be a N₂O surface emitter.

As a conclusion, the overall uncertainty on the volume mixing ratio, calculated as the square root of the quadratic sum of the random and the systematic uncertainties, is estimated to be around 3% (under dry conditions) without taking into account the potential systematic uncertainty of 5% on the line intensity S (or about 6% with the later contribution).

Intercomparisons have been performed between volume mixing ratios obtained from Gas phase Chromatography (GC) and standard pressurized cylinders vs our infrared spectrometer (SPIRIT), over a concentration range representative from flux field measurements. N₂O and CH₄ mixture glass bulbs were used, one filled with ambient air (N₂O: 325 ppb and CH₄: 1990 ppb) and two others filled with N₂O/CH₄ doped dry air (bulb 1: 1.208 ppm N₂O and 4.980 ppm CH₄; bulb 2: 6.028 ppm N₂O and 10.17 ppm CH₄). These volume mixing ratio values derived from SPIRIT were prepared from multiple dilution steps in a vacuum system using certified Baratron® pressure gauges and pure pressurized N₂O and CH₄ cylinders. N₂O and CH₄ were sampled from the bulbs via a 6 mm o.d. PFA Teflon line in which a T Swagelok holds a septum for simultaneous sampling from gas-syringes in order to perform off line GC analysis. Simultaneously with SPIRIT measurements, and for each bulb, four gas samples of 20 mL were injected into 12-mL glass vials sealed with a rubber septum and a silicone septum (Exetainers®, Labco, High Wycombe, UK). Tubes were stored under ambient conditions for less than a month and then analyzed using a gas chromatograph (Model 3800, Varian Inc., Walnut Creek, CA) equipped with ECD (N₂O) and FID (CH₄) detectors, according to the protocols described in Rochette and Bertrand (2007). Figure 6a shows that GC data scattering (Y axis) are significantly larger than SPIRIT data (X axis) for N₂O. This Y axis scattering is better shown on figure 6b where the averaged volume mixing ratios measured for N₂O by GC were (2.7±3.1%, 1σ precision) higher than the ones measured by SPIRIT (0.2%, 1σ precision). SPIRIT’s precision and accuracy have also been checked from two N₂O standards independently prepared from Air Liquide® in dry N₂ pressurized cylinders.
demonstrating good agreement and reproducibility of SPIRIT values over a 6 months period time (figure 6 b). The volume mixing ratio values derived from SPIRIT agree within 2% with the ones calculated from multiple dilution steps in bulbs and provided from pressurized N₂O standard cylinders. GC data for CH₄, not presented here, showed lower values (-6.7±3.5%, 1σ precision), which are also confirmed from bulbs and a certified pressurized 2 ppm (1% precision) cylinder from Air Liquide®. The reproducibility of the N₂O and CH₄ mixing ratios values measured in the laboratory ambient air and during the several field campaigns has also been shown to be very good (variations < 2-3%: 323±8 ppb and 1.90±0.05 ppm), over 30 months period from August 2008 to February 2011.

To summarize, SPIRIT data are accurate by 6 % due to non accurate spectroscopic parameters (line strength and shape) and this accuracy can be reduced to 3% when more accurate spectroscopic parameters will be released in the Hitran database. Correction for water vapour presence should be taken into account in the presence of more than 1% of this species. According to the volume mixing ratio overall uncertainties given by SPIRIT for N₂O and CH₄ (6%, 1σ), by GC-ECD for N₂O (3%, 1σ) and by GC-FID for CH₄ (1%, 1σ) (Rochette and Bertrand, 2007), data are within the accuracy range of the measurements obtained by the two analytical methods. Despite the good accuracy claimed by GC, the off line GC protocol is shown to be less precise than the online SPIRIT infrared spectrometry for volume mixing ratio retrieval as shown in figure 6.

3. Methodology for flux retrieval

3.1. Rationale for the use of a closed chamber method

Two main methods for flux measurements of gases at the soil-atmosphere interface are available (Guenther et al., 1996; Laville et al., 1997). The first ones are chamber methods that are put at the soil surface during the measurement period, usually varying from several to twenty minutes. The flux is derived either (i) from the rate of accumulation of the gas concentration as a function of time for the closed chambers, or (ii) from the gas concentration difference between the inside of the chamber and the ambient air for the open chambers. The second ones are micrometeorological methods: (i) the gradient method (diffusive transport predominating) consists in calculating fluxes from both vertical profiles of gas concentration and horizontal mean wind speed in the first few meters above the surface, (ii) the Eddy Covariance (EC) method (turbulence transport predominating) consists in calculating the vertical flux by the mean product of a gas concentration c and the vertical wind velocity w sampled in one location at high frequency (10 Hz), the time lag between the air sampling and analysis being determined as that giving the maximum correlation coefficient between w and c.

It is now well established that the chamber method remains the best technique to assess spatial variability of emissions and also if a detailed process-oriented study of flux emissions needs to be achieved at the scale of the vegetation type (1 m²) such as in peatlands (Hendriks et al., 2010).
peatlands, strong space and time variations for CH\textsubscript{4} emissions do exist and emissions often need to be studied as a function of biotic (e.g. vegetation, microbiology) or abiotic (e.g. soil temperature, pH, humidity) parameters with the aim to investigate biogeochemical processes involved in the C cycles. EC is known to be the best method for assessing time variability of emissions at an upper scale (100 m\textsuperscript{2}) (Hendriks et al., 2010) and is mainly appropriate if a uniform distribution of the emission sources does exist (such as for N\textsubscript{2}O in a homogenous cultivated field). It has also been shown that soil gradient method for CH\textsubscript{4} emission from peatlands should be use with care and the results are likely reliable only in areas with low water table and shallow roots. According to the strong space variability of the emissions from the ecosystem studied here, to the frequent changes of emissions pathways (diffusive versus sudden strong bubbling events) for some plots of peatlands and water bodies, and to the micro-scale objectives of our investigations, chamber methods, described hereafter, were used. As a consequence, the current frequency of SPIRIT for concentration measurement (0.67 Hz) is needed to assess possible chambers artefacts and emissions pathways despite fluxes are inferred from an accumulation period in the order of minutes. More specifically the dynamic closed chamber method is used, meaning that the optical cell of SPIRIT is connected to the chamber via a PFA Teflon tubes where the headspace chamber air is sampled from and re-injected to the chamber (before and during enclosure) and passing through the SPIRIT optical cell. A sufficient sampled volume rate (3 STP L min\textsuperscript{-1}) is maintained in order to keep a short delay between sampling and analysis, but low enough to avoid a pressure gradient inside the chamber when enclosed.

3.2. Experimental conditions for the derivation of a flux using the closed chamber method

The flux of CH\textsubscript{4} or N\textsubscript{2}O (named \textit{i}) per surface unit \(F_{S_{i}}\) (molecule m\textsuperscript{-2} s\textsuperscript{-1}) is inferred from the slope of the stabilized linear increase of the \(i\) volume mixing ratio \(X_{i}\) with time (from a 3 to 10 minutes period usually) starting from the observable linear increase after enclosure and is given by Eq. (2),

\[
F_{S_{i}} = \frac{P}{k_{B}.T}.h.\frac{dX_{i}}{dt}\]

where \(P\) is the pressure in the chamber (Pa), \(k_{B}\) the Boltzmann constant (1.381×10\textsuperscript{-23} J K\textsuperscript{-1}), \(T\) the absolute temperature (K), \(h\) the mean height of the chamber above the soil surface (m) or more precisely, the volume of the closed loop (reduced to the chamber pressure) to the surface ratio of the soil emitter, \(X_{i}\) the volume mixing ratio of \(i\) in the chamber (unit-less), \(dX_{i}/dt\) the rate of accumulation of \(i\) in the chamber (s\textsuperscript{-1}). Many kinds of units are given to express fluxes of gases according (i) to the field of investigations, such as atmospheric sciences, agronomy, biology, global geochemical cycles models using mass balances of exchanges, and (ii) to the time and space scales where the fluxes are
considered, which depend mainly on their existing time and space stabilities in the considered ecosystem. \( F^s_i \) can also be given by Eq. (3) using practical units:

\[
F^s_i = A \cdot \frac{P}{T} \cdot h \cdot \frac{dX_i}{dt}
\]  

(3)

where the \( A \) constant value provides the desired unit for the \( F^s_i \), as given in Table 1; \( P \) is given in hPa, \( T \) in K, \( h \) in cm, and \( \frac{dX_i}{dt} \) in parts per billion per second (ppb s\(^{-1}\)).

To derive the flux, the slope \( \frac{dX_i}{dt} \) of the gas mixing ratio curve is calculated using the linear regression method because according to the short integration period usually used (6 minutes) no significant difference is observed between the linear regression method, the slope intercept method and the exponential regression method (see Kroon et al., 2008, for accurate derivation methods of fluxes). This approach is used to derive \( \text{CH}_4 \) flux emission from peatlands (Schrier-Uijl et al., 2010). However SPIRIT instrument time resolution (1.5 s) for the online concentration measurement provides a much more precise monitoring of \( \frac{dX_i}{dt} \) compared to the work from Schrier-Uijl et al. (2010) in which \( \frac{dX_i}{dt} \) is derived from an integration period of 5 minutes based on 5 concentration data points after enclosure of the chamber, using a photo acoustic gas monitor with a limited time resolution (1 min).

For flux measurements in peatlands and water bodies a large volume plexiglass transparent chamber (height = 29 cm, internal diameter = 30 cm, volume = 20.5 L) is used due to the size of some vegetation present in peatlands (\( \text{Betula} \) spp or \( \text{Molinia caerulea} \)). A small fan is installed on top of the chamber to homogenize the inside air, as recommended for high chambers, which otherwise could be affected by poor air mixing (Rochette and Eriksen-Hamel, 2008). To insure reliability and high quality of each flux measured with the closed chamber, all tests (\( i.e. \) headspace chamber air ventilation and flow rate sampling) recommended by Davidson et al. (2002) were performed for all kinds of emitting surfaces and all the criteria given by Rochette and Eriksen-Hamel (2008) were also taken into account.

All chosen peatland slots where gas emissions are measured are equipped with a permanent PVC cylinder collar (internal diameter = 30 cm) sunk into the soil (at approximately 10 cm below the water surface). The top of the emerging part of the collar is levelled to the top of the \( \text{sphagnum cuspidatum} \), leading to a typical mean height chamber of 45 cm. The seal between the inside and the outside air is ensured by the weight of the chamber fitted on the top of the collar by the mean of a wetted silicon o-ring sealed on the chamber.

Care was taken to avoid known chamber artefacts as described by Davidson et al. (2002) which may affect significantly the natural emission such as (i) pressure changes during chamber enclosure, (ii) inside and outside chamber pressures drift during accumulation, or (iii) alteration of the natural concentration gradient within the soil (or within the water table) due to the concentration rise in the
headspace of the chamber. To avoid these respective artefacts, (i) the top of the chamber is connected to a short 4 mm outlet exit line open to the outside air and closed few seconds after enclosure, (ii) the pressures inside and outside the chamber are continuously checked during accumulation with a manometer (the stability of pressures and the pressure differential are kept below 0.1 hPa thanks to a vent located on top of the chamber as recommended by Davidson et al. (2002) and Hutchinson and Livingston (2001)), and (iii) the flux measurement integration period is short and usually takes into account the first 6 minutes after enclosure. For flux measurements from water bodies, such as the Loire river or the Pavin volcanic lake, the same chamber is used, fitted to a similar PVC collar partially sunk inside the water and stabilized by a floating belt.

3.3. Precision and accuracy for fluxes

The flux detection limit for N\textsubscript{2}O and CH\textsubscript{4} emission is calculated for a 6 minutes accumulation rate using the two standard deviations of 1.2 ppb and of 8 ppb obtained for the sensitivity to the concentration variation of N\textsubscript{2}O and CH\textsubscript{4}, leading then to a rate of accumulation detection limit for N\textsubscript{2}O and CH\textsubscript{4} of \(3.3 \times 10^{-3}\) ppb s\(^{-1}\) and \(2.2 \times 10^{-2}\) ppb s\(^{-1}\), respectively (corresponding also to the minimum observable positive slope for \(dX/dt\)). Assuming a pressure of 1013 hPa, an absolute temperature of 298 K, a maximum mean height of the chamber above the soil surface of 45 cm (for peatlands), the upper calculated/measured flux detection limit for N\textsubscript{2}O and CH\textsubscript{4}, expressed in usable units for atmospheric sciences, are \(3.7 \times 10^9\) molecule cm\(^{-2}\) s\(^{-1}\) (6.1 \times 10\(^{-2}\) nmol m\(^{-2}\) s\(^{-1}\)) and \(2.5 \times 10^{10}\) molecule cm\(^{-2}\) s\(^{-1}\) (4.1 \times 10\(^{-1}\) nmol m\(^{-2}\) s\(^{-1}\)), respectively; or, when expressed in units found in agronomic, biologic or earth sciences, 1.5 g N-N\textsubscript{2}O ha\(^{-1}\) day\(^{-1}\) (1.7 ng N-N\textsubscript{2}O m\(^{-2}\) s\(^{-1}\)) and 4.2 g C-CH\textsubscript{4} ha\(^{-1}\) day\(^{-1}\) (4.9 ng C-CH\textsubscript{4} m\(^{-2}\) s\(^{-1}\)) for our 45 cm high chamber.

The precision for N\textsubscript{2}O and CH\textsubscript{4} for an individual measured flux, mainly given by the precision on the slope during linear increase of N\textsubscript{2}O and CH\textsubscript{4}, is better than 1% above 10 times flux detection limits. The overall uncertainty for an individual measured flux reach at most 10%, mainly given from (i) the uncertainty of the mean height of the chamber (3 to 8%) explained by the non homogenous ground surface due to the presence of the vegetation in case of peatlands, and from (ii) the overall uncertainty of the volume mixing ratio of N\textsubscript{2}O and CH\textsubscript{4} (< 6%).

4. In situ measurements of N\textsubscript{2}O and CH\textsubscript{4} fluxes in different ecosystems
4.1. CH₄ from peatlands

4.1.1. The research project. The world's peatlands influence the global carbon (C) cycle. In their natural state, peatlands have a cooling effect by sequestering and storing huge amounts of atmospheric CO₂ via peat accumulation, and emit CH₄. Peatlands are the only ecosystem that have stored in the last thousands years a C stock comparable to the total C present in the atmosphere (Falkowski et al., 2000). Northern Sphagnum dominated peatlands contain 20-30% of the world’s soil C stock (Gorham, 1991) and have the greatest peat accumulation potential due to low rates of plant residue decomposition (Thormann et al., 1999). However, strong uncertainties remain on their response to climate change or to anthropogenic perturbations as regard to net carbon exchanges with the atmosphere. The main reason is that the associated response of the global C cycle is still not understood (Belyea and Malmer, 2004). Human activities (drainage, fertilisation, peat extraction) and climate change (temperature increase, drought) expose large stocks of C to less constrained decomposition conditions. Thus peatlands may become significant sources of CO₂ and at the same time do not totally stop to emit CH₄, particularly in sub-arctic areas. Drained peatlands, especially after fertilization, can also become an important source of N₂O. A selected peatland has been chosen to investigate the effect of anthropogenic perturbations on biodiversity changes over time and on net carbon exchanges. The site is the La Guette peatland, a Sphagnum-dominated acidic fen located in the centre of France (Neuvy sur Barangeon, Cher; see Gogo et al., 2010a, for a detailed description of the site). The increased invasion of this peatland in the last decades by native vascular plants (Betula spp and Molinia caerulea) is due to a set of interacting causes, which comprise drainage by pine trees planted over-slope of the peatland, water works probably affecting the water regime, changes in land-use (decreasing ecosystem renewing and increasing forest influence) and increasing nutrient input. As a result faster decaying litter may increase CO₂ release without necessarily decrease CH₄ emissions.

4.1.2. Field measurements. A first measurement campaign was conducted in La Guette’s peatland on March 19, 2009. Space and time distributions of CH₄ emissions as well as efflux pathway (diffusion transport versus bubbling events) were studied, thanks to the high time resolution of SPIRIT. No significant N₂O emission was observed in that site. Figure 7 shows the two efflux pathways. Diffusive emission is the main process of CH₄ emission. It is observed as a linear increase of CH₄ concentration as a function of time resulting in a constant emission flux during the 10 minutes of CH₄ accumulation, as observed in figure 7 for the Sphagnum cuspidatum part of the plot. For flux derivation when only diffusive emission occurs, \( dX/dt \) is calculated using the linear regression method. The increase of CH₄ concentration is sometimes broken by a sharp rise corresponding to a bubble event, as represented by the arrows in figure 7 for the Eriophorum vaginatum part of the plot. Due to the large increase of CH₄ concentration inside the headspace of the chamber during bubble events, inhibition of the diffusive flux may be observed by the attenuation of \( dX/dt \) over time for long accumulation period (20 minutes).
as visually observable in this *Eriophorum vaginatum* part of the plot. Exponential regression method may be used to derive more accurately the diffusive flux. Table 2 shows that CH$_4$ diffusive emissions ranged from 0.29 mg CH$_4$-C m$^{-2}$ h$^{-1}$ to 1.71 mg CH$_4$-C m$^{-2}$ h$^{-1}$ (70-410 g CH$_4$-C ha$^{-1}$ J$^{-1}$) during that day for the 4 types of vegetation. These values, as well as emissions processes, are commented in details in Gogo et al. (2010b). These values are in the range of previously published data and agree with the plant community where CH$_4$ is the most produced and emitted in the wettest area. A second measurement campaign was conducted in La Guette’s peatland on May 26 and 27, 2009 to get a better knowledge of the CH$_4$ dependence emission with vegetation. Four vegetation species (addition of *Molinia caerulea* plots) in four different zones of the peatland were studied. Higher emissions occur for all vegetations compared to March 19 due to the temperature rise of the water layer (14°C) responsible for the stronger micro-biological activities than in March. Gogo et al. (submitted, 2010c) demonstrated the importance of the interaction between vegetation and microbiological activities in the CH$_4$ production. Under conditions of high photosynthesis activities, CH$_4$ emissions become correlated to the soil temperature mainly for vascular plants, *Molinia caerulea* and *Betula* spp. Through their root, exudates vascular plants inject high enough labile carbon to allow for an increase of the microbial activity correlated with temperature, which is more intense for methanogens than for methanotrophs.

4.2. *N$_2$O from fertilized soils*

4.2.1. *The research project.* Crutzen et al. (2008) pointed out that the amount of nitrous oxide emitted by agricultural nitrate fertilizers has been seriously underestimated. N$_2$O emission predictions over different time and regional scales are very difficult to assess because relationships between soil properties, cultivated vegetation species, biotic and abiotic factors and emission magnitude are poorly understood. It is now established that N$_2$O is mostly formed in soil through the microbial processes of nitrification and denitrification (Firestone and Davidson, 1989; Granli and Bøckman, 1994). These processes are controlled by several soil variables, such as the availability of mineral nitrogen (NH$_4^+$ and NO$_3^-$), aeration, temperature, organic/labile carbon content (for denitrification) and pH (Firestone and Davidson, 1989; Davidson et al., 2000). N$_2$O emissions from cultivated field are known to increase with inputs of N in form of mineral fertilizers and organic manure, and with humidity such as anoxic soils. CH$_4$ may also be released if the anoxia is well pronounced. In order to assess (i) the N$_2$O emissions budget from cultivated ecosystems and (ii) the reduction efficiency of N$_2$O emissions by improving agricultural practices according to the Intergovernmental Panel on Climate Change (IPPC 2007), a pilot site is studied at a regional scale near Villebon (Beauce Chartraine and Faux Perche). This site has been chosen because a large spatial distribution of soil properties, humidity conditions,
agriculture kinds and practices, does exist in this 10 km² small territory. Field studies have started in spring 2010 during crop growth and fertilization.

4.2.2. $\text{N}_2\text{O}$ emission tests at INRA-Orléans. $\text{N}_2\text{O}$ emission tests have been performed on the field of INRA-Orléans to prepare the future campaigns on the site of Villebon. For those $\text{N}_2\text{O}$ specific flux measurements, the closed chambers are made of a disk dark lid fitted on a PVC or on a stainless steal collar (internal diameter = 47-49 cm; height = 7.5 cm) as described in Hénault et al (2005). The chambers are sealed with foam rubber compressed with grips. Three parcels have been prepared in order to test strong different level of possible emissions. The soil is sandy (76.5 % of sand) and treatment before first measurements (September 26, 2008) are described on Table 3. The unfertilized soil is the reference parcel. Parcels 1 and 2 had input of organic carbon (horse manure was incorporated on 25 cm depth) and nitrogen (NH$_4$NO$_3$) fertilizers. Soil was compacted with a tractor to obtain a high bulk density in the superficial layer (1.62±0.05 Mg m$^{-3}$). In order to increase artificially $\text{N}_2\text{O}$ emissions and to test possible CH$_4$ emissions under extreme conditions, the parcel 2 has been (i) enriched with a loamy soil sampled from Villebon and (ii) put under strong anoxia by the installation of an impermeable sheet installed at 30 cm below the ground surface. Figure 8 shows that we obtained a high range of $\text{N}_2\text{O}$ flux, from very low to very high, as a function of fertilisation supply and level of anoxia. $\text{N}_2\text{O}$ emissions ranged from 0.2 g N-N$_2$O h$^{-1}$ day$^{-1}$ for the unfertilized soil to 650 g N-N$_2$O h$^{-1}$ day$^{-1}$ for the parcel 2 (highest level of nitrogen supply and strongest anoxia) on September 26 (shortly after fertilization) at the warmest time of the day. The lowest value corresponds to the detection limit for the N$_2$O flux obtained for the 7.5 cm mean height of the chamber. Observation that $\text{N}_2\text{O}$ emission peaks were associated with rainfall events (in our case with irrigation) shortly after fertilizer application is in agreement with the previous reports that emission from agricultural soils is largely controlled by soil aeration and mineral N content (Sehy et al., 2003; Rochette et al., 2008). $\text{N}_2\text{O}$ emissions are strongly dependent on the diurnal cycle in September 26 and October 14, both. The soil temperature increases with the solar irradiation stimulating microbiological activities and thus $\text{N}_2\text{O}$ productions by opposition to peatland flooded soils, where the soil temperature remains constant at a daily scale. Average $\text{N}_2\text{O}$ emissions are also 7 times less intense on October 14 compared to September 26. This drop is more explained by the lack of available nitrate after 19 days rather than the lower air and soil temperature measured on October 14 (20°C for the air temperature at 14h L.T.) than on September 26 (23°C for the air temperature at 14h L.T.).
4.3. CH$_4$ from lakes and rivers

4.3.1. The research project. Although largely documented in tropical floodplains and reservoirs, emission of methane by lakes, rivers and wetlands was long underestimated in part due to large spatial and temporal variability (Pavel et al. 2009). Several recent surveys have indicated high fluxes of methane to the atmosphere notably from temperate water bodies. For instance, between 7 to 80 µmol m$^{-2}$ h$^{-1}$ for wetlands and lakes in the Danube Delta (Pavel et al. 2009), from 8 to 40 µmol m$^{-2}$ h$^{-1}$ for different shallow water lakes studied by Bastviken et al. (2008) and even more than 190 µmol m$^{-2}$ h$^{-1}$ from a hydropower reservoir (Delsongro et al. 2010). In addition, CH$_4$ emissions from lakes account for 6–16% of the non-anthropogenic emissions (Bastviken et al., 2004; St. Louis et al., 2000). Most data on CH$_4$ emission come from the measurement of dissolved gas in water. Continuous rapid measurement by laser absorption spectrometry allowed us to directly determine gas flux across the water surface and potentially distinguish between diffusive and bubbling fluxes.

4.3.2. Field measurements. A preliminary test was realised on the river Loire at Orléans on November 10$^{th}$, 2009. Discharge was about 122 m$^{3}$ s$^{-1}$ (less half the monthly mean discharge). Measured fluxes ranged from 8 to 28 µmol m$^{-2}$ h$^{-1}$ (22-80 g C-CH$_4$ ha$^{-1}$ d$^{-1}$) which compare with the data mentioned above for more organic productive water-bodies. Sources of methane in lakes and floodplains are attributed to sediments, and anoxic bottom layers or plant materials when present. In the globally oxic waters of a turbulent stream like Loire River at Orléans, methane may originate from bank sediments and abandoned channels in less hydrodynamical active area likely influenced by urban waste-waters (Abril et al 2007). Intensive spatial and temporal surveys would be necessary to better document this case study. Preliminary test also performed on a small river nearby the experimental cultivated field of the Villebon (Beauce Chartraine and Faux Perche) in March 18, 2010, show low emissions of CH$_4$ as expected, i.e. 2.8 µmol m$^{-2}$ h$^{-1}$ (7.9 g C-CH$_4$ ha$^{-1}$ d$^{-1}$), but significant emissions of N$_2$O, i.e. 1.8 µmol m$^{-2}$ h$^{-1}$ (11.7 g N-N$_2$O ha$^{-1}$ d$^{-1}$), likely due to drainage of N$_2$O from surrounded fertilized soils (figure 9a). For the first time, flux measurement of CH$_4$ have been performed during daytime on June 23$^{th}$ and 24$^{th}$, 2010, on Lake Pavin, demonstrating the high sensitivity of SPIRIT for low CH$_4$ emission measurements. Figure 9b shows representative linear CH$_4$ rises versus time during flux measurements performed in two places: (1) where the lowest emission was observed (close to the littoral, right part of the plot) and for the highest one (nearby the spillway of the lake, left plot) leading respectively to 0.88 to 4.9 µmol m$^{-2}$ h$^{-1}$ (2.5 and 14.0 g C-CH$_4$ ha$^{-1}$ d$^{-1}$). Average flux from the lake was 2.4 µmol m$^{-2}$ h$^{-1}$ (6.9 g C-CH$_4$ ha$^{-1}$ d$^{-1}$). Low CH$_4$ emissions form Lake Pavin is explained by the very low percentage of shallow epilimnetic sediments due to sharp border slopes, and high average depth.

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SPIRIT has demonstrated its ability to measure a wide range of fluxes in various ecosystems emitting N\textsubscript{2}O and CH\textsubscript{4} with high time and space resolutions, including efflux pathways determination. The essential advantages of this new infrared spectrometer come from the coupling of two last technology advances leading to compactness, ease use, sensitivity, accuracy and robustness: (i) the very high spectral resolution (< 1.5\times10^{-4} \text{ cm}^{-1}) laser sources (QCL) working near room temperature associated with a Stirling cycle cooled detector avoiding the use of liquid nitrogen, and (ii) the multipass optical cell patented by the authors (Robert, 2007).

Thus, SPIRIT is well suited to investigate and to understand detailed processes of production, transport, and emissions of N\textsubscript{2}O and CH\textsubscript{4} as a function of biotic and abiotic parameters including symbiosis between different part of the ecosystems (vegetation, organic matter, bacteria). This is of great importance to get a better understanding of the C and N cycles and to asses their budget in an ecosystem, as a function of soil properties, climate change and anthropogenic forcing. As a consequence, SPIRIT will evolve to support two more QCLs in order to monitor, in addition to N\textsubscript{2}O and CH\textsubscript{4}, fluxes of CO\textsubscript{2} (for C balance determination) as well as trace gases uneasily measurable by other instruments that are of a great importance in climate change. These could be HCHO and HONO as they play a key role in the oxidation capacity of the atmosphere. By addition of a reference cell as described in section 3, SPIRIT could be also upgraded for the direct measurement of isotopic ratios, and then for the $\delta$ value of $^{13}\text{C}/^{12}\text{C}$ isotopic ratio determination for CO\textsubscript{2} (Bowling et al., 2003; Mondelain et al., 2008) and CH\textsubscript{4} (Tsuji et al., 2006), which is of prime importance to study biogeochemical mechanisms involving CO\textsubscript{2} (Rochette et al., 1999) and CH\textsubscript{4} (Whiticar 1999).

We have demonstrated that the use of a high temporal resolution spectrometer (1.5 s) is needed to assess on line measurement artefacts and emission pathways, even for fluxes derived from chamber methods when investigations are made at the vegetation scale. To investigate gas emissions at an upper scale, SPIRIT can be adapted to the Eddy Covariance technique. A sampling rate of 10 Hz is generally preferred for EC in order to obtain the best estimates of fluxes for CH\textsubscript{4} (Hendriks et al., 2008) and for N\textsubscript{2}O (Neftel et al. 2010). Measurements performed at 1 Hz can also generate reliable results (Neftel et al. (2010) and can be sufficient at measurements heights above 2 m for CH\textsubscript{4} and N\textsubscript{2}O in natural and agricultural ecosystems (Edwards et al. 2003). By the installation of a smaller optical absorption cell and minor modifications of the instrument, SPIRIT can meet criteria for EC. SPIRIT’s software and hardware are currently configured to operate with 3 switched lasers with a period of measurement of averaged spectra of 1.5 s. When using a single laser in a continuous mode, this period can be reduced to 0.5 s allowing for a 2 Hz frequency for CH\textsubscript{4} and N\textsubscript{2}O volume mixing ratio without any lost for precision. This frequency can be also increased up to 3 or 4 Hz with only few
modifications of the software by reducing (i) the period of each saw-tooth ramp, implying the decrease of the number of points for each ramp (this solution would decrease slightly the resolution of the spectra) or (ii) the number of averaged saw-tooth ramp (this solution would reduce the signal-to-noise ratio).

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Table 1. Unit expressions for the flux of an i species ($F^S_i$) according to Eq. (3) and associated values of $A$.

<table>
<thead>
<tr>
<th>Value of $A$</th>
<th>Unit expression of $F^S_i$</th>
</tr>
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<tbody>
<tr>
<td>$7.241 \times 10^9$</td>
<td>molecule cm$^{-2}$ s$^{-1}$</td>
</tr>
<tr>
<td>0.1202</td>
<td>n mol m$^{-2}$ s$^{-1}$</td>
</tr>
<tr>
<td>3.366</td>
<td>ng N-N$_2$O m$^{-2}$ s$^{-1}$</td>
</tr>
<tr>
<td>2.910</td>
<td>g N-N$_2$O ha$^{-1}$ J$^{-1}$</td>
</tr>
<tr>
<td>$1.213 \times 10^{-2}$</td>
<td>mg N-N$_2$O m$^{-2}$ h$^{-1}$</td>
</tr>
<tr>
<td>1.443</td>
<td>ng C-CH$_4$ m$^{-2}$ s$^{-1}$</td>
</tr>
<tr>
<td>1.247</td>
<td>g C-CH$_4$ ha$^{-1}$ J$^{-1}$</td>
</tr>
<tr>
<td>$5.196 \times 10^{-3}$</td>
<td>mg C-CH$_4$ m$^{-2}$ h$^{-1}$</td>
</tr>
</tbody>
</table>
### Table 2. Range of methane emission on March 19, in La Guette’s peatland.

<table>
<thead>
<tr>
<th>Vegetation</th>
<th>Flux range</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>mg C m⁻² h⁻¹</td>
<td>(g C ha⁻¹ day⁻¹)</td>
</tr>
<tr>
<td><em>Eriophorum</em></td>
<td>1.08-1.71</td>
<td>(260-410)</td>
</tr>
<tr>
<td><em>Sphagnum cuspidatum</em></td>
<td>0.88-1.00</td>
<td>(210-240)</td>
</tr>
<tr>
<td><em>Betula spp.</em></td>
<td>0.58-0.67</td>
<td>(140-160)</td>
</tr>
<tr>
<td><em>Sphagnum rubellum</em></td>
<td>0.29-0.54</td>
<td>(70-130)</td>
</tr>
</tbody>
</table>
Table 3. Soil treatment before first measurements on September 26, and October 14, 2008.

<table>
<thead>
<tr>
<th>Parcel label</th>
<th>Reference</th>
<th>1</th>
<th>2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ploughing date</td>
<td>15/06/08</td>
<td>15/06/08</td>
<td>15/06/08</td>
</tr>
<tr>
<td>Horse manure on 26/06/08 (t/ha)</td>
<td>0</td>
<td>65</td>
<td>65</td>
</tr>
<tr>
<td>Horse manure on 29/08/08 (t/ha)</td>
<td>0</td>
<td>34</td>
<td>34</td>
</tr>
<tr>
<td>Nitrogen fertilizers on 29/08/08</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>(kg N /ha, solid form)</td>
<td>0</td>
<td>75</td>
<td>75</td>
</tr>
<tr>
<td>Compaction date</td>
<td>08/09/08</td>
<td>08/09/08</td>
<td>08/09/08</td>
</tr>
<tr>
<td>Cylinder collar installation date</td>
<td>15/09/08</td>
<td>15/09/08</td>
<td>15/09/08</td>
</tr>
<tr>
<td>Nitrogen fertilizers on 24/09/08</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(kg N /ha, liquid form)</td>
<td>0</td>
<td>100</td>
<td>150</td>
</tr>
<tr>
<td>Watering (mm)</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>24/09/08</td>
<td>30</td>
<td>30</td>
<td>30</td>
</tr>
<tr>
<td>25/09/08</td>
<td>5</td>
<td>10</td>
<td>10</td>
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<tr>
<td>26/09/08 (8h30 L.T.)</td>
<td>3</td>
<td></td>
<td>3</td>
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<tr>
<td>09/10/2008</td>
<td>15</td>
<td>15</td>
<td>15</td>
</tr>
<tr>
<td>10/10/2008</td>
<td>5</td>
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<tr>
<td>13/10/2008</td>
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</tr>
<tr>
<td>14/10/2008</td>
<td>3</td>
<td>3</td>
<td>3</td>
</tr>
</tbody>
</table>

a mixte with faux Perche loamy soil and put under strong anoxia by the installation of an impermeable sheet installed at 30 cm below the ground surface.
Figure 1. Schematics of the instrument principle
Figure 2. Schematics of the optical bench. The path of the laser beam is symbolized by a red ray.
Figure 3. Average raw signals of 110 measurements (black curve), with 8 superimposed Fabry-Pérot reference signal (gray curve), recorded at 35.80 hPa and 15.90 °C.
Figure 4. (a) Example of experimental (black curve) and simulated (gray squares) transmission spectra with (b) the associated residual, i.e. (experiment – simulated) signal; (c) Experimental (black curve) and simulated second derivatives of the CH₄ (grey diamonds), N₂O (grey triangles) and H₂O (grey squares) transmission spectra with (d) the associated residual. This analysis leads to volume mixing ratios of 2001.1 ppb CH₄, 323.25 ppb N₂O and 0.7379% H₂O, with 1σ statistical uncertainties of 0.50%, 0.42 % and 2.48%, respectively. Pressure was 35.80 hPa and temperature was 15.90 °C.

The high residual for the left H₂O line (around 1261.92 cm⁻¹) is explained by a poor line fit because its spectroscopic parameters are not well characterized in the Hitran database, and the retrieval includes the simultaneous fitting of three H₂O lines (with those at 1262.12 and 1262.27 cm⁻¹).
Figure 5. \( \text{N}_2\text{O} \) Allan plot showing long term precision and drift behaviour using room laboratory air.
Figure 6. (a) N₂O volume mixing ratios: Gas phase Chromatography (GC) and certified N₂O pressurized cylinders versus infra red spectrometry (SPIRIT) (b) Relative difference in per cent versus SPIRIT to show the Y axis scattering of the data.

Diamonds and triangles: GC versus SPIRIT from N₂O prepared bulbs in April 2009; diamonds: 4 samples from N₂O ambient air; triangles: 8 samples from N₂O doped dry air. Good agreement between GC (accuracy < 3%) and SPIRIT (accuracy < 6%) are obtained (more details given in the text). Data from GC are likely overestimated because a high ambient air averaged mixing ratio is
found (341 ppb) for GC diamond data compared to SPIRIT (324 ppb), where the latter one agree more with the expected ambient air volume mixing ratio from IPCC (2007).

Circles and crosses: Certified N₂O pressurized cylinders standards vs SPIRIT; N₂O standards were independently prepared from Air Liquide® in dry N₂ pressurized cylinders where mixing ratios (397±8 ppb and 1490±30 ppb) are certified from the BIPM (Bureau International des Poids et Mesures); Circles data from September 2010 (390 and 1515 ppb given by SPIRIT) and Crosses data from February 2011 (392 and 1520 ppb given by SPIRIT) show excellent reproducibility over 6 months and agreement between standards values (accuracy < 2%) and SPIRIT values (accuracy < 6%).

The full line (slope = 1) in figure 6a represents the line of perfect agreement between GC and standard pressurized cylinders vs our infrared spectrometer (SPIRIT).
Figure 7. CH$_4$ volume mixing ratio (ppb) as a function of Local Time during accumulation in the closed chamber (*Eriophorum vaginatum* plot and *Sphagnum cuspidatum* plot, successively), on March 19, in La Guette’s peatland.
Figure 8. Time series of N$_2$O fluxes for unfertilized soil (reference) and for 2 fertilized soils (parcels 1 and 2) with input of organic carbon (horse manure) and nitrogen (NH$_4$NO$_3$) fertilizers. Parcel 2 is enriched with a loamy soil and put under strong anoxia compared to parcel 1.
Figure 9a: $\text{N}_2\text{O}$ (upper plot) and $\text{CH}_4/7$ (lower plot) volume mixing ratios (ppb) versus time (U.T.) from a river nearby the Villebon experimental cultivated site (Beauce Chartraine and Faux Perche), March 18, 2010.
Figure 9b: CH$_4$ concentrations (ppb) versus time (U.T.) from Lake Pavin in June 24, 2010