Vertical partitioning and controlling factors of gradient-based soil carbon dioxide fluxes in two contrasted soil profiles along a loamy hillslope

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Abstract. In this study we aim to elucidate the role of physical conditions and gas transfer mechanism along soil profiles in the decomposition and storage of soil organic carbon (OC) in subsoil layers. We use a qualitative approach showing the temporal evolution and the vertical profile description of CO\textsubscript{2} fluxes and abiotic variables. We assessed soil CO\textsubscript{2} fluxes throughout two contrasted soil profiles (i.e. summit and footslope positions) along a hillslope in the central loess belt of Belgium. We measured the time series of soil temperature, soil moisture and CO\textsubscript{2} concentration at different depths in the soil profiles for two periods of 6 months. We then calculated the CO\textsubscript{2} flux at different depths using Fick’s diffusion law and horizon specific diffusivity coefficients. The calculated fluxes allowed assessing the contribution of different soil layers to surface CO\textsubscript{2} fluxes. We constrained the soil gas diffusivity coefficients using direct observations of soil surface CO\textsubscript{2} fluxes from chamber-based measurements and obtained a good prediction power of soil surface CO\textsubscript{2} fluxes with an \textit{R}^2 of 92\%.

We observed that the temporal evolution of soil CO\textsubscript{2} emissions at the summit position is mainly controlled by temperature. In contrast, at the footslope, we found that long periods of CO\textsubscript{2} accumulation in the subsoil alternates with short peaks of important CO\textsubscript{2} release. This was related to the high water filled pore space that limits the transfer of CO\textsubscript{2} along the soil profile at this slope position. Furthermore, the results show that approximately 90 to 95\% of the surface CO\textsubscript{2} fluxes originate from the first 10 cm of the soil profile at the footslope. This indicates that soil OC in this depositional context can be stabilized at depth, i.e. below 10 cm. This study highlights the need to consider soil physical properties and their dynamics when assessing and modeling soil CO\textsubscript{2} emissions. Finally, changes in the physical environment of depositional soils (e.g. longer dry periods) may affect the long-term stability of the large stock of easily decomposable OC that is currently stored in these environments.

1 Introduction

Soils play a major role in the global C budget, as they contain 2 to 3 times more C than the atmosphere (Eswaran et al., 1993; Lal, 2003). However, current assessments of the exchange of C between the soil and the atmosphere in response to environmental change are associated with large uncertainties (e.g. Peters et al., 2010). One of the sources of this uncertainty is related to our poor understanding of C dynamics in the deeper layers of the soil profile. Rumpel and Kögel-Knabner (2011) showed that deep soil OC (organic carbon) is highly processed, but that subsoil C fluxes from C input, stabilization and destabilization processes are still poorly constrained. In addition to this, recent work has highlighted the significance of buried OC in depositional setting for the C cycle (e.g. Berhe et al., 2007; Van Oost et al., 2012; Wang et
al., 2014; Wiaux et al., 2014a, b). More specifically, buried OC that is stored in colluvial soils at the bottom of eroding hillslopes (e.g. Stallard, 1998) cannot be assumed to be inert to loss as it can decompose as a result of continued degradation or disturbances such as global warming, desiccation of saturated soils, land use change, and re-excision by gullying (e.g. Van Oost et al., 2012). Some studies suggested an erosion-induced C source along hillslope ranging from 0.37 (Jacinthe and Lal, 2001) to 0.8–1.2 petagram C yr\(^{-1}\) (Lal, 2003). This shows that more quantitative information on the contribution of deep C to soil-atmosphere C exchange as well as an increased understanding of the controlling factors is needed.

There is now significant concern about the contribution of soil organic carbon (OC) to future climate change where a climate change driven acceleration of soil OC decomposition could represent a positive feedback on climate (e.g. Davidson and Janssens, 2006; Frey et al., 2013). Under our temperate climate, temperature increase as well as summer drought would constitute potential climatic changes (IPCC, 1990, 1992) which are supposed to increase OC turnover (e.g. Davidson and Janssens, 2006). Recent studies highlight the importance of soil biophysical conditions that may vary substantially with time and across landscapes (e.g. Dai et al., 2012). These studies have shown that, in addition to the effects of soil moisture, temperature and OC quality, soil physical properties (e.g. gas diffusion barriers) may also exert an important control on soil microbial activity and soil CO\(_2\) fluxes (e.g. Wiaux et al., 2014b; Ball, 2013; Maier et al., 2011). Furthermore, there is empirical evidence suggesting that physical protection (i.e. soil aggregates) is a key factor controlling the long-term stability of OC in soils (e.g. Schmidt et al., 2011). Schmidt et al. (2011) also argued that physical conditions may prevent decomposition of deep OC even if this OC would be easily decomposable under optimal conditions. However, other process studies indicate that subsoil OC represents an important C store that interacts actively with the atmosphere (e.g. Rumpel and Kögel-Knabner, 2011). Understanding the soil physical controls on soil CO\(_2\) fluxes is thus particularly relevant in landscapes with complex topography where buried OC in depositional areas represent a significant part of the total OC stored (e.g. Van Oost et al., 2012; Wang et al., 2014 and Wiaux et al., 2014a).

In a forest ecosystem, Goffin et al. (2014) showed that the upper first 30 cm of a soil profile contributes substantially to the total surface CO\(_2\) flux. However, to our knowledge, a vertical partitioning has not been evaluated in agro-ecosystems or in systems with contrasting soil physical and/or chemical properties. Agro-ecosystems differ from forest ecosystems as litter and A horizons in forest ecosystems are characterized by both a high amount and quality of OC (e.g. Brahy et al., 2002; Goffin et al., 2014), while these horizons have disappeared in crop soils due to erosion, plowing, and export of plant residues (e.g. Wiaux et al., 2014a). Hence, deep OC in forest soils may have a lower contribution relatively to surface CO\(_2\) fluxes given that surface soil horizons enriched in fresh organic matter are more likely to emit more CO\(_2\) than soils in croplands. In addition, the roots network in forests is dense and difficult to remove when installing in situ measurement settings compared to crop soils. This creates interferences when measuring heterotrophic CO\(_2\) fluxes as an indicator of OC turnover (e.g. Davidson et al., 1998; Epron et al., 2006; Fiener et al., 2012).

In this study, we aim to elucidate the role of physical controls on soil–atmosphere CO\(_2\) fluxes and its variation with soil depth for a cultivated soil. To this end, we present a comparative analysis between two contrasting soil profiles along an eroded and cultivated hillslope. Previous work (i.e. Wiaux et al., 2014b), has shown that soil surface CO\(_2\) respiration is highly variable along this hillslope, with 30 % more respiration at the downslope and 50 % more at the backslope, relative to the uneroded summit position. Why some controlling factors have been identified, the role of soil physical controls and of the significance of subsoil OC contributions remain unknown. The specific objectives of this study are as follows: (i) to quantify the relative contribution of soil surface and subsoil OC to CO\(_2\) fluxes through a vertical partitioning of these fluxes; and (ii) to identify the role of soil physical properties using the time series of soil moisture measurements and gas diffusivity at different depths. The selected study site is characterized by two contrasting soils in terms of soil hydrological regimes and soil structure and is representative of the cultivated soils of the Belgian loam belt.

2 Material and methods

2.1 Study site description

The study was carried out in the Belgian loam belt along a cultivated hillslope of 150 m in length (50.6669° N, 4.6331° W). The site has a maritime temperate climate, with an average annual temperature of 9.7 °C and an average annual precipitation of 805 mm. The slope percentage in the backslope area ranges between 8.5 and 16 %, with a mean slope of 12 %. The slope percentage in the convex shoulder area ranges between 4 and 8.5 %, with an average of 6 %. The field was plowed (0–30 cm soil surface layer) every year. Each year, manure and nitrate fertilization was carried out. The previous crop rotation was winter wheat, maize and spring wheat. The study site has been described in detail in Wiaux et al. (2014a, b). For this study, we selected two measurement stations along the hillslope: one at the summit and one at the footslope position. The soil is a Dystric Luvisol type at the summit and a Colluvic Regosol in the depositional area at the footslope (IUSS Working Group WRB, 2007; Wiaux et al., 2014a, b).
In order to characterize the physical and bio-chemical properties of these two soil profiles, we measured soil porosity and soil water retention (SWR) curves. Total OC, labile OC and soil porosity were already characterized by Wiaux et al. (2014a, b) and are illustrated in Fig. 1. Total C (i.e. the sum of organic and inorganic C) was analyzed using an elemental analyzer (Variomax, Elementar GmbH). Instrument precision for total C analyses is 0.05 % C concentration. The samples were then treated with 1 % HCl in order to remove inorganic CaCO3 and were analyzed again with the elemental analyzer. Soil OC concentration was then deduced from the difference between total carbon analyses before and after 1 % HCl treatments. Stable OC was defined as the pool of NaOCl-resistant OC (Siregar et al., 2005). We quantified the stable OC by mixing 3 g of air dried soil with 30 mL of 6 wt % NaOCl (adjusted to pH 8). The NaOCl-treated soil was then washed (shaken and centrifuged) with de-ionized water until the solution was chloride free (i.e. no reaction with AgNO3 occurred). The samples were then dried at 105 °C and homogenized before collecting a subsample for total C measurement by dry combustion. The labile OC pool was defined as the residual OC pool that was not resistant to NaOCl oxidation. Hence, this labile OC pool should be interpreted as easily mineralizable OC under ideal conditions where no other factors play a role in stabilization (e.g. anoxic environment, aggregation, etc).

The total porosity ($\phi$) was already characterized by Wiaux et al. (2014a, b) and is illustrated in Fig. 2. Porosity was measured in the laboratory by weighing 100 cm$^3$ of undisturbed soil cores both at saturation and after oven drying at 105 °C for 48 h. We deduced $\phi$ from the mass of water needed to fill sample pores. We calculated the air-filled porosity ($\varepsilon$) as the difference between $\phi$ and volumetric water content (VWC). We calculated average and standard deviation values on triplicate samples for each depth.

The assessment of SWR curves was carried out following the widely used pressure plate technique: undisturbed soil samples were submitted to several increasing and discrete pressure values inside a closed chamber, with a precise monitoring of soil water content for each pressure level (Richards and Fireman, 1943). We used undisturbed soil cores at 10, 25, 35, 70 and 95 cm depth, with 3 replicates at each depth. We obtained the $\varepsilon_{100}$ and $b$ parameters of the Campbell (1974) SWR model by fitting the model to the SWR observations (Moldrup et al., 2000).

### 2.3 Monitoring of soil CO$_2$, water and temperature

We measured soil CO$_2$ concentrations using custom-built soil CO$_2$ probes. The CO$_2$ sensor in the probe is based on the CARBOCAP® Single-Beam Dual Wavelength non-dispersive infrared (NDIR) technology (GMM221, Vaisala corp., Vantaa, Finland). The analytical precision is a function of both the probe characteristic and the value of the observation. This can be calculated as the sum of 1.5 % of the measurement range and 2 % of the observed value. The sampling head of the CO$_2$ probe is a cylinder of 18.5 mm diameter and 40 mm long, covered with a PTFE (polytetrafluoroethylene) membrane, enabling gas exchange and protection against water infiltration. Since the GMM221 sensors were not designed for wet soil conditions, the sensors were encapsulated into an additional perforated PVC tube, providing additional protection against water (Fig. 1). This tubing method is an adaptation of the technique presented by Young et al. (2009). We inserted these tubes vertically into the soil, after creating boreholes with a diameter that equals the diam-
eter of the PVC tubes. This approach avoids the need to back-fill the bore hole, which will disturb the soil structure and diffusion process. Two rubber stoppers, one at 155 mm from the tube head, and another at the top of the tube, prevented atmospheric air from penetrating into the gas sampling volume. Petroleum jelly on these two rubber stoppers ensured a perfect air- and water-tightness and we verified this under laboratory conditions before using the probes. We used a nylon membrane to avoid soil particles entering the perforated tube and to limit further water infiltration.

We adjusted the concentration ranges of the CO$_2$ probe for each soil depth and for each slope position. This allowed an optimal fit of the probes to the local concentrations. Each probe has to characterize the entire range of values encountered during the seasons while at the same time it should have a sufficiently narrow measurement range to ensure measurement precision. At the summit position, measurements ranged between 0 and 2% at 12, 25, 45 cm depth and between 0 and 5% at 85 cm depth. At the footslope position, measurements ranged between 0 and 5% at 12 cm depth, between 0 and 10% at 25 and 45 cm depth and between 0 and 20% at 85 cm depth.

To avoid vegetation growth and any autotrophic contribution to the soil respiration, we covered the measurement plots with a synthetic permeable geotextile during the complete measurement period. To increase the quality of the soil CO$_2$ concentration data time series, we removed observations where the battery voltage was lower than 11.5 V. We also corrected soil profile CO$_2$ concentrations measurements for temperature variations using the empirical formulas described by Tang et al. (2003). This allowed removing the impact of temperature on the CO$_2$ reading of the CO$_2$ probe, since the CARBOCAP® technology is temperature dependent. The probe manufacturer (Vaisala corp., Vantaa, Finland) provided probe specific parameters values for the correction formulas. We also obtained observations of surface CO$_2$ fluxes by means of a portable infrared gas analyzer with an automated closed dynamic chamber (LI-8100A system, LI-COR, United States), following Davidson et al. (2002). The sampling design of these surface chamber CO$_2$ fluxes measurements on the same study site has been described in Wiaux et al. (2014b).

We monitored soil temperature using a thermistor probe (Therm107, Campbell Scientific Lt., UK). Analytical precision is 0.4°C. We monitored soil volumetric water content (VWC) using Time Domain Reflectometry (TDR) probes based on Topp’s equation (Topp et al., 1980) calibrated in the close vicinity of our study site (Heimovaara, 1993; Garré et al., 2008; Beff et al., 2013).

We recorded water, temperature and CO$_2$ concentration profiles measurements with an automatic data logger (CR1000, Campbell Scientific Lt., UK), connected to a multiplexer (AM16/32, Campbell Scientific, Campbell Scientific Lt., UK).

### 2.4 Overall sampling design

The sampling design is shown in Fig. 4. At each of the two slope positions, we measured soil VWC and CO$_2$ concentrations profiles with three replicates on each measurement depth (Fig. 4). We averaged these triplicates, providing an average value for each soil depth and slope position. This allows to account for the spatial variability of VWC and CO$_2$ concentrations (Maier and Schack-Kirchner, 2014) by extending the measurement footprint to an area of ca. 5 m$^2$.

18 VWC measurement points (six soil depths, three replicates) were collected at each of the two slope positions. VWC was measured at depths of 10, 25, 35, 50, 70 and 95 cm (Fig. 4). CO$_2$ concentrations was measured at depths of 10, 25, 45 and 85 cm. Soil temperature was measured at the same depths (10, 25, 45, 85 cm) but without replicates (Fig. 4). Soil temperature and VWC profiles were calculated using a linear interpolation between the depth specific values within
the profile. We kept the values constant between the sampling point at the top of the profile and the soil surface. The estimation of CO$_2$ concentration profiles is described below (Sect. 2.5).

In order to obtain an equilibrated soil environment around the soil VWC, temperature and CO$_2$ probes, measurements started 1 month after the installation of the probes. At the footslope position, hourly time series of VWC, temperature and CO$_2$ concentrations were recorded from 12 May to 13 December 2012 and from 14 May to 22 November 2013. At the summit position, measurements were recorded for the period from 2 June to 13 December 2012 and from the 14 June to 22 November 2013.

We also performed surface CO$_2$ fluxes measurements at 16 dates (profile and surface sampling time was within a 30 min time interval). Note that the averaged values of CO$_2$ concentration for each observation depth cover the same area as the IRGA chamber network located at the soil surface (Fig. 4). These reference surface CO$_2$ fluxes allowed calibrating parameters of the soil gas diffusion model, ensuring the accuracy of profile CO$_2$ fluxes (Sect. 2.4).

We calculated soil temperature and VWC profiles using a linear interpolation between the depth specific values within the profile. We kept the values constant between the sampling point at the top of the profile and the soil surface.

2.5 Calculation of the CO$_2$ fluxes profiles

We calculated the CO$_2$ flux using Fick’s first law of diffusion according to the gradient method (Eq. (1), e.g. Maier and Schack-Kirchner, 2014):

\[ F_{\text{CO}_2} = -D_s \frac{\partial \text{CO}_2}{\partial z}, \]

where \( F_{\text{CO}_2} \) is the soil CO$_2$ flux [µmol m$^{-2}$ s$^{-1}$], \( D_s \) the diffusivity of CO$_2$ in soil [m$^2$ s$^{-1}$], CO$_2$ the soil CO$_2$ concentration [µmol m$^{-3}$] and \( \frac{\partial \text{CO}_2}{\partial z} \) the vertical soil CO$_2$ gradient (with \( z \) representing the soil depth).

In order to calculate the vertical soil CO$_2$ gradient, we suggest an equation that accounts for curve concavity variations (Eq. 2). Variations in curve concavity in CO$_2$ concentration profiles have already been reported in the literature (e.g. Maier and Schack-Kirchner, 2014). In this study, we built Eq. (2) to consider this issue and improve the model fit to CO$_2$ concentration profiles. We evaluated the performance of this fitting by means of the regression coefficient ($R^2$). When the $R^2$ values were lower than a threshold value of 95%, we considered the CO$_2$ concentration profile as unreliable and we did not retain the resulting CO$_2$ fluxes in the final analysis.

\[ \text{CO}_2(z) = 0.04 + A \left( \frac{1}{1 + e^{-\gamma_1 z}} \right) + \left( \frac{1}{1 + e^{-\gamma_2 (z-d)}} \right) - \left( \frac{1}{2} + \frac{1}{e^{\gamma_2 d} + 1} \right), \]

where \( z \) is the soil depth [cm], \( d \) is the soil depth [cm] at which the sharpness of the curve changes due to a diffusion barrier, \( \gamma_1 \) and \( \gamma_2 \) [cm$^{-1}$] are fitted parameters which characterize the sharpness of the curve above and below the soil depth \( d \), and A [%] is a reference value used to define the fitted asymptotic value of the CO$_2$ concentration at infinite depth. We fitted the \( A, d, \gamma_1 \) and \( \gamma_2 \) parameters for each CO$_2$ profile using the trust-region-reflective optimization algorithm in Matlab$^R$. The derivative of Eq. (2) provided the CO$_2$ gradient ($\frac{\partial \text{CO}_2}{\partial z}$) used in Eq. (1) to calculate the CO$_2$ fluxes. The diffusivity of CO$_2$ in soil, \( D_s \) in Eq. (1), is a function of the diffusivity of CO$_2$ in free air (varying with temperature \( T \) and pressure, e.g. Davidson et al., 2006) and of the gas tortuosity factor (\( \xi \)) (Eq. 3):

\[ D_s = \xi \times 10^{-5} \left( \frac{T + 273}{273} \right)^{1.75}, \]

where \( \xi \) depends on soil physical and hydrological properties. We used the Moldrup et al. (2000) model (Eq. 4) which was shown to provide the most accurate and precise results (Davidson et al., 2006; Goffin et al., 2014);

\[ \xi = \left( 2 \epsilon_{100}^3 + 0.04 \epsilon_{100} \right) \left( \frac{\epsilon}{\epsilon_{100}} \right)^{2 + 3/b}, \]

where \( \xi \) is the gas tortuosity factor, \( \epsilon \) [m$^3$ m$^{-3}$] is the soil air-filled porosity, \( b \) is the slope of the Campbell (1974) soil water retention curve model between \(-100\) and \(-500\) cm H$_2$O water suction, and \( \epsilon_{100} \) [m$^3$ m$^{-3}$] is the soil air-filled porosity at a soil water potential of \(-100\) cm H$_2$O.

CO$_2$ fluxes, as assessed by the gradient based method, were calculated on an hourly timescale, and then integrated on a daily basis. Temperature, VWC, diffusivity and CO$_2$ concentration values were also averaged on a daily basis.

In contrast to other studies (e.g. Pingintha et al., 2010; Turcu et al., 2005), we did not aggregate the soil diffusivity coefficient for the entire soil profile or for an entire soil layer. We considered the vertical distribution explicitly, and integrated Eq. (4) in the finite difference numerical solution of Eq. (1). In this numerical integration, we used a depth increment of 0.1 cm and constrained the surface CO$_2$ concentrations with atmospheric CO$_2$ levels (i.e. 0.04 %). In addition, and contrary to Goffin et al. (2014) and Maier and Schack-Kirchner (2014), we did not calculate the CO$_2$ fluxes from each soil slice based on the difference of CO$_2$ concentrations between the top and the bottom of soil horizons, instead we assessed a continuous profile of CO$_2$ fluxes and production.
We calibrated the diffusion model by adjusting the parameters related to the gas diffusion coefficient (i.e., $b$ and $\varepsilon_{100}$) in such a way that calculated fluxes fit instantaneous CO$_2$ fluxes observations at 16 dates spread along the measurement period. This calibration ensures the consistency, and consequently the precision, of the calculated CO$_2$ fluxes. Comparing the gradient-based CO$_2$ fluxes with directly measured IRGA CO$_2$ fluxes, we obtained a good precision with an $R^2$ of 92% for all soil profiles together (Fig. 5). In addition, the slope of the fit (i.e., 1.05 and 1.22, respectively in 2012 and 2013, Fig. 5) was used to correct the estimated fluxes. The comparison between gradient-based calculation and observed surface CO$_2$ fluxes, which allowed the optimization of the calculated fluxes, is illustrated in Fig. 5.

### 2.7 Vertical partitioning of CO$_2$ fluxes

We partitioned the continuous CO$_2$ flux profiles obtained using Eq. (2) into 10 slides of 10 cm along the soil profile. For each soil slide, we calculated the difference between the top and bottom fluxes. We divided this difference by the total CO$_2$ flux (e.g., the value at the soil surface). This provides the relative contribution in terms of both CO$_2$ production and transfer (in %) of each soil slide to the surface CO$_2$ flux (e.g., Goffin et al., 2014; Maier and Schack-Kirchner, 2014).

In order to allow an easy representation of the temporal dynamic of this vertical partitioning, we averaged values on a timescale of 1 month and a half, representing the beginning or the end of a season. Standard deviation values reflect the variability overtime during each half season.

### 3 Results

#### 3.1 Spatio-temporal analysis of measured soil variables

Figures 6 to 10 show the spatio-temporal variation of soil temperature, moisture, CO$_2$ fluxes, concentrations and diffusion, respectively. All of these values correspond to in situ measurements during a 6 month period in 2013. Similar measurements have been carried out in 2012 and display similar spatio-temporal trends (data not shown). Here, we focus on the temporal dynamics of the measured variables, as well as the shape of the vertical distribution along the soil profile. The relationship between these variables was previously analyzed in Wiaux et al. (2014b) and this is not further discussed here. It should be noted that the comparison of the profile distribution at different dates or of temporal dynamics at different depths is done in a qualitative manner.

During the observation period, the soil temperature (Fig. 6) shows a rather similar evolution at the summit and the footslope, although higher temperatures were observed at the summit profile for some shorter periods (e.g., days 180 to 220 of the year when temperatures are approximately 2 to 3°C higher). The mean daily temperatures at the soil surface ranges between 4 to 28°C at the summit, and between 4 to 25°C at the footslope.

The space-time dynamics of the soil volumetric water content (VWC, Fig. 7) differ substantially between the summit and the footslope profiles. At the footslope, the observed soil VWC at different soil depths varied in a narrow range (0.36 to 0.39 cm$^3$ cm$^{-3}$). In contrast, soil VWC at the summit varied between 0.23 and 0.34 cm$^3$ cm$^{-3}$ for the plow layer (0–30 cm depth) and higher values (approximately 0.39 cm$^3$ cm$^{-3}$) were observed for the rest of the soil profile. The soil at the summit position was wettest during early spring and late fall and driest in the summer. At the footslope, soil VWC reached the saturation level in the early summer after an important rainfall event and then slowly decreased until the early fall and reached saturation again in the late autumn.

In contrast to the VWC, and as expected given the physical dependence of diffusivity on soil water content (Eq. (4), Sect. 2.5), the soil gas diffusivity (Fig. 8) reached its maximum value in the summer at the summit while it was low at the footslope. Soil gas diffusivity was approximately 10 times lower at the footslope than at the summit.

The soil CO$_2$ concentrations at both the summit and the footslope increased gradually from spring to late summer...
Figure 6. Space-time dynamic of soil temperature at the summit (red) and the footslope (black) position in 2013: (a) time series at different depths; (b) profile at different dates. The legend of the curves in relation to the different depths and dates is given in Fig. 7.

Figure 7. Space-time dynamic of soil moisture at the summit (red) and the footslope (black) position in 2013: (a) time series at different depths; (b) profile at different dates.

(Fig. 9a). Thereafter, concentrations dropped again and lowest values were observed in the late autumn.

The ranges of CO$_2$ fluxes obtained for the footslope and summit profiles were very similar (Fig. 10a). However, their temporal distribution was different: the periods characterized by high CO$_2$ fluxes did not occur at the same time and had a different duration. More precisely, at the summit, peaks of CO$_2$ fluxes appear at the early summer and disappear after 1 month, while at the footslope, peaks of CO$_2$ fluxes appear at the early fall and are 30% lower than at the summit but remain constant during 2 months. For all soil profiles, CO$_2$ fluxes decreased with depth and reached null values at approximately 30 cm depth at the summit and approximately 15 cm depth at the footslope.

3.2 Shape and variability of CO$_2$ concentrations and fluxes profiles

The observed soil CO$_2$ concentrations increased with soil depth (Fig. 9b), from the atmospheric value of 0.04% at the surface to concentrations which were two orders of magnitude higher at 100 cm depth (CO$_2(z)$ in Eq. 2). For the measurement period of 6 months considered here, CO$_2$ concentration values at 100 cm depth were three to four times higher at the footslope position than at the summit position. In 2013, these values ranged from 0.86 to 3.46% at the summit position and from 3.68 to 9.12% at the footslope position.

The observed CO$_2$ concentration profiles (Fig. 9b) followed a double exponential trend (Eq. 2). This particular model built in this study to represent soil CO$_2$ concentration profiles (Eq. 2) fits our observations relatively well, with regression coefficients ranging between 97 and 100%. These exponential curve starts approximately at the middle of the profile, and is particularly pronounced at the footslope, reflecting a shift of nearly 4% CO$_2$ between 44 and 100 cm depth. Standard deviations around averaged values of observed hourly CO$_2$ concentrations at each depth are given in Table 1. The small-scale spatial variability is low relative to the mean values of CO$_2$ concentrations, the only exception being the footslope at 25 cm depth where the maximum standard deviation exceeded the maximum mean value.
The CO$_2$ fluxes (Fig. 10) were calculated based on both CO$_2$ concentrations and diffusivity. For all soil profiles (Fig. 10a), CO$_2$ fluxes decreased with depth and reached null values at ca. 30 cm depth at the summit and ca. 15 cm depth at the footslope.

### 3.3 Vertical partitioning of CO$_2$ fluxes

The distribution of the soil CO$_2$ fluxes in the profile is illustrated in Fig. 11. At the summit (Fig. 11a), the relative contribution of the different soil layers was more dynamic in time, with a contribution of the first 10 cm of the soil profile ranging from 80% at the late spring, decreasing to 60% in the early summer, and reaching 40% from late summer to late fall. At the summit (Fig. 11a), the first 30 cm of the soil profile significantly contributed to surface fluxes. This contribution decreased with depth in the late spring and the early summer, but is homogeneously distributed with depth for the rest of the time. At the summit (Fig. 11a), soil layers deeper than 30 cm depth sometimes contributed for up to 20% of the total flux, especially in the autumn. At the footslope (Fig. 11b), 90 to 95% of the surface CO$_2$ fluxes were generated in the first 10 cm of the soil profile. The soil layer between 10 and 20 cm contributed for only 5 to 10%, and the deeper layers did not significantly contribute to the surface fluxes.

### 4 Discussion

#### 4.1 Soil physical control on CO$_2$ emissions

The observed differences between the footslope and summit soil profiles, in terms of the temporal evolution of surface soil CO$_2$ fluxes (Fig. 10), indicate that the controlling factors are not the same. At the summit, the evolution of surface soil CO$_2$ fluxes (Fig. 10) clearly follows the temperature variations (Fig. 6, maximum during the summer). At the footslope, the soil surface CO$_2$ flux was small even when temperature increased and remained relatively small throughout the summer period (Fig. 10). This is most likely related to the high VWC values observed at the footslope (Fig. 7), as it is well known that VWC negatively impacts soil CO$_2$ emis-
Figure 10. Space-time dynamic of soil CO$_2$ fluxes, at the summit (red) and the footslope (black) position in 2013: (a) time series at different depths; (b) profile at different dates. The legend of the curves in relation to the different depths and dates is given in Fig. 7.

Table 1. Range of standard deviation (SD) and mean values of triplicated measured hourly CO$_2$ concentrations at each depth, both at the summit and at the footslope position. This range is indicated by minimum (Min) and maximum (Max) values encountered along time (hourly time series) during the 6 months measurement period. NI means No Information (i.e. due to a lack of replicates to allow reliable mean and SD).

<table>
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<th>Soil depth [cm]</th>
<th>Summit mean [%]</th>
<th>Summit Max [%]</th>
<th>Summit Min SD [%]</th>
<th>Summit Max SD [%]</th>
<th>Footslope mean [%]</th>
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<td>0.30</td>
<td>3.93</td>
<td>0.00</td>
<td>5.32</td>
</tr>
<tr>
<td>45</td>
<td>NI</td>
<td>NI</td>
<td>NI</td>
<td>NI</td>
<td>0.12</td>
<td>3.96</td>
<td>0.00</td>
<td>1.96</td>
</tr>
<tr>
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<td>0.15</td>
<td>2.83</td>
<td>0.00</td>
<td>1.42</td>
<td>0.48</td>
<td>7.52</td>
<td>0.00</td>
<td>2.48</td>
</tr>
</tbody>
</table>

sions (e.g. Webster et al., 2008b; Perrin et al., 2012; Wiaux et al., 2014b; also illustrated by Tran et al., 2015). More precisely, we suggest that VWC is not the only factor controlling CO$_2$ emissions at the footslope, but that the difference between the VWC and the water saturation level of the soil pore spaces, i.e. the water-filled pore spaces, also plays an important role. While the VWC at the footslope remained high throughout the year, we observed that the soil surface CO$_2$ flux dramatically increased when the air-filled pore spaces becomes high enough, which is illustrated by the gas diffusivity exceeding a threshold value of c. 0.1 cm$^2$ d$^{-1}$ (i.e. from day 255 to 305 of year 2013, Fig. 10). Hence, we argue that the occasionally low CO$_2$ emissions at the footslope profile are related to the high VWC, as described in the literature by the bimodal effect of VWC on CO$_2$ emissions (e.g. Davidson et al., 1998; Perrin et al., 2004; Webster et al., 2008b; Castellano et al., 2011; Bauer et al., 2012; Wiaux et al., 2014b).

Indeed, according to these authors, when a threshold VWC value is exceeded, this: (i) strongly limits the transfer of biotic CO$_2$ along the soil profile, and (ii) reduces the production of CO$_2$ in itself due to the lack of oxygen for the microbial community. In both cases, the lower CO$_2$ emissions at the footslope profile relative to the summit are due to gas diffusion limitations (even indirectly in the case of a lack of oxygen), as also suggested by Ball (2013). This stands in sharp contrast to the summit profile where gas can easily diffuse throughout the year and along the entire soil profile (Fig. 8).

In the period preceding the important CO$_2$ emissions (i.e. from day 255 to 305 of year 2013, Fig. 10), the soil CO$_2$ cannot move along the soil profile and accumulates within soil pores. This results in an increase in the CO$_2$ concentration during the early and the late summer, especially below 50 cm depth (Fig. 9), where a compacted soil layer appears (see porosity profile in Fig. 1). This suggests that gas diffusion barriers strongly impact the CO$_2$ concentration profile at the footslope. As a result of these gas diffusion barriers, 90 to 95 % of fluxes occur from the top soil (i.e. the first 10 cm) at this location (Fig. 11). This suggests that contributions of deep soil layers could be higher without these diffusion barriers. This may occur in dry conditions where even compacted soil layers can display a low proportion of water in pore spaces. The permanently high water content (Fig. 7), at least during the period of observations, measured at this downslope location prevents the contribution of deeper soil layers. While this soil profile remains wet all the time, the temporal dynamics of VWC and gas diffusion at the footslope (Figs. 7–8) control the time-dynamic behavior of soil
surface CO$_2$ fluxes (Fig. 10). This is in agreement with recent studies (e.g. Maier et al., 2011; Schmidt et al., 2011; Ball, 2013; also illustrated by Tran et al., 2015) that show that soil physical properties are key to understanding the mechanisms regulating the soil gases emissions. Our study brings new insights by demonstrating the strong linkages between soil physical properties and CO$_2$ emissions based on in situ and depth-explicit observations. However, further work is still needed to better understand the processes controlling microbial inhibition and the gas transfer inhibition incase of soil diffusion barriers.

As a consequence, we argue that the significantly higher CO$_2$ concentrations observed at the footslope, especially for deeper soil layers, are not only related to the large amount of labile OC that was found at this position (shown in Wiaux et al., 2014a, b), but more likely result from the long term accumulation (i.e. during periods with a very low diffusivity) of the CO$_2$ produced by the mineralization of this large labile OC stock. Maier et al. (2011) showed that the CO$_2$ efflux (observed CO$_2$ flux resulting from all transfer and production mechanisms together) can deviate in time from the instantaneous soil respiration (due to micro-organisms metabolic activity) because of the CO$_2$ storage into soil pore spaces. Hence, our data suggest that at the footslope, soil physical properties are the dominant control on surface CO$_2$ fluxes. In other words, while the footslope profile contains more labile OC in the subsoil relative to the summit (Fig. 1, Wiaux et al., 2014a), there is a lower contribution from the subsoil to the overall respiration fluxes due to physical limitations (both low diffusivity and lack of O$_2$).

In summary, our study highlights that the mechanisms that govern soil surface CO$_2$ emissions are highly variable in both space and time. On a well-drained soil at the summit of a hillslope, the observed soil CO$_2$ emissions were directly related to soil microbial respiration and CO$_2$ production (e.g. Wiaux et al., 2014b). However, at the footslope of the hillslope, which is characterized by a different hydrological regime, we observed that the temporal dynamic of soil CO$_2$ emissions were more closely related to physical transfer mechanisms: long periods of CO$_2$ production and accumulation alternate with periods of important release at the soil surface. When considering a situation where gas diffusion is limited, and as a result, also oxygen supply for micro-organisms is low, we argue that oxygen concentration in soil pore spaces is not completely null. Hence, the remaining oxygen allows CO$_2$ production through microbial respiration, especially at the footslope due to the high amount of labile soil OC (Wiaux et al., 2014b). This CO$_2$ then accumulates under the soil diffusion barriers. This accumulated CO$_2$ is then later emitted when VWC decreases under a threshold value which allows a significant gas diffusion, as suggested by Maier et al. (2011) and Ball (2013). The main implication of these observations is that if hydrologic regimes change and that footslope soils become drier (reaching moisture conditions favorable for micro-organisms respiration and gas transfer), there is a large amount of potentially easily decomposable OC stored at depth that can suddenly decompose and be emitted to the atmosphere.

### 4.2 Soil organic carbon storage in downslope deposits

The soil respiration rate can be interpreted as an indicator of soil OC persistence (e.g. Gregorich et al., 1994; Wiaux et al., 2014a, b). However, a further analysis of what occurs along the soil profile is needed to thoroughly answer the question of the persistence of OC. The vertical partitioning of the soil CO$_2$ fluxes, as illustrated in Fig.11, shows that during the observation period, 90 to 95% of the surface CO$_2$ flux originated from the first 10 cm of the soil profile at the footslope. Given the important amount of OC until up to 100 cm depth in our study site (Fig. 1, Wiaux et al., 2014a), this observation is not in agreement with the study of Goffin et al. (2014), who suggested that the relative contribution of a soil layer to the surface CO$_2$ fluxes is related to OC distribution along the soil profile. However, while similarities exist in the physical controls and the method used to calculate the vertical partitioning, the study of Goffin et al. (2014) reports on CO$_2$ production in forest soils. Comparing forest and crop soils is difficult because the important part of the autotrophic respiration originates from roots in forest while this is less important in cropland soils (e.g. Davidson et al., 1998; Epron et al., 2006; Martin and Bolstad, 2009; Webster et al., 2008b; Goffin et al., 2014). Hence, in the case of forest ecosystems, the dense roots network in soil creates interferences when measuring heterotrophic CO$_2$ fluxes, and this has been shown to explain an important part of the vertical distribution of CO$_2$ production along soil profiles in forest ecosystems (Goffin et al., 2014). In addition, the estimation of CO$_2$ production in forest soils is more difficult as turbulent advection needs to be accounted for (i.e. the predominance of non-diffusive transport in the litter layer, Goffin et al., 2014). All these elements make a direct and quantitative comparison between forest and agro-ecosystems difficult. However, we can observe some qualitative similarities between our observations and those of Goffin et al. (2014) in forest soils: (i) surface soil VWC values and dynamics were shown to be a critical factor in accurately estimating soil CO$_2$ production, and (ii) the vertical distribution of CO$_2$ concentration increased with depth while CO$_2$ production decreased with depth. In addition, the substantial contribution of the upper soil layers found here was not related to higher temperatures (Fig. 6), contrary to what was suggested by Takahashi et al. (2004). According to the CO$_2$ concentration and diffusivity profiles (Fig. 8), the relative contribution of the soil layers to the surface CO$_2$ flux is more likely governed by soil physical controls (Ball, 2013) rather than by biological production dependent on thermal energy and OC substrate. Here, soil gas diffusivity strongly decreases from 10 to 40 cm depth (where diffusivity is null) at the two slope positions, and the profile
of CO₂ concentration displays no gradient between 10 and 40 cm depth, particularly at the footslope (Fig. 9).

Our data showed that despite the fact that the footslope profiles generate CO₂ fluxes which exceed those observed at the summit position (demonstrated in Wiaux et al., 2014b), the contribution of soil layers below 10 cm depth is very small (Fig. 11). The OC in the top layer of the soil profile (i.e. 0–10 cm) contributed for approximately 90 % of the total CO₂ flux at the footslope position (Fig. 11). This can be explained by environmental conditions specific to this 0–10 cm layer playing in favor of both microbial respiration and gas diffusion. There are no limitations related to both diffusion barriers and access to the oxygen close to the soil surface. Hence, the only impact of soil VWC on soil respiration is its positive effect as it provides a more easy access for soil micro-organisms to their OC substrate, and to the enhancement of their metabolic activities by water (Akinremi et al., 1999; Castellano et al., 2011; Herbst et al., 2008; Howard and Howard, 1993; Šimůnek and Suarez, 1993). The combination of this high amount and high quality of soil OC (Fig. 1, as described by Wiaux et al., 2014a) with this net positive effect of soil VWC results in a strong increase in microbial respiration rates.

Finally, our results suggest that buried soil OC in colluvial deposits is effectively protected from mineralization below 10 cm depth, which corroborates the assumption of a long-term stabilization of buried OC in colluvial soils as suggested in the literature (e.g. Doetterl et al., 2012; Berhe et al., 2008, 2012; Chaopricha and Marín-Spiotta, 2014). This also corroborates the notion of Schmidt et al. (2011), suggesting that deep soil OC may be protected because of unfavorable physical conditions rather than substrate limitations.

5 Conclusions

In this study, we evaluated the factors controlling soil carbon dioxide fluxes for two soil profiles along a hillslope characterized by contrasting physical and chemical characteristics. At the summit position of the hillslope, the time course of surface soil CO₂ fluxes was strongly related to soil temperature and maximum CO₂ fluxes were observed during the summer. Here, the observed soil CO₂ emissions are directly related to soil micro-organisms respiration and associated with biotic CO₂ production. In contrast, the higher levels of water filled pore space observed at the footslope profiles strongly limited the transfer of biotic CO₂ throughout the soil profile and likely the transfer of O₂ to deeper soil depths. The soil surface CO₂ flux increased substantially during short periods when the gas diffusivity exceeded a threshold value related to sufficient air-filled pore spaces. As a result, the time course of observed soil CO₂ emissions was to a large extent explained by physical transfer mechanisms: long periods of accumulation alternate with shorter periods of important CO₂ release. The vertical partitioning of the soil CO₂ fluxes for the footslope profiles showed that, during the observation period, 90 to 95 % of the surface CO₂ fluxes originated from the first 10 cm of the soil profile. This study highlights the need to consider soil physical properties and their dynamics when estimating and modeling soil CO₂ emissions. When considering changes in hydrologic regimes, e.g. the footslope soils become drier (reaching moisture conditions favorable for micro-organisms respiration and gas transfer), there is a large amount of potentially easily decomposable OC stored at depth that can result in an additional emission of C to the atmosphere.

Author contributions. F. Wiaux designed the experiments, and carried out the research. M. Vanclooster, K. Van Oost and F. Wiaux analyzed the results. F. Wiaux wrote the main part of the paper and prepared the manuscript with contributions from all co-authors.

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References


