Carbon isotopic signature of coal-derived methane emissions to atmosphere: from coalfication to alteration

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Abstract. Currently, the atmospheric methane burden is rising rapidly, but the extent to which shifts in coal production contribute to this rise is not known. Coalbed methane emissions into the atmosphere are poorly characterised, and this study provides representative δ13CCH4 signatures of methane emissions from specific coalfields. Integrated methane emissions from both underground and opencast coal mines in the UK, Australia and Poland were sampled and isotopically characterised. Progression in coal rank and secondary biogenic production of methane due to incursion of water are suggested as the processes affecting the isotopic composition of coal-derived methane. An averaged value of -65 ‰ has been assigned to bituminous coal exploited in open cast mines and of -55 ‰ in deep mines, whereas values of -40 ‰ and -30 ‰ can be allocated to anthracite opencast and deep mines respectively. However, the isotopic signatures that are included in global atmospheric modelling of coal emissions should be region or nation specific, as greater detail is needed, given the wide global variation in coal type.

1 Introduction

Methane emissions from the energy sector have been driven in recent years by the impact of a shift from coal to natural gas, in the US and Europe, whereas in China coal production has increased in this century. Currently, the atmospheric methane burden is rising rapidly (Nisbet et al., 2014), but the extent to which shifts in coal production contribute to this rise is not known. Coalbed methane emissions into the atmosphere are poorly characterised, as they are dispersed over large areas and continue even after the mines’ closure (IPCC, 2006). Methane is emitted in coal processing (crashing and pulverisation) and during the initial removal of the overburden; it can be diluted and emitted through ventilation shafts in underground coal mines, or directly emitted to atmosphere from open-cut coal mining, where releases may occur as a result of deterioration of the coal seam.

For the UN Framework Convention on Climate Change, national emissions are estimated by a “bottom-up” approach, based upon a general equation where the coal production data are multiplied by an emission factor that takes into account the mine’s gassiness, which in turn is related to the depth of the mine and the coal rank (i.e. carbon content of coal) (U.S. EPA, 2013). These modelled estimates are often reported without an error assessment, and therefore the level of accuracy of the emissions is not known. “Top-down” assessment of methane
emissions can be made by chemical transport models constrained by atmospheric measurements (Bousquet et al., 2006; Locatelli et al., 2013). However, this top-down approach provides the total amount of methane emissions into the atmosphere, which has to be distributed among the different methane sources in order to quantify each source contribution.

For methane emissions from fossil fuels (coal and natural gas), the source partitioning is mainly “bottom-up”, based on energy use statistics and local inventories, which might be highly uncertain. Conversely, the “top-down” study of the carbon isotopic composition of methane, which is indicative of the methane origin, provides a valuable constraint on the budget appraisal, allowing different sources in a source mix to be distinguished and their individual strength to be evaluated (Hein et al., 1997; Liptay et al., 1998; Lowry et al., 2001; Townsend-Small et al., 2012).

Measurements of methane mole fractions can be complemented in atmospheric models by typical δ13CCH4 signatures of the main methane sources in order to estimate global and regional methane emissions and assess emissions scenarios throughout the past years (Fung et al., 1991; Miller, 2004; Whiticar and Schaefer, 2007; Bousquet et al., 2006; Monteil et al., 2011; Mikaloff Fletcher et al., 2014). However, even though isotopic values are fairly distinctive for specific methanogenic processes, the variety of production pathways and local environmental conditions that discriminate the methane formation process leads to a wide range of δ13CCH4 values. The global isotopic range for coal is very large, from -80 to -17 ‰ (Rice, 1993), but it can be narrowed down when a specific basin is studied. While there are several studies of isotopic composition of methane generated from coal in Australia, U.S.A. and China (Smith and Rigby, 1981; Dai et al., 1987; Rice et al.; 1989; Aravena et al., 2003; Flores et al., 2008; Papendik et al., 2011), there is a significant lack of information about the isotopic characterisation of methane emissions from coal mines in Europe.

The purpose of this study was to determine links between δ13CCH4 signatures and coal rank and mining setting, and to provide representative 13C signatures to be used in atmospheric models in order to produce more accurate methane emission estimates for the coal exploitation sector.

Process of coalification and parameters affecting the δ13C signature of methane emissions

The process of coalification involves both biochemical and geochemical reactions. The vegetal matter firstly decays anaerobically under water; the simple molecules derived from initial decomposition (i.e. acetate, CO2, H2, NH4+, HS-, long chain fatty acids) are metabolised by fermentative archaea, which produce methane via two methanogenic paths: acetoclastic reaction or CO2 reduction (Whiticar, 1999). Different pathways lead to diverse δ13CCH4 isotopic signatures - methane from acetate is 13C enriched relative to methane from CO2 reduction, ranging from -65 ‰ to -50 ‰ and -110 ‰ to -50 ‰ respectively (Levin et al., 1993; Waldron et al., 1998). With increasing burial and temperatures, coal is subjected to thermal maturation, which implicates more geochemical changes.

As coalification proceeds, the carbon content increases, accompanied by a relative depletion in volatile compounds, such as hydrogen and oxygen, emitted in the form of water, methane, carbon dioxide and higher hydrocarbons through decarboxylation and dehydration reactions (Stach and Murchison, 1982). At higher degrees of coalification and temperature, the liquid hydrocarbons formed in previous stages are thermally cracked to methane, increasing the amount of methane produced (Faiz and Hendry, 2006). Peat and brown coal represent the first stage of the coalification process. The vertical pressure exerted by accumulating sediments converts peat into
lignite. The intensification of the pressure and heat results in the transition from lignite to bituminous coal, and eventually to anthracite, the highest rank of coal (O’Keefe et al., 2013) (see Figure 1).

During peat and brown coal stages, primary biogenic methane is formed and it is mainly dissolved in water or released during burial, as coal is not appropriately structured for gas retention (Kotarba and Rice, 2001). At more mature stages, thermogenic methane is produced by thermal modification of sedimentary organic matter, which occurs at great depths and intensive heat. Following the basin uplift, methane production can be triggered in the shallower sediments by the meteoric water inflow into the coal (secondary biogenic gas) (Rice, 1993; Scott et al., 1994).

The isotopic signature of the methane produced during the coalification process is controlled by the methane origin pathway (Whiticar, 1996). Thermogenic methane is isotopically enriched in $^{13}\text{C}$ ($\delta^{13}\text{C} > -50 \, \%$) compared to biogenic methane, as methanogens preferentially use the lightest isotopes due to the lower bond energy (Rice, 1993). Intermediate isotopic compositions of methane might reflect a mixing between microbial and thermogenic gases or secondary processes. Indeed, many controlling factors co-drive the fractionation process, and several contentions about their leverage still persist in literature. Deines (1980) asserts that no significant trend is observed in the isotopic signature of methane in relation to the degree of coalification. Conversely, Chung et al. (1979) observed that the composition of the parent material does affect the isotopic composition of the methane accumulated. While the link between isotopic composition and coal rank is not that straightforward, studies carried out in different worldwide coal seams confirm a stronger relationship between coal bed gas composition and depth. Rice (1993), using data from Australian, Chinese and German coal beds, shows that shallow coal beds tend to contain relatively isotopically lighter methane when compared to those at greater depths. In the presence of intrusions of meteoric water, secondary biogenic methane, isotopically lighter, can be generated and mixed with the thermogenic gas previously produced. Colombo et al. (1966) documented a distinct depth correlation in the Ruhr Basin coal in Germany, with methane becoming more $^{13}\text{C}$-depleted towards the surface zone, independently from coalification patterns. This tendency can be explained either by bacterial methanogenesis, or by secondary processes such as absorption-desorption of methane. Also Scott (2002), in a study about coal seams in the Bowen Basin, Australia, ascribes the progressive methane $^{13}\text{C}$-enrichment with depth to the meteoritic recharge in the shallowest seams, associated with a higher bacterial activity and a preferential stripping of $^{13}\text{C}$-CH$_4$ by water flow.

The migration of methane from the primary zone as a consequence of local pressure release can affect the isotopic composition, since $^{12}\text{CH}_4$ diffuses and desorbs more readily than $^{13}\text{CH}_4$ (Deines, 1980), but the fractionation effect due to migration is less than 1 % (Fuex, 1980). A much larger variation in the isotopic composition is associated with different methanogenic pathways (variation of ~30 %) and thermal maturation stage (variation of ~25 %) (Clayton, 1998). The measurement of Deuterium, coupled with $\delta^{13}\text{C}$CH$_4$ values, would help to distinguish the pathways of secondary biogenic methane generation, acetoclastic reactions or CO$_2$ reduction (Faiz and Hendry, 2006), but such distinction is beyond the scope of this study.

Overall, the $\delta^{13}\text{C}$ values of methane from coal show an extremely wide range and understanding the processes driving the methane isotopic composition needs to focus on the particular set of geological conditions in each sedimentary basin.

Here we analyse the isotopic signatures of methane plumes emitted to atmosphere, from the dominant bituminous and anthracite mines in Europe and Australia, of both deep and open cut type, to test the theory of isotopic change due to coal rank.
2 Material and Methods

2.1 Coal basins investigated and type of coal exploited

2.1.1 English and Welsh coal mines

Coal-derived methane emissions in the UK are estimated at 66 Gg in 2013 (naei.defra.gov.uk), representing 0.4% of global emissions from mining activity (US EPA, 2012). The major coalfields of England and Wales belong to the same stage in the regional stratigraphy of northwest Europe (Westphalian Stage, Upper Carboniferous). Mining has ceased in many areas. Of those remaining some are located in South Yorkshire (Hatfield and Maltby collieries), where 50% of the coalfield's output came from the Barnsley seam, which includes soft coal overlaying a semi-anthracitic coal and bituminous coal in the bottom portion.

The coal of the South Wales basin exhibits a well-defined regional progression in rank, which varies from highly volatile bituminous coal in the south and east margin to anthracite in the north-west part, and the main coal-bearing units reach 2.75 km in thickness toward the south-west of the coalfield (Alderton et al., 2004). The coal is now preferentially extracted in opencast mines, as the extensive exploitation of the coal has left the accessible resources within highly deformed structures (e.g. thrust overlaps, vertical faults) that cannot be worked by underground mining methods (Frodsham and Gayer, 1999). Emissions from two deep mines, Unity and Aberpergwm, were investigated, where mine shafts reach depths up to approximately 750 m (http://www.wales-underground.org.uk/pit/geology.shtml).

2.1.2 Upper Silesian Coal Basin in Poland

The Upper Silesian Coal Basin extends from Poland to the Czech Republic and is one of the largest coal basins in Europe, with an area of ~7400 km² (Jureczka and Kotas, 1995), contributing 1.3% to global coal-derived methane emissions (US EPA, 2012). Emissions from the Silesian Region of Poland are estimated to be in the range of 450-1350 Gg annually (Patyńska, 2013). The upper Carboniferous coal-bearing strata of this region are associated with gas deposits of both thermogenic and microbial origin, and the methane content and spatial distribution are coal rank related (Kędzior, 2009) - i.e. the sorption capacity of coal in the basin is found to increase with coal rank. Most of the methane generated during the bituminous stage in the coalification process has escaped from the coal source following basin uplift during Paleogene (Kotarba and Rice, 2001) and diffused through fractures and faults occurring in tectonic zones. The late-stage gas generated by microbial reduction of CO₂ in the coal seams at the top of the Carboniferous sequence accumulated under clay deposition in the Miocene (Kędzior, 2009).

2.1.3 Australia: Hunter Coalfield (Sydney Basin)

The Hunter Coalfield is part of the Sydney Basin, on the east coast of New South Wales in Australia, and consists of 3 major coal measures. The deepest is the early Permian Greta Coal Measures, which is overlain by the late Permian Whittingham Coal Measures and upper Newcastle Coal Measures. Throughout the Hunter coalfield all sedimentary strata are gently folded, and the same coal seam can be mined at the ground surface and at depths of several hundred meters. In the region surveyed both the opencast and underground mines are extracting coal from the Whittingham coal measures, which are generally high volatile bituminous coals, although some medium to low bituminous coals are extracted (Ward and Kelly, 2013). The mining operation is on a much larger scale than
in the UK - the total coal production for the Hunter Coalfield was 123.63 Mt in 2011, of which 88.24 Mt was
saleable (State of New South Wales, 2013) - and methane emissions are estimated to contribute 4.6% to global
coal-derived methane (US EPA, 2012). Several studies have attested to the dispersion of thermogenic methane
formed at higher degrees of coalification (i.e. high temperatures and pressures) during the uplift and the
subsequent erosion of the basin, followed by the replenishment of the unsaturated basin with more recently formed
methane of biogenic origin (Faiz and Hendry, 2006; Burra, 2010). Gas emplacement is related to sorption capacity
of coal strata, in particular to the pore pressure regime, which is influenced by the local geological features
(compressional or extensional) of the basin. The south of the Hunter Coalfield is characterized by higher gas
content and enhanced permeability than the northern area, with a large potential for methane production, mainly
biogenic (Pinetown, 2014).

2.2 Sampling and measurement methodology

For isotopic characterisation of the methane sources, integrated methane emissions were assessed through
detection of the offsite downwind plume. In fact, even when emissions are focused on defined locations, such as
vent pipes in underground mines, the methane provenance cannot be localised, since most of collieries are not
accessible. For sample collection and measurements of methane emissions downwind of coal mines in the UK
and Australia the mobile system described by Zazzeri et al. (2015) has been implemented. The system utilises a
Picarro G2301 CRDS (Cavity Ring-Down Spectroscopy) within the survey vehicle, for continuous CH₄ and CO₂
mole fraction measurements, and a mobile module including air inlet, sonic anemometer and GPS receiver on the
roof of the vehicle. The entire system is controlled by a laptop, which allows methane mole fractions and the
methane plume outline to be displayed in real time on a Google Earth platform during the survey to direct plume
sampling. When the plume was encountered, the vehicle was stopped and air samples collected in 3L Tedlar bags,
using a diaphragm pump connected to the air inlet. Samples were taken at different locations along the plume
transect in order to obtain a wide range of methane mole fractions and isotopic signatures in the collected air. The
Upper Silesian basin was surveyed with a Picarro 2101-i measuring continuous CO₂ and CH₄ mole fractions and
δ¹³CO₂ isotopic ratio. Samples were collected on site for analysis of δ¹³CCH₄ isotopic ratio.

The carbon isotopic ratio (δ¹³CCH₄) of bag samples was measured in the greenhouse gas laboratory at RHUL
(Royal Holloway University of London) in triplicate to high precision (±0.05 ‰) by continuous flow gas
chromatography isotope ratio mass spectrometry (CF GC-IRMS) (Fisher et al., 2006). CH₄ and CO₂ mole fractions
of samples were measured independently in the laboratory with a Picarro G1301 CRDS analyser, calibrated
against the NOAA (National Oceanic and Atmospheric Administration) WMO-2004A and WMO-X2007
reference scales respectively. The δ¹³CCH₄ signature for each emission plume was calculated using the Keeling
plot approach, according to which the δ¹³C isotopic composition of samples and the inverse of the relative mole
fractions have a linear relationship, whose intercept represents the isotopic signature of the source (Pataki et al.,
2003). Source signatures were provided with the relative uncertainty, computed by the BCES (Bivariate
Correlated Errors and intrinsic Scatter) estimator (Akritas and Bershady, 1996), which accounts for correlated
errors between two variables and calculates the error on the slope and intercept of the best interpolation line. Mole
fraction data and co-located coordinates were used to map the mole fraction variability using the ArcGIS software.
3 Results and Discussion

While most of the emissions from deep mines come specifically from ventilation shafts, which are point sources, emissions from open-cut mines are wide-spread, and difficult to estimate. However, the objective of this study is not the quantification of emissions, but the assessment of the overall signature of methane released into the atmosphere, made through the sampling of integrated emissions from the whole area. Therefore, even though onsite access to collieries was not possible, by driving around the contiguous area, methane emissions could be intercepted and their mole fractions measured. Table 1 summarises δ^{13}C_{CH4} signatures of all the coal mines and coal basins surveyed with the Picarro mobile system.

3.1 English coal mines

Methane plumes from Hatfield colliery, one of the few UK deep mines still open at the time of this study, were detected during surveys on 10th July and 26th September 2013. The methane desorbed from coal heaps within the colliery area and methane vented from shafts might explain the relatively high mole fractions measured (up to 7 ppm). Keeling plots based on the samples collected (4 on the first and 5 on the second survey) give intercept values of -48.3 ±0.2 and -48.8 ±0.3 ‰ (2SD).

Maltby colliery is one of the largest and deepest mines in England and was closed in March 2013. The coal mine methane was extracted and used for electricity, but over 3 ppm mole fractions were detected in ambient air during both surveys in July and September 2013, giving evidence of methane releases from the recovery system and ventilation shafts. The Keeling plot intercept based on the samples collected in July and in September are -45.9 ±0.3 and -45.4 ±0.2 ‰ (2SD) respectively, ~3 ‰ heavier for Hatfield Colliery on the same sampling days. Both source signatures are in good agreement with the value of -44.1 ‰ observed for desorbed methane of the Barnsley coal seam, in a study conducted by Hitchman et al. (1990b).

Keelingley colliery is a still open deep mine situated in North Yorkshire. A surface drainage plant for electrical power generation has been implemented (Holloway et al., 2005), and methane releases from the power plant might justify the mole fractions peak of 9 ppm that was observed while driving on the north side of the coal mine on 10th July 2013. The Keeling plot analysis of the samples collected indicates a source signature of -46.5 ±0.3 ‰ (2SD).

The highest mole fractions detected around Thoresby mine, in Nottinghamshire, approached 5 ppm. Nine samples were collected, giving a source signature of -51.2 ±0.3 ‰ (2SD).

During the sampling of methane plumes from Daw Mill colliery, the highest methane mole fractions (~5 ppm) were recorded close to the edge of the colliery, whereas, driving downwind of the site at further distances, only background values were measured. Indeed, the estimated methane content of the coal seam exploited in Daw Mill colliery, closed in March 2013, is low (typically about 1.7 m^3/tonne) and, furthermore, a ventilation system is implemented, so that the coal mine methane potential is curtailed (Drake, 1983). Keeling plot analysis reveals a source signature of -51.4 ±0.2 ‰ (2SD).

3.2 Welsh coal mines

Methane emissions from coal mines in Wales were sampled in order to characterise methane release from a different rank of coal. The area investigated extended from Cwmllynfell to Merthyr Tydfil. The South Wales coalfield is estimated to have the highest measured seam gas content in the UK (Creedy, 1991).
3.2.1 Deep Mines

The deep mine Aberpergwm, which closed in December 2012, did not operate coal mine methane schemes and methane was vented up to the surface as part of standard operation systems (Holloway et al., 2005). That is consistent with methane mole fractions peaks of 6 ppm observed when approaching the colliery. Air samples were also collected near Unity deep mine, Wales’ largest drift mine, which reopened in 2007 and is located in the town of Cwmgwrach, only 1.5 km away from Aberpergwm. Isotopic source signatures of \(-33.3 \pm 1.8 \%_0\) and \(-30.9 \pm 1.4 \%_0\) result from the Keeling plots based on samples collected respectively near Aberpergwm and Unity colliery, both highly $^{13}$C enriched relative to all English collieries.

3.2.2 Opencast Mines

The Picarro mobile system was driven around opencast mines at Cwmllynfell and Abercrave, in the Swansea Valley. Up to 3 ppm methane mole fractions were recorded near the two mines, which were closed in the 1960s as drift mines and are currently exploited as opencast mines. Our measurements confirm that they are still emitting methane, even though the methane mole fractions recorded downwind of the mines were not significantly above background. $^{13}$C signatures between \(-41.4 \pm 0.5\) and \(-41.2 \pm 0.9 \%_0\) (2SD) result from isotopic analysis of samples collected downwind of the opencast mines, approximately 10 \% lower than the isotopic signature characterizing Welsh deep anthracite mines.

3.3 Polish coal mine

A Picarro mobile survey of the Upper Silesian basin took place on 10th June 2013 and 12 air samples were collected for isotopic analysis. All the mines in this area are deep mines, exploiting the coal at depths ranging from 300 to 900 m. The Keeling plot analysis includes 8 samples collected around the area of Radoszow, downwind of the KWK Wujek deep mine shafts. Methane mole fractions in the range of 3-5 ppm were measured in the majority of the area of Katowice and over 20 ppm mole fractions were detected when transecting the plume originating from the exhaust shafts, which confirms the high level of methane that the mine contains. A source signature of \(-50.9 \pm 0.6 \%_0\) (2SD) was calculated by Keeling plot analysis, which is consistent with the values obtained for the deep mined English bituminous coal.

3.4 Australia: Hunter Coalfield

On 12th and 18th March 2014 12 samples in total were collected along the route in the Hunter Coalfield, where the bituminous coal strata of the Sydney Basin are extracted both in opencast and underground mines. The methane plume width was in the range of 70 km (Fig. 2). A maximum mole fraction of 13.5 ppm was measured near a vent shaft associated with the Ravensworth underground mine (see white star in Fig. 2b). The source signature calculated by the Keeling plot analysis based on all the samples collected during both surveys (grey markers in Fig. 3c) is \(-66.4 \pm 1.3 \%_0\) (2SD) and this signature likely includes a mixture of methane derived both from underground and opencast mines. Two samples collected downwind of the Ravensworth ventilation shaft (red pushpins in Fig. 2b) fall off the Keeling plot trend for March 2014 and are not included in the calculation, because they are likely dominated by methane from the vent shaft and are not representative of the regional mixed isotopic signature.
In January 2016, 10 samples were collected downwind of a ventilation fan in the Bulga mine (second white star in Fig. 2b). The Keeling plot for these (black circles in Fig. 3c) indicates a δ\(^{13}\)C source signature of -60.8 ±0.3 ‰ (2SD). The samples collected next to the Ravensworth ventilation shaft in 2014 fit on this Keeling plot, suggesting that the δ\(^{13}\)CCH\(_4\) isotopic signature of emissions from underground works in the Hunter Coalfield is consistent.

### 3.5 New representative δ\(^{13}\)CCH\(_4\) isotopic signatures for coal-derived methane

The δ\(^{13}\)CCH\(_4\) isotopic values for coal have been found to be characteristic of single basins, but general assumptions can be made to characterise coal mines worldwide. Table 2 provides the literature δ\(^{13}\)CCH\(_4\) isotopic values characteristic of specific coal basins. The isotopic signatures of emissions from English bituminous coal are consistent with the range of -49 to -31 ‰ suggested by Colombo et al. (1970) for in situ coal bed methane in the Ruhr basin in Germany, which contains the most important German bituminous coal of Upper Carboniferous age and low volatile anthracite (Thomas, 2002). Progression in coal rank might explain the value of -50 ‰ for emissions from English underground mined bituminous coal and -30 ‰ for anthracite deep mines, followed by a 5-10 ‰ \(^{13}\)C-depletion likely caused by the incursion of meteoric water in the basin and the subsequent production of secondary biogenic methane (Scott et al., 1994), resulting in -40 ‰ for methane plumes from Welsh open-cut anthracite mines.

The link between coal rank and δ\(^{13}\)CCH\(_4\), isotopic signature is appreciable in the study of UK coal mines, but differences in the \(^{13}\)C/\(^{12}\)C isotopic ratio within the same coal sequences could be ascribed to other parameters, such as the depth at which the coal is mined - as methane migrates towards the surface, diffusion or microbial fractionation may occur, especially if there is water ingress (see section 1). In particular, emissions from Thoresby are more \(^{13}\)C depleted than those measured around Maltby, although both mines exploit the Parkgate seam (see Table 1), meaning that different isotopic signatures cannot be entirely linked to the coal rank. Biogenic methane produced in a later stage due to water intrusion might have been mixed with the original thermogenic methane formed during the coalification process.

Differences in methane emissions and their isotopic signature between opencast and deep mines have been assessed by surveying both surface and underground mines, in the Welsh anthracite belt, and in the Hunter Coalfield. The shallower deposits are more exposed to the weathering and meteoric water, most likely associated with the production of some isotopically lighter microbial methane. Mole fractions up to 2.5 ppm were measured around opencast mines in the Hunter Coalfield, in Australia, within a methane plume of more than 70 km width. The highest methane mole fractions were consistently measured downwind of vent shafts in underground mines.

The difference in the source isotopic signature for methane emissions between the two types of mining in the Hunter Coalfield (from -61 to -66 ‰) and in Wales (from -31 to -41 ‰) reflects the isotopic shift of 5-10 ‰ that might be attributed to the occurrence of secondary biogenic methane.

The δ\(^{13}\)C signatures for coalbed methane emissions from the Upper Silesian basin are highly variable, with the most \(^{13}\)C depleted methane associated with diffusion processes or secondary microbial methane generation (Kotarba and Rice, 2001), but the value of -51 ‰ measured for methane emissions from the KWK Wujek deep mine is consistent with the value of -50 ‰ inferred for emissions from English bituminous coal extracted in underground mines.
4 Conclusions

By measuring the isotopic signatures of methane plumes from a representative spread of coal types and depths, we show that the δ¹³C isotopic value to be included in regional and global atmospheric models for the estimate of methane emissions from the coal sector must be chosen according to coal rank and type of mining (opencast or underground). For low resolution methane modelling studies an averaged value of -65 ‰ is suggested for bituminous coal exploited in open cast mines and of -55 ‰ in deep mines, whereas values of -40 ‰ and -30 ‰ can be assigned to anthracite opencast and deep mines respectively.

Global methane budget models that incorporate isotopes have used a δ¹³C signature of -35 ‰ for coal or a value -40 ‰ for total fossil fuels (e.g. Hein et al., 1997; Mikaloff Fletcher et al., 2004; Bousquet et al., 2006; Monteil et al., 2011), but, given the relative rarity of anthracite global coal reserves and the dominance of bituminous coal (1% and 53% respectively) (World Coal Institute, 2015), it seems likely that a global average emission from coal mining activities will be lighter, with the -50 ‰ recorded for deep-mined bituminous coal in Europe being a closer estimate. However, for detailed global modelling of atmospheric methane, isotopic signatures of coal emissions should be region or nation specific, as greater detail is needed given the wide global variation. The assignment of an incorrect global mean, or a correct global mean but inappropriate for regional scale modelling, might lead to incorrect emissions estimates or source apportionment. The new scheme gives the possibility for an educated estimate of the δ¹³C signature of emission to atmosphere to be made for an individual coal basin or nation, given information on the type of coal being mined and the method of extraction.

In conclusion, high-precision measurements of δ¹³C in plumes of methane emitted to atmosphere from a range of coal mining activities have been used to constrain the isotopic range for specific ranks of coal and mine type, offering more representative isotopic signatures for use in methane budget assessment at regional and global scales.

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<td></td>
<td></td>
<td>Sept-2013</td>
<td>-48.8 ±0.3</td>
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<tr>
<td>Thoresby Colliery (active)</td>
<td>Nottinghamshire/UK</td>
<td>Parkgate seam, ~650 m in depth</td>
<td>Bituminous coal</td>
<td>Deep Mine</td>
<td>Nov-2013</td>
<td>-51.2 ±0.3</td>
<td>9</td>
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<tr>
<td>Daw Mill Colliery (closed since March 2013)</td>
<td>Warwickshire/UK</td>
<td>Warwickshire Thick seam, from 500 to 1000 m in depth.</td>
<td>Bituminous coal</td>
<td>Deep Mine</td>
<td>Nov-2013</td>
<td>-51.4 ±0.2</td>
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<tr>
<td>Cwmllynfell Colliery (active)</td>
<td>Wales/UK</td>
<td>South Wales Coalfield*</td>
<td>Anthracite</td>
<td>Surface Mine</td>
<td>Oct-2013</td>
<td>-41.2 ±0.9</td>
<td>4</td>
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<tr>
<td>Abercrave Colliery (active)</td>
<td>Wales/UK</td>
<td>South Wales Coalfield</td>
<td>Anthracite</td>
<td>Surface Mine</td>
<td>Oct-2013</td>
<td>-41.4 ±0.5</td>
<td>5</td>
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<tr>
<td>Aberpergwm Colliery (closed since December 2012)</td>
<td>Wales/UK</td>
<td>South Wales Coalfield</td>
<td>Anthracite</td>
<td>Deep Mine</td>
<td>Oct-2013</td>
<td>-33.3 ±1.8</td>
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<td></td>
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<tr>
<td>Unity Colliery (active)</td>
<td>Wales/UK</td>
<td>South Wales Coalfield</td>
<td>Anthracite</td>
<td>Deep Mine</td>
<td>Oct-2013</td>
<td>-30.9 ±1.4</td>
<td>3</td>
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<tr>
<td>Hunter Coalfield (active)</td>
<td>Australia</td>
<td>Whittingham Coal Measures</td>
<td>Bituminous coal</td>
<td>Surface and Deep Mines</td>
<td>Mar-2014</td>
<td>-66.4 ±1.3</td>
<td>12</td>
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<tr>
<td>Bulga Colliery (active)</td>
<td>Australia</td>
<td>Blakefield South Seam, from 130 to 510 m in depth (Bulga Underground Operation mining, 2015)</td>
<td>Bituminous coal</td>
<td>Deep Mine</td>
<td>Jan-2016</td>
<td>-60.8 ±0.3</td>
<td>10</td>
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<tr>
<td>Upper Silesian Basin (active)</td>
<td>Poland</td>
<td>502 (590-634m) and 510 seam (763-812m)</td>
<td>Bituminous coal</td>
<td>Deep Mine</td>
<td>Jun-2013</td>
<td>-50.9 ± 1.2</td>
<td>8</td>
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</tr>
</tbody>
</table>

Table 1 $\delta^{13}$CH$_4$ signatures of all the coal mines and coal basins surveyed with the Picarro mobile system. Errors in the $\delta^{13}$C signatures are calculated as 2 standard deviations. *Only the coalfield is indicated, as the name of the seams exploited could not be retrieved.
<table>
<thead>
<tr>
<th>Site</th>
<th>Coal Rank</th>
<th>$\delta^{13}$C (‰)</th>
<th>Author</th>
<th>$\delta^{13}$C (‰) measured in this study</th>
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</thead>
<tbody>
<tr>
<td>Powder River Basin, U.S.A.</td>
<td>Sub-bituminous coal</td>
<td>-79.5 to -56.8</td>
<td>Bates et al. (2011)</td>
<td></td>
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<tr>
<td>Australia</td>
<td>From brown coal to low volatile bituminous coal.</td>
<td>-73.0 to -43.5</td>
<td>Smith et al. (1981)</td>
<td>-66.4 ±1.3 to -55.5 ±1.3</td>
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<tr>
<td>Velenje Basin, Slovenia</td>
<td>Lignite</td>
<td>-71.8 to -43.4</td>
<td>Kanduč et al. (2015)</td>
<td></td>
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<tr>
<td>Powder River Basin, U.S.A.</td>
<td>Sub-bituminous coal</td>
<td>-68.4 to -59.5</td>
<td>Flores et al. (2008)</td>
<td></td>
</tr>
<tr>
<td>Illinois Basin, U.S.A.</td>
<td>Bituminous coal</td>
<td>-66.6 to -56.8‰</td>
<td>Strapoč et al. (2007)</td>
<td></td>
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<tr>
<td>Elk Valley Coalfield, Canada</td>
<td>Bituminous coal</td>
<td>-65.4 to -51.8</td>
<td>Aravena et al. (2003)</td>
<td></td>
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<tr>
<td>Bowen basin, Australia</td>
<td>Bituminous coal</td>
<td>-62.2 to -48</td>
<td>Kinnon et al. (2010)</td>
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<tr>
<td>Queensland Basin, Australia</td>
<td>From Sub-bituminous to high volatile bituminous coal</td>
<td>-57.3 to -54.2</td>
<td>Papendick et al. (2011)</td>
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<tr>
<td>Tara region, Queensland Basin, Australia</td>
<td>From Sub-bituminous to high volatile bituminous coal</td>
<td>-56.7</td>
<td>Maher et al. (2014)</td>
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<tr>
<td>San Juan basin, New Mexico and Colorado</td>
<td>High-volatile Bituminous coal</td>
<td>-43.6 to -40.5</td>
<td>Rice et al. (1989)</td>
<td></td>
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<td>UK, Barnsley seam</td>
<td>Bituminous coal</td>
<td>-44.1</td>
<td>Hitchman et al. (a, b)</td>
<td>-48.8 ±0.3 to -45.4 ±0.2</td>
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<tr>
<td>Upper Silesian Coal Basin, Poland</td>
<td>Sub-bituminous coal to anthracite</td>
<td>-79.9 to -44.5</td>
<td>Kotarba and Rice (2001)</td>
<td>-50.9 ±0.6</td>
</tr>
<tr>
<td>Eastern China</td>
<td>Sub-bituminous to anthracite</td>
<td>-66.9 to -24.9</td>
<td>Dai et al. (1987)</td>
<td></td>
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<tr>
<td>Ruhr basin, Germany</td>
<td>Bituminous coal, Anthracite</td>
<td>-37 (-60 to -14)</td>
<td>Deines (1980)</td>
<td></td>
</tr>
<tr>
<td>Western Germany</td>
<td>High-volatile bituminous to anthracite</td>
<td>-70.4 to -16.8</td>
<td>Colombo et al. (1970)</td>
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<tr>
<td>Eastern Ordos Basin, China</td>
<td>Bituminous Coal</td>
<td>-42.3 to -39.8</td>
<td>Guo et al. (2012)</td>
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<td>Qinshui, China</td>
<td>Anthracite</td>
<td>-41.4 to -34.0</td>
<td>Qin et al. (2006)</td>
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<td>Qinshui Basin, China</td>
<td>Semi-anthracite to Anthracite</td>
<td>-36.7 to -26.6</td>
<td>Su et al. (2005)</td>
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<tr>
<td>Sichuan Basin, China</td>
<td>Bituminous to anthracite</td>
<td>-36.9 to -30.8</td>
<td>Dai et al. (2012)</td>
<td></td>
</tr>
<tr>
<td>Wales, UK</td>
<td>Anthracite</td>
<td></td>
<td>Xiong et al. (2004)</td>
<td>-33.3 ±1.8 to -30.9 ±1.4</td>
</tr>
</tbody>
</table>

Table 2 Literature isotope values obtained by methane samples CBM (Coalbed Methane) wells across the basin. Errors are not included in the sources.
Figure 1 Coal maturation process.

Figure 2 Methane mole fractions recorded in the Hunter Coalfield on 12th March 2014 (a) and during a more detailed survey of the area highlighted by the black square on 18th March 2014 (b). Source: Esri, DigitalGlobe, GeoEye, i-cubed, Earthstar Geographics, CNES/Airbus DS, USDA, USGS, AEX, Getmapping, Aerogrid, IGN, IGP, swisstopo and the GIS User Community.
Figure 3 Keeling Plots based on samples collected around English and Welsh coal mine (a), coal mines in Poland (b) and Australia (c). Errors on the y-axis are within 0.05 ‰ and on the x-axis 0.0001 ppm⁻¹, and are not noticeable on the graph.