ELSEVIER

Contents lists available at ScienceDirect

Applied Geochemistry

journal homepage: www.elsevier.com/locate/apgeochem



Carbonate precipitation in artificial soils as a sink for atmospheric carbon dioxide

P. Renforth a,b,*, D.A.C. Manning a,b, E. Lopez-Capel b,c

ARTICLE INFO

Article history: Received 26 February 2009 Accepted 10 May 2009 Available online 18 May 2009

Editorial handling by R. Fuge

ABSTRACT

Turnover of C in soils is the dominant flux in the global C cycle and is responsible for transporting 20 times the quantity of anthropogenic emissions each year. This paper investigates the potential for soils to be modified with Ca-rich materials (e.g. demolition waste or basic slag) to capture some of the transferred C as geologically stable CaCO₃. To test this principal, artificial soil known to contain Ca-rich minerals (Ca silicates and portlandite) was analysed from two sites across NE England, UK. The results demonstrate an average C content of $30 \pm 15.3 \, \text{Kg C} \, \text{m}^{-2}$ stored as CaCO₃, which is three times the expected organic C content and that it has accumulated at a rate of $25 \pm 12.8 \, \text{t C} \, \text{ha}^{-1} \, \text{a}^{-1}$ since 1996. Isotopic analysis of the carbonates gave values between -6.4% and -27.5% for δ^{13} C and -3.92% and -20.89% for δ^{18} O, respectively (against V-PDB), which suggests that a combination of carbonate formation mechanisms are operating including the hydroxylation of gaseous CO₂ in solution, and the sequestration of degraded organic C with minor remobilisation/precipitation of lithogenic carbonates. This study implies that construction/development sites may be designed with a C capture function to sequester atmospheric C into the soil matrix with a maximum global potential of 290 Mt C a⁻¹.

© 2009 Elsevier Ltd. All rights reserved.

1. Introduction

The global C cycle transports approximately 210 Gt of C per year between a multitude of pools including the biosphere, geological sediment, the ocean and the atmosphere. During this cycle, 60% of all the C flux passes between the atmosphere and the terrestrial system (120 Gt C a⁻¹; Dupre et al., 2003). Thus within the context of the possible use of geoengineering procedures to compensate for artificial CO_2 emissions (e.g. Lal, 2003), it is highly appropriate to consider the role of the coupled plant–soil system in C capture, and to develop ways of enhancing natural processes artificially.

In temperate climates, the role of soils as C sinks is often associated with the accumulation of soil organic C (SOC) in agricultural soils (Smith, 2004). Documentation of soil inorganic C (SIC) is usually confined to soils formed in arid climates, where unbroken 'calcrete' structures can cover an area of several km². Schlesinger (1985) suggests that soils of this type may contain between 24.5 and 33 kg C m $^{-2}$.

Recent work has demonstrated substantial variability of organic C content in urban soils which can be as high as 28.5 kg C m^{-2} , although the average C content is expected to be between 8 and 10 kg C m^{-2} (Pouyat et al., 2002, 2006, Banaitis et al., 2007). Pouyat

E-mail address: Philip.Renforth@ncl.ac.uk (P. Renforth).

et al. (2002) have demonstrated the influence of landuse on SOC and suggest that soils beneath impervious surfaces or in clean engineered fill material will return the lowest values for organic C. Moreover, the turnover of C in urban soils and the characterisation of the C pool (labile to refractory) have yet to be determined. By investigating the formation of inorganic C (SIC) as carbonate minerals, the aim of this study is to enhance the understanding of the C cycle in urban soils.

Soils at two brownfield sites in NE England, UK were investigated for the accumulation of SIC as geologically stable CaCO₃. Brownfield sites are commonly but not exclusively characterised by the presence of waste material arising from the historical use of the site, a proportion of which becomes part of the soil matrix. Some of this is derived from construction materials, including Ca-rich components (artificial mortars, plaster, concrete, natural basic rock aggregates, slag (e.g. Fredericci et al., 2000). It is hypothesised that weathering of Ca-rich minerals (silicates, hydroxides, sulphates) within these materials will result in precipitation of CaCO₃ within soils through Eq. (1) by reaction with C cycled through plant roots (Manning, 2008).

$$Ca^{2+} + 2HCO_3^- \leftrightarrow CaCO_3 + H_2O + CO_2 \tag{1}$$

The availability of Ca depends on the stability of the Ca minerals in the soil system. The rate at which these materials weather and release Ca is dependant on various factors including mineralogy, physical grain properties (including surface area), solution pH,

^a Institute for Research on Environment and Sustainability, Newcastle University, Newcastle upon Tyne, NE1 7RU, United Kingdom

^b School of Civil Engineering and Geosciences, Newcastle University, Newcastle upon Tyne, NE1 7RU, United Kingdom

^cJoseph Swan Institute for Energy Research, Newcastle University, Newcastle upon Tyne, NE1 7RU, United Kingdom

^{*} Corresponding author. Address: Institute for Research on Environment and Sustainability, Newcastle University, Newcastle upon Tyne, NE1 7RU, United Kingdom. Tel.: +44 0 191 246 4894.

the presence of organic complexes and flow rate, but is estimated to vary between 10^{-15} and 10^{-6} mol cm⁻² s⁻¹ (Blum and Stillings, 1995; Berg and Banwart, 2000; van Hees et al., 2002).

Carbon is transferred into the soil through dissolution in rainwater or through biological processes. During growth, plants exude large quantities of C through their roots as organic compounds which ultimately degrade to CO₂ and return to the atmosphere (Kuzyakov and Domanski, 2000; Ryan et al., 2001). This is the dominant conveyor in the global C cycle, and transports approximately 120 Gt C a⁻¹ compared to 6 Gt C a⁻¹ produced by anthropogenic sources (Lal, 2003). The rate at which plants turn over C is difficult to measure, but Manning (2008) suggests between 1.2 and 16.1 mg C g⁻¹ of fresh weight per year is exuded from plant roots based on in-vitro experiments (Ryan et al., 2001). However, Moulton et al. (2000) have monitored the carbonate concentration of waters issuing from a groundwater system in Iceland, showing that plant-derived HCO₃ concentrations of waters draining forested areas were between 911 and 999 mol ha⁻¹ a⁻¹, or approximately 500 mol kg⁻¹ a⁻¹ when normalised against biomass per ha (Manning, 2008). This is an order of magnitude larger than the exudation rates calculated from the laboratory experiments.

Carbon dating of pedogenic (soil formed) carbonates indicates long residence times in soils (>30 ka; Kalin et al., 1997) and field studies of carbonates in ancient soils similarly support the stable nature of carbonates (up to 2.6 Ga; e.g. Watanabe et al., 2004). Furthermore, Kuzyakov et al. (2006) demonstrated, in controlled conditions, the dissolution and reprecipitation of carbonate minerals under the influence of plant roots. They concluded that a 100% turnover of the carbonate can take between 0.4 and 2 ka. Both the field and laboratory scale investigations have demonstrated the stable nature of pedogenic carbonates on human time scales.

The aim of the research reported here is to investigate the extent to which soils modified by the artificial introduction of Ca minerals can capture C exuded from plants to give a semi-permanent sink, and the significance of this process as compensation for artificial C emissions.

2. Study sites

An urban brownfield site was chosen in Newcastle upon Tyne approximately 2.5 km east of the city centre (GB National Grid: NZ275649 – see insert of Fig. 1) which was previously occupied

by a concrete office complex that was demolished in 1996. The site has remained unused since then, apart from storage of a soil heap excavated from an adjacent development site (approximate location shown in Fig. 1). A contaminated land report completed by Newcastle City Council in 1998 presented soil profiles that show the presence of demolition waste throughout the site to 2–3 m in depth. Furthermore, the study found pH levels of up to 11.8 and a Sulphate content of up to 20313 mg kg⁻¹ towards the north of the site indicating portlandite (Ca(OH)₂) and gypsum (CaSO₄·2H₂O) dissolution, respectively. Vegetation on the site comprised of C₃ grasses (including species typical of a restoration seed mix) and C₄ ornamental garden escapes. The bedrock geology of the area is dominated by Carboniferous sandstone which is superficially overlain by glacial till. The site has been heavily modified by industrial activity and a substantial thickness (>3 m) of made ground is present.

A second site was selected adjacent to a former steel works in Consett (GB National Grid: NZ094492), County Durham, England which was closed in the 1980s creating 290 ha of derelict land. In a review of the contamination problems typically associated with iron and steel making, Harber and Forth (2001) describe remodelling of the ground profile at the Consett site with the emplacement of 'clean slag'. At the site, slags of different types have accumulated as steel making processes evolved since the early 1800s. The basic slag used for the cover is a lime-rich Ca silicate glass, containing Ca silicate minerals (such as merwinite, melilite and larnite; Fredericci et al., 2000), Ca oxide as Ca ferrite and portlandite as well as Fe oxides (Harber and Forth, 2001). The final stage in restoration involves emplacement of top soil, from which the samples were taken. Subsequent analysis of the area (Mayes et al., 2006) has discovered pH levels of 12.5 within leachate draining the site which was attributed to the weathering of portlandite and Ca silicate minerals. The waters were supersaturated with respect to calcite and showed a relationship between precipitation and biological activity, coupled with attenuation of pH through a pond and calcareous wetland which has developed on a watercourse draining the former steel works area. A calcareous hardpan has formed at the surface adjacent to the pond and covers an area of approximately 400 m². The bedrock geology of the area is dominated by the Carboniferous mudstones of the Lower Coal Measures, but the landscape has been considerably altered by industrial activity.

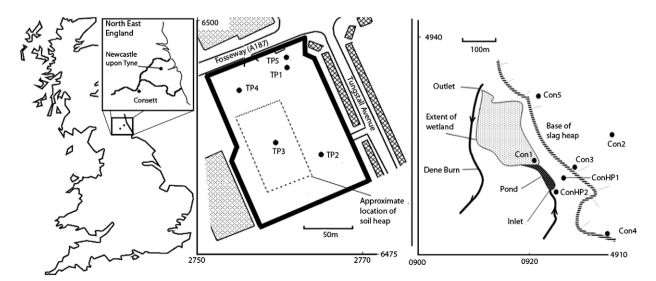


Fig. 1. Sampling locations (●) at an urban brownfield site in Newcastle upon Tyne (left) and a former steelworks in Consett (right) (Grid based on the Ordinance Survey National Grid of Great Britain – tile NZ).

3. Methodology

At the brownfield site in Newcastle upon Tyne, Ian Farmer Associates (www.ianfarmerassociates.co.uk) were contracted to excavate five trial pits to 3.5 m. At the former steelworks at Consett, soil samples were collected with a hand auger to depths up to 20 cm prior to contact with large fragments of slag. Furthermore, sediment was collected from the bottom of the high pH pond (locations are shown in Fig. 1). Samples were collected through the soil profile at both sites, air-dried and sieved to <2 mm. Calcium carbonate content was determined using an Eijkelkamp calcimeter to BS 7755-3.10:1995 (reproducibility better than ±0.6%), the residue was collected, washed, filtered and oven dried at 80 °C and used for the determination of organic C isotope ratios. Soil pH was analysed according to ISO 10390 1994, using a Jenway 3020 pH meter.

Isotope ratio mass spectrometry was conducted using a Europa Scientific 20-20 IRMS (Iso-Analytical, Cheshire UK) for $^{13}\text{C}/^{12}\text{C}$ and $^{18}\text{O}/^{16}\text{O}$ for CaCO $_3$ and $^{13}\text{C}/^{12}\text{C}$ for organic C collected from the calcimeter residues. IA-R022, NBS-18, NBS-19 were used twice as reference materials for carbonate (C and O) isotope analysis and IA-R001, IA-R005 and IA-R006 were used for organic C isotope analysis. Replication had a standard deviation better than $\pm 0.1\%$ and all but one of the six reference materials were within two standard deviations of the expected results (<0.2% difference for NBS-18; two separate analysis batches). The results were recorded relative to the Vienna Peedee Belemnite scale (V-PDB).

4. Results

4.1. Urban brownfield, Newcastle upon Tyne

In some trial pits, visible precipitates of CaCO $_3$ occurred as coatings and concretions on demolition rubble. Chemical analysis determined CaCO $_3$ quantities within the soil to be between 2.2 and 19.9 wt% ($\bar{x}=9.5\%$ s = 5.1). Trial pits TP1, TP4 and TP5 showed negative correlation of CaCO $_3$ with depth ($r^2=0.78$, 0.99 and 0.50, respectively, after removal of outliers from the basal clay in TP1, for which a carbonate content of 4.3% was measured compared with values between 11% and 20% returned for other samples from the same trial pit, and unconsolidated sand in TP5, which had a similar lower value than the other samples from the same pit). There was no variation in carbonate content with depth in TP2 and TP3 (see Fig. 2 – note error bars are within the data point symbols).

Carbon and O isotope analysis of the carbonates gave δ^{13} C values between -6.3% and -27.5% and δ^{18} O between -3.9% and

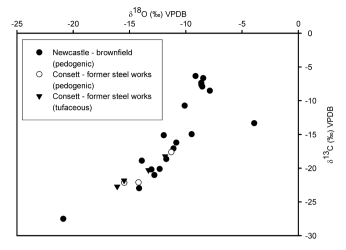


Fig. 3. Isotopic ratios from carbonates formed in an urban brownfield soil (\bullet) and within the drainage regime of the former steelworks (\bigcirc) .

-20.9% (Fig. 3). The most negative carbon isotope values (between -13.3% and -27.5%) were detected in precipitates on demolition rubble. There is a positive correlation between inorganic O and C isotope values ($r^2 = 0.66$ or 0.86 after the removal of outliers). Analysis of the organic C isotope ratios gave values between -19.0% and -24.0% δ^{13} C (all but one value lies within 1 standard deviation of the mean of 22.9%).

Comparison of pH values recorded in this study with those presented in the contaminated land report suggest a decline in soil pH over time (from 12 to 7 at the present day). However, a comparison is speculative due to variation in sampling locations.

4.2. Former steelworks, Consett

Calcium carbonate occurred in both the soil (pedogenic carbonate) and within the drainage system of the site (tufaceous carbonate). Quantities of CaCO₃ ranged between 0% and 93.4% ($\overline{x}=29.6\%$ s = 36.6) over both systems. The largest concentration of CaCO₃ was recorded in the sediment of the pond (>90%) and the hardpan (48–87%). The concentration of pedogenic carbonate was extremely variable (between 0% and 38.3% $-\overline{x}=7.0\%$ s = 14.0) and there appears to be no relationship between concentration and depth (see Fig. 2). Carbonate isotope ratios were between -17.6% and -22%. 7% for δ^{13} C and -11.3% and -16.1% for δ^{18} O (see Table 1 and Fig. 3).

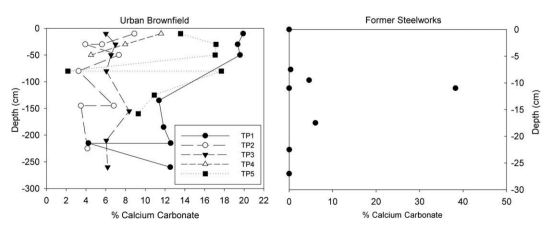


Fig. 2. Carbonate concentrations with depth at both sites.

Table 1Isotopic values and sample depth for pedogenic carbonates at both sites.

Sample no.	Sample depth (cm)	δ^{13} C	δ^{18} O
Urban brownfield (NZ275649)		
A2TP1 1	0–20	-7.7	-8.6
A2TP1 2	40-60	-20.2	-13.1
A2TP1 3	200-230	-18.6	-11.7
A2TP2 1	0–20	-15.1	-12.0
A2TP2 2	40-60	-10.7	-10.1
A2TP2 3	200-250	-7.9	-8.5
A2TP3 1	0–20	-6.7	-8.5
A2TP3 2	40-60	-6.4	-9.1
A2TP3 3	190-230	-7.4	-8.6
A2TP4 1	0–20	-8.5	-7.9
A2TP4 2	40-60	-15.0	-9.5
A2TP5 1	0–20	-16.2	-10.8
A2TP5 2	40-60	-20.1	-12.3
A2TP5 3	150–170	-17.1	-11.1
A2TP1 RS 1	100-170	-27.5	-20.9
A2TP1 RS 2	170–200	-23.0	-14.2
A2TP2 RS 1	200-250	-21.0	-12.8
A2TP2 RS 2	100-190	-13.3	-3.9
A2TP5 RS 1	40-60	-18.9	-13.9
Former steelworks	(NZ094492)		
CON 01 1	0-35	-22.1	-14.2
CON 01 2	Surface	-22.7	-16.1
CON 04 1	0–19	-17.6	-11.3
CON HP1 1	0–8	-22.1	-15.5
CON HP1 2	8–15	-18.2	-11.8
CON HP2 1	>15	-20.3	-13.3
CON HP2 2	0–15	-21.8	-15.5

5. Discussion

5.1. $\delta^{13}C$ and $\delta^{18}O$ stable isotopes of carbonates

Stable isotope analysis has been used to study carbonates since the 1950s (Craig, 1953) primarily to investigate diagenesis conditions in limestone and the influence of organic C during the remobilisation of CaCO3 in soils (Hudson, 1977). Early work by Salomons and Mook (1976) and Cerling (1984) demonstrate the incorporation of organic C in pedogenic carbonates which characteristically have isotopic values between -2% and -10% for δ^{13} C and 0% and -5% for δ^{18} O, respectively. Similar values have been found in more recent studies from a range of environments. It is clear that carbonate isotope values are controlled by precipitation conditions, including climate, rainfall, temperature, underlying geology and continentality (Andrews, 2006). The data obtained for the artificial soils investigated here are compared with published data from both natural and artificial soil in Figs. 4 and 5.

In Fig. 4, Region A represents the precipitation of carbonates in 'natural' sedimentary environments. In Fig. 5, the expected range of δ^{13} C values is shown for carbonates formed under the influence of C_3 vegetation. Measurements from several studies fall within this range (Salomons and Mook, 1976; Cerling, 1984; Zanchetta et al., 2000; Bajnoczi et al., 2005; Kovda et al., 2006; Boguckyj et al., 2006; Wang and Greenberg, 2007; Singh et al., 2007 and Łącka et al., 2008). According to Andrews (2006) there is a general decrease in δ^{18} O with increasing continentality (from δ^{18} O = -4%0 to -14%0). Many authors (Boguckyj et al., 2006; Singh et al., 2007; Wang and Greenberg, 2007; Łącka et al., 2008) record the occurrence of rhizoliths, which are carbonates formed in close proximity to plant roots and are dominated by organic C.

The provenance of C within pedogenic carbonates is usually a combination of that which is derived from organic C, and remobilisation of lithogenic carbonates. Pedogenic carbonates formed under these influences will return an isotopic signature which is between the organically-dominated region and the lithogenically-dominated region contiguous with $\delta^{13}C = 0\%$ (Hudson, 1977).

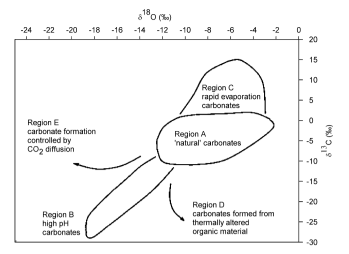


Fig. 4. δ^{13} C and δ^{18} O isotope values returned from carbonates formed in a range of environments. Region A depicts carbonates formed in 'natural' conditions (Salomons and Mook, 1976; Cerling, 1984; Liu et al., 1996; Zanchetta et al., 2000; Knauth et al., 2003; Piovano et al., 2004; Bajnoczi et al., 2005; Boguckyj et al., 2006; Kovda et al., 2006; Singh et al., 2007; Wang and Greenberg, 2007; Sikes and Ashley, 2007; Łącka et al., 2008; Yanes et al., 2008). Region B depicts the range of values returned from carbonates in high pH environments (Macleod et al., 1991; Dietzel et al., 1992; Andrews et al., 1997; Krishnamurthy et al., 2003; Boguckyj et al., 2006; Fléhoc et al., 2006). Region C depicts values returned from evaporation dominated environments (Achyuthan et al., 2007; Knauth et al., 2003). Region D includes values from carbonates which have been formed under the influence of thermally modified organic carbon (Ohlsson, 2000; Fourcade et al., 2007). Region E depicts carbonates formed under closed/semi closed systems where the diffusion of CO₂ becomes rate limiting (van Strydonck et al., 1989 and Kosednar-Legenstein et al., 2008).

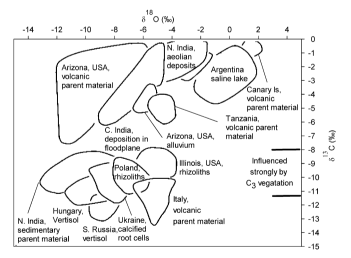


Fig. 5. Detailed breakdown of isotope values in Region A.

However, the use of isotope ratio data to determine geological source is potentially ambiguous, as demonstrated by Zanchetta et al. (2000), Sikes and Ashley (2007) and Yanes et al. (2008), who have independently found differing values for carbonates formed in soils on igneous parent rocks.

Several studies have published isotopic data for carbonates formed in anthropogenic environments (Region B in Fig. 4). van Strydonck et al. (1989), Macleod et al. (1991), Dietzel et al. (1992), Krishnamurthy et al. (2003) and Kosednar-Legenstein et al. (2008) have investigated the formation of carbonates in alkaline environments associated with concrete and attribute the observed negative isotopic signatures to kinetic fractionation when

 ${\rm CO_2}$ gas is dissolved in solution, which rapidly reacts with OH $^-$ ions (from portlandite dissolution) to form carbonate through Eq. (2).

$$OH^- + CO_2 \leftrightarrow CO_3^- + H^+ \tag{2}$$

The rate of this is governed by (Dietzel et al., 1992):

$$r = C_o \cdot (D \cdot k \cdot [OH^-])^{0.5} \tag{3}$$

where k is the rate constant (cm³ mol⁻¹ s⁻¹) for hydroxylation, D is the diffusion coefficient of CO_2 through the liquid (cm² s⁻¹) and C_0 the CO₂ concentration in solution (mol cm⁻³). Assuming typical values for k and D of $10^{-4.83}$ cm² s⁻¹ and $10^{6.41}$ cm³ mol⁻¹ s⁻¹, respectively, it can be seen that the primary rate controlling step is the reaction between hydroxide and dissolved CO2 gas. Dietzel et al. (1992) have associated this phenomenon with an isotope fractionation of -18.8% for δ^{13} C, and a similar fractionation is experienced by O isotopes. Further to this, Andrews et al. (1997), Boguckyj et al. (2006) and Fléhoc et al. (2006) have published data which conform to this model. Collectively, the isotopic signatures observed in this study fall within a tight regression ($r^2 = 91.8\%$) against the line δ^{13} C = 1.6 δ^{18} O + 2.8, which corresponds to carbonates formed in a combination of natural and high pH conditions. Indeed, the position on this line in comparison to the two approximate end members $(\delta^{18}O = -6.75\%, \delta^{13}C = -8\%)$ for organically derived C (Cerling, 1984) and $(\delta^{18}O = -17.6\%, \delta^{13}C = -25.3\%)$ for high pH carbonate (Dietzel et al., 1992), can be used to estimate the proportion of C derived organically and that which has been sequestered through hydroxylation.

Results from the isotopic analysis carried out in this study suggest that three mechanisms of carbonate formation (lithogenic remobilisation, hydroxylation of gaseous CO₂ and organic C sequestration) were operating in the urban brownfield site and two (hydroxylation and organic C sequestration) were operating at the former steelworks. The relative proportions of each process

can be deduced from proximity to each end member, as described previously, and is summarised in Fig. 6. The recorded isotope values at both sites comply with the trend observed for other published data, and there is no statistical difference between slopes (p < 0.05) in Figs. 3 and 4. Remobilisation of lithogenic carbonate was assumed to be negligible in high pH waters.

5.2. Carbonate precipitation in soil modified with demolition rubble

The analysis of the urban brownfield site suggests that an average of 56.8% of the carbonate C is of organic origin, 40.5% is derived from hydroxylation and 2.7% from lithogenic carbonate. Extending these figures spatially for 10% average carbonate accumulation to 2.5 m depth and assuming 1 t m⁻³ density of crushed concrete (Dhir et al., 1999), it can be theorised that this particular urban brownfield site has a SIC content of $30 \pm 15.3 \text{ kg C m}^{-2}$, 97.3% of which has been sequestered ultimately from the atmosphere. The SIC content is three times greater than the value reported for the average organic C content in urban areas (Pouvat et al., 2006). Extrapolating these figures through the life of the brownfield site, it can be estimated that the soil has accumulated C at a rate of approximately $25 \pm 12.8 \text{ t C ha}^{-1} \text{ a}^{-1}$. It can be speculated that the accumulation of carbonate is accordingly matched by a decrease in soil pH. Indeed, the pH levels recorded in both this study and the contaminated land report completed in 1998 were not sufficiently high to cause hydroxylation. Therefore, it can be hypothesised that the accumulation of hydroxylated carbonate was rapid (within 2 a) and the subsequent carbonate accumulation is the result of Ca silicate weathering and organic C sequestration. However, the clast-supported nature of the soil suggests that the site drains freely; therefore, it is possible that the micro-environment around individual soil grains experience high pH levels.

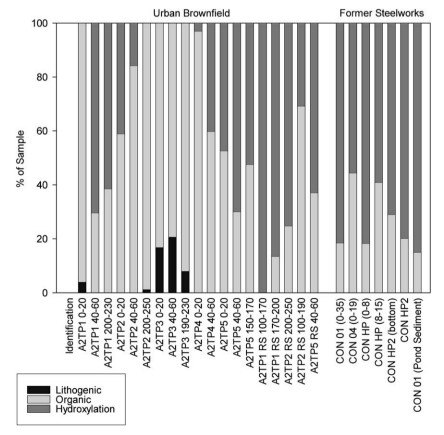


Fig. 6. Proportions of carbonate precipitation mechanisms active at both study sites.

5.3. Carbonate precipitation in soil modified with basic slag

Similar analysis of the carbonate C formed within soil modified with basic slag suggests that 31.4% of the carbonate C is derived from organic C and 68.6% is the product of hydroxylation of soil CO₂. Carbonate concentration is heterogeneous throughout the site and is clearly dependent on the drainage regime, with high concentrations at the pond/wetland and lower concentration at the brow of the slope. High pH levels are still present within the drainage waters on site almost 30 a after the steelworks were closed, which suggests sustained weathering of portlandite.

The rate of carbonate formation within the aqueous regime at the former steelworks was calculated by Mayes et al. (2006) using the Dreybrodt limestone tablet method. They found that precipitation rates decrease across the flow path of the wetland between 0.6 and 1.8 t C ha $^{-1}$ a $^{-1}$ over an area of 1500 m 2 . The high variability of soil carbonate concentration means that it is not possible to carry out spatial accumulation calculations.

5.4. Implications for carbon capture and geoengineering

Brownfield sites are ubiquitous in the UK as a consequence of the country's industrial heritage, and cover approximately 42,000 ha of land (see Table 2 for details). Extrapolating 30 kg C m⁻² for carbonate concentration found in this study, it can be estimated that the UK stores approximately 12.7 Mt C as carbonate in brownfield soil. It is interesting to speculate the C capture potential of urban soils if they were designed for that purpose. For example, the 625 ha occupied by London Olympics 2012 by the authors' calculations could sequester 180,000 t C. The formation of CaCO₃ within soils by reaction with construction materials is one way to compensate for the production of CO2 during cement manufacture. Given the importance of concrete in construction (an industry which contributes 8% of the UK economy; DBERR, 2008), a full understanding of the possible value of concrete as a C sink at the end of its life allows the full life cycle impact of C emissions associated with construction to be understood in the context of sustainably mitigating climate change.

The maximum capacity of inorganic C capture technology is limited by the availability of Ca-rich minerals. DCLG (2007b) suggests approximately 88.6 Mt of construction and demolition waste is produced annually in the UK and 46.5 Mt of this is currently

landfilled or spread on demolition sites. Assuming a CaO content of 20%, the maximum potential of carbonate capture is estimated to be 2 Mt C $\rm a^{-1}$. These figures can be extrapolated to the global 'geoengineering' scale, under the assumption of a global production to waste ratio for concrete similar to the UK, and suggest that the upper limit for C capture using this technology is approximately 290 Mt C $\rm a^{-1}$ which is equivalent to 90% of the emissions associated with cement manufacture (Hendriks et al., 2004).

6. Conclusions

It is believed that this is the first reported case that demonstrates how artificial soils (i.e. made ground) can act as C sinks by accumulating CaCO₃. Precipitation of CaCO₃ was found to be associated with the weathering of portlandite at both sites, but it is unknown to what extent other Ca-rich minerals (Ca silicates, gypsum) are responsible for the formation of CaCO₃. The capacity for the OH⁻ ion to buffer pH in soil at the urban brownfield site has diminished over time as a result of hydroxylation of gaseous CO2, biological activity and site hydrology, which facilitated a decrease in soil pH. It is possible that a similar pH regime is progressing at the former steelworks site but a larger initial concentration of portlandite has sustained the process. With this in mind, the prevalence of hydroxylation on site will be replaced by the incorporation of biologically influenced carbonates as the weathered cations from silicates exert greater influence on the precipitation of CaCO₃. To that extent, further research should be undertaken to investigate the weathering and contribution of silicate minerals on carbonate formation. It is also important to note that the study area at the former steelworks is only a small fraction of the original site and it is impossible to gauge the efficiency of C capture without additional work to quantify the extent of the Ca-rich material using extensive groundwater/hydrological analysis and soil profile analysis.

Brownfield soils are extremely heterogeneous and additional studies are required to investigate how other artificial soils accumulate C. The results of this study suggest that demolition waste recycled into the soil will sequester C at a rate of $25 \pm 12.8 \text{ t C ha}^{-1} \text{ a}^{-1}$. However, additional work is required to investigate carbonate dynamics and stability over time. Furthermore, investigations are needed to ascertain the C capture potential of soils specifically engineered for that purpose.

Table 2UK national and global limits of inorganic carbon capture technology.

Brownfield land and concrete production in the UK and globally				
JK figures				
Jrban settlement area	1,286,000 ha ^a	General Register Office for Scotland (2000), The Northern Ireland Statistics and Research Agency (NISRA) (2001) and DCLG (2008)		
Brownfield/vacant land	42,200 ha ^b	(Scottish Executive – Statistical Bulletin, 2002, DCLG, 2007a)		
Carbon currently stored in brownfield sites	12.7 MtC			
Cement production	15.7 Mt	British Cement Association (2009) - 2007 figures		
Estimated concrete production	110 Mt ^c			
Concrete waste production (not recycled)	46.5 Mt	DCLG (2007b)		
Maximum capture potential	2 Mt C y ⁻¹			
Global figures				
Jrban settlement coverage	44,237,400 ha	Demographia (2008)		
Brownfield/vacant land	1,451,000 ha ^d			
Estimated current storage	435.3 Mt C			
Cement production	2310 Mt	USGS (2004)		
Stimated concrete production	16,200 Mt ^c			
stimated concrete waste production	6800 Mt ^e			
Maximum capture potential	290Mt C y^{-1}			

- ^a Statistics for England and Wales amended with data for Glasgow, Edinburgh, Dundee, Motherwell and Belfast.
- b Brownfield contribution estimated based on relative proportions of urban area to brownfield in England, Wales and Scotland.
- ^c Based on a concrete mix of 1:2:4 w/w of cement, sand and aggregate.
- ^d Calculated by extrapolating the ratio of brownfield to urban land in Scotland, England and Wales for the total global urban area.
- ^e Estimated using the UK concrete production to waste ratio.

Decoupling economic development from the production of greenhouse gases is the most important step in sustainably mitigating climate change. Economically developed countries support a strong construction sector which is a direct physical manifestation of development. One of the primary materials of construction is cementitious products (i.e. concrete), which is produced by calcining CaCO₃ and is responsible for 8% of the world's CO₂ emissions (Wilson, 1993) and steel which is responsible for approximately 5% (von Scheele, 2006). Reforming CaCO₃ in soils would partially close the loop on the C footprint from steel and cement production and decouple a substantial part of the construction industry from greenhouse gas emission.

Acknowledgements

This work was funded by ESPRC (EP/F02777X/1) and NERC (studentship NE/F008716/1). The authors thank Philip Hartley and Newcastle City Council for providing information and access to one of the field sites. Dr. Simon Peacock and Dr. William Mayes at Newcastle University should also be thanked for their help during fieldwork. Comments from Professor Julian Andrews and an anonymous reviewer were appreciated.

References

- Achyuthan, H., Quade, J., Roe, L., Placzek, C., 2007. Stable isotopic composition of pedogenic carbonates from the eastern margin of the Thar Desert, Rajasthan, India. Quatern. Int. 162 (163), 50–60.
- Andrews, J.E., 2006. Palaeoclimatic records from stable isotopes in riverine tufas: synthesis and review. Earth-Sci. Rev. 75, 85–104.
- Andrews, J.E., Gare, S.G., Dennis, P.F., 1997. Unusual isotopic phenomena in Welsh quarry water and carbonate crusts. Terra Nova 9, 67–70.
- Bajnoczi, B., Horvath, Z., Demeny, A., Mindszenty, A., 2005. Stable isotope geochemistry of calcrete nodules and septarian concretions in a Quaternary 'red clay' paleovertisol from Hungary. In: 8th Isotope Workshop of the European-Society-for-Isotope-Research, Leipzig, Germany, June 25–30. Taylor & Francis Ltd.
- Banaitis, M.R., Langley-Turnbaugh, S.J., Aboueissa, A., 2007. Variations of soil organic carbon in three urban parks: a Maine case study. Int. J. Appl. Environ. Sci. 2, 119–128.
- Berg, A., Banwart, S.A., 2000. Carbon dioxide mediated dissolution of Ca-feldspar: implications for silicate weathering. Chem. Geol. 163, 25–42.
- Blum, A.E., Stillings, L.L., 1995. Feldspar dissolution kinetics. In: White, A.F., Brantley, S.L. (Eds.), Chemical Weathering Rates of Silicate Minerals. Reviews in Mineralogy, vol. 31. Mineralogical Society of America, pp. 291–351.
- Boguckyj, A.B., Lanczont, M., Lacka, B., Madeyska, T., Zawidzki, P., 2006. Stable isotopic composition of carbonates in Quaternary sediments of the Skala Podil'ska sequence (Ukraine). Quatern. Int. 152 (153), 3–13.
- British Cement Association, 2009. Table 2. Quarterly Cementitious BCA Web Statistics.
- Cerling, T.E., 1984. The stable isotopic composition of modern soil carbonate and its relationship to climate. Earth Planet. Sci. Lett. 71, 229–240.
- Craig, H., 1953. The geochemistry of the stable carbon isotopes. Geochim. Cosmochim. Acta 3, 53–92.
- DBERR, 2008. Strategy for Sustainable Construction. Crown Copyright, UK.
- DCLG, 2007a. Previously-Developed Land That may be Available for Development: England 2006. Crown Copyright (07HC04713), UK.
- DCLG, 2007b. Survey of Arisings and Use of Alternatives to Primary Aggregates in England, 2005 Construction, Demolition and Excavation Waste. Crown Copyright, UK.
- DCLG, 2008. Urban Settlement 2001: England and Wales.
- Demographia, 2008. Demographia World Urban Areas (World Agglomerations): 2008.
- Dhir, R.K., Limbachiya, M.C., Leelawat, T., 1999. Suitability of recycled concrete aggregate for use in BS 5328 designated mixes. Proc. Inst. Civ. Eng.-Struct. Build. 134 257-274
- Dietzel, M., Usdowski, E., Hoefs, J., 1992. Chemical and ¹³C/¹²C- and ¹⁸O/¹⁶O-isotope evolution of alkaline drainage waters and the precipitation of calcite. Appl. Geochem. 7. 177–184.
- Dupre, B., Dessert, C., Oliva, P., Godderis, Y., Viers, J., Francois, L., Millot, R., Gaillardet, J., 2003. Rivers, chemical weathering and Earth's climate. Compt. Rend. Geosci. 335, 1141–1160.
- Fléhoc, C., Girard, J.P., Piantone, P., Bodénan, F., 2006. Stable isotope evidence for the atmospheric origin of CO₂ involved in carbonation of MSWI bottom ash. Appl. Geochem. 21, 2037–2048.
- Fourcade, S., Trotignon, L., Boulvais, P., Techer, I., Elie, M., Vandamme, D., Salameh, E., Khoury, H., 2007. Cementation of kerogen-rich marls by alkaline fluids released during weathering of thermally metamorphosed marly sediments. Part

- I: Isotopic (C, O) study of the Khushaym Matruk natural analogue (central Jordan). Appl. Geochem. 22, 1293–1310.
- Fredericci, C., Zanotto, E.D., Ziemath, E.C., 2000. Crystallization mechanism and properties of a blast furnace slag glass. J. Non-Cryst. Solid. 273, 64–75.
- General Register Office for Scotland, 2000. Scottish Settlements: Urban and Rural Areas in Scotland.
- Harber, A.J., Forth, R.A., 2001. The contamination of former iron and steel works sites. Environ. Geol. 40, 324–330.
- Hendriks, C., Worrell, E., deJager, D., Blok, K., Riemer, P., 2004. Emission Reduction of Greenhouse Gases from the Cement Industry. Greenhouse Gas Control Technologies Conference Paper. 23/08/2004. Emission Reduction of Greenhouse Gases From the Cement Industry: International Energy Agency.
- Hudson, J.D., 1977. Stable isotopes and limestone lithification. J. Geol. Soc. 133, 637–660
- Kalin, R.M., Dardis, G., Lowndes, J., 1997. Secondary carbonates in the Antrim basalts: geochemical weathering at 35KyBP. In: Geofluids II Conference Extended Abstracts, pp. 22–25.
- Knauth, L.P., Brilli, M., Klonowski, S., 2003. Isotope geochemistry of caliche developed on basalt. Geochim. Cosmochim. Acta 67, 185–195.
- Kosednar-Legenstein, B., Dietzel, M., Leis, A., Stingl, K., 2008. Stable carbon and oxygen isotope investigation in historical lime mortar and plaster results from field and experimental study. Appl. Geochem. 23, 2425–2437.
- Kovda, I., Mora, C.I., Wilding, L.P., 2006. Stable isotope compositions of pedogenic carbonates and soil organic matter in a temperate climate vertisol with gilgai, southern Russia. Geoderma 136, 423–435.
- Krishnamurthy, R.V., Schmitt, D., Atekwana, E.A., Baskaran, M., 2003. Isotopic investigations of carbonate growth on concrete structures. Appl. Geochem. 18, 435–444.
- Kuzyakov, Y., Domanski, G., 2000. Carbon input by plants into the soil review. J. Plant Nutr. Soil Sci. 163, 421–431.
- Kuzyakov, Y., Shevtzova, E., Pustovoytov, K., 2006. Carbonate re-crystallization in soil revealed by C-14 labelling: experiment, model and significance for paleoenvironmental reconstructions. Geoderma 131, 45–58.
- Łącka, B., Łanczont, M., Komar, M., Madeyska, T., 2008. Stable isotope composition of carbonates in loess at the Carpathian margin (SE Poland). Studia Quaternaria 25, 3–21.
- Lal, R., 2003. Global potential of soil carbon sequestration to mitigate the greenhouse effect. Crit. Rev. Plant Sci. 22, 151–184.
- Liu, B., Phillips, F.M., Campbell, A.R., 1996. Stable carbon and oxygen isotopes of pedogenic carbonates, Ajo Mountains, southern Arizona: implications for paleoenvironmental change. Palaeogeog. Palaeoclimatol. Palaeoecol. 124, 233–246.
- Macleod, G., Fallick, A.E., Hall, A.J., 1991. The mechanism of carbonate growth on concrete structures, as elucidated by carbon and oxygen isotope analyses. Chem. Geol. 86, 335–343.
- Manning, D.A.C., 2008. Biological enhancement of soil carbonate precipitation: passive removal of atmospheric CO₂. Mineral. Mag. 72, 639–649.
- Mayes, W.M., Younger, P.L., Aumonier, J., 2006. Buffering of alkaline steel slag leachate across a natural wetland. Environ. Sci. Technol. 40, 1237–1243.
- Moulton, K.L., West, J., Berner, R.A., 2000. Solute flux and mineral mass balance approaches to the quantification of plant effects on silicate weathering. Am. J. Sci. 300 (7), 539–570.
- NISRA The Northern Ireland Statistics and Research Agency, 2001. Area Measurements in Northern Ireland.
- Ohlsson, K.E.A., 2000. Carbonation of wood ash recycled to a forest soil as measured by isotope ratio mass spectrometry. Soil Sci. Soc. Am. J. 64, 2155–2161.
- Piovano, E.L., Ariztegui, D., Bernasconi, S., McKenzie, J.A., 2004. Stable isotopic record of hydrological changes in subtropical Laguna Mar Chiquita (Argentina) over the last 230 years. Holocene 14, 525–535.
- Pouyat, R.V., Groffman, P., Yesilonis, I., Hernandez, L., 2002. Soil carbon pools and fluxes in urban ecosystems. Environ. Pollut. 116 (suppl. 1). \$107–\$118.
- Pouyat, R.V., Yesilonis, I.D., Nowak, D.J., 2006. Carbon storage by urban soils in the United States. I. Environ. Oual. 35. 1566–1575.
- Ryan, P.R., Delhaize, E., Jones, D.L., 2001. Function and mechanism of organic anion exudation from plant roots. Ann. Rev. Plant Physiol. Plant Mol. Biol. 52, 527–560
- Salomons, W., Mook, W.G., 1976. Isotope geochemistry of carbonate dissolution and re-precipitation in soils. Soil Sci. 122, 15–24.
- Schlesinger, W.H., 1985. The formation of caliche in soils of the Mojave-desert, California. Geochim. Cosmochim. Acta 49, 57–66.
- Sikes, N.E., Ashley, G.M., 2007. Stable isotopes of pedogenic carbonates as indicators of paleoecology in the Plio-Pleistocene (upper Bed I), western margin of the Olduvai Basin, Tanzania. J. Human Evol. 53, 574–594.
- Singh, B.P., Lee II, Y., Pawar, J.S., Charak, R.S., 2007. Biogenic features in calcretes developed on mudstone: examples from Paleogene sequences of the Himalaya. India Sed. Geol. 201, 149–156.
- Sinha, R., Tandon, S.K., Sanyal, P., Gibling, M.R., Stuben, D., Berner, Z., Ghazanfari, P., 2006. Calcretes from a late Quaternary interfluve in the Ganga Plains, India: Carbonate types and isotopic systems in a monsoonal setting. Palaeogeog. Palaeoclimatol. Palaeoecol. 242, 214–239.
- Smith, P., 2004. Soils as carbon sinks: the global context. Soil Use Manage. 20, 212–218.
- USGS (U.S. Geological Survey), 2004. Minerals Yearbook [Online] http://minerals.usgs.gov/minerals/pubs/country/index.html#pubs (accessed 14.01.09).

- van Hees, P.A.W., Lundstrom, U.S., Morth, C.M., 2002. Dissolution of microcline and labradorite in a forest O horizon extract: the effect of naturally occurring organic acids. Chem. Geol. 189, 199–211.
- van Strydonck, M.J.Y., Dupas, M., Keppens, E., 1989. Isotopic fractionation of oxygen and carbon in lime mortar under natural environmental conditions. Radiocarbon 31, 610–618.
- von Scheele, J., 2006. Short-term opportunities for decreasing $\rm CO_2$ emissions from the steel industry. Int. J. Green Energy 3, 139–148.
- Wang, H., Greenberg, S.E., 2007. Reconstructing the response of C_3 and C_4 plants to decadal-scale climate change during the late Pleistocene in southern Illinois using isotopic analyses of calcified rootlets. Quatern. Res. 67, 136–142.
- Watanabe, Y., Stewart, B.W., Ohmoto, H., 2004. Organic- and carbonate-rich soil formation 2.6 billion years ago at Schagen, East Transvaal district, South Africa. Geochim. Cosmochim. Acta 68, 2129–2151.
- Wilson, A., 1993. Cement and Concrete: Environmental Considerations. Environmental Building News.
- Yanes, Y., Delgado, A., Castillo, C., Alonso, M.R., Ibáñez, M., De la Nuez, J., Kowalewski, M., 2008. Stable isotope (δ^{18} O, δ^{13} C, and δD) signatures of recent terrestrial communities from a low-latitude, oceanic setting: Endemic land snails, plants, rain, and carbonate sediments from the eastern Canary Islands. Chem. Geol. 249, 377–392.
- Zanchetta, G., Vito, M.D., Fallick, A.E., Sulpizio, R., 2000. Stable isotopes of pedogenic carbonates from the Somma-Vesuvius area, southern Italy, over the past 18 kyr: palaeoclimatic implications. J. Quatern. Sci. 15, 813–824.