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Solubility trapping in formation water as dominant CO2 sink in natural gas fields

Stuart M. V. Gilfillan*, Barbara Sherwood Lollar, Greg Holland, Dave Blagburn, Scott Stevens, Martin Schoell, Martin Cassidy, Zhenju Ding, Zheng Zhou, Georges Lacrampe-Couloume and Chris J. Ballentine

*Author to whom correspondence should be addressed:

Scottish Carbon Capture and Storage, School of GeoSciences, The University of Edinburgh, Grant Institute, The King's Buildings, West Mains Road, Edinburgh, EH9 3JW, UK.

Email: stuart.gilfillan@ed.ac.uk

Injecting CO₂ into deep geological strata is proposed as a safe and economically favourable means to store CO₂ captured from industrial point sources¹⁻³. However, it is difficult to assess the long term consequence of CO₂ flooding in the subsurface from decadal observations of existing disposal sites^{1,2}. Both the site design and long term safety modelling critically depend on how and where CO2 will be stored in the site over its lifetime $^{2-4}$. Natural gas fields dominated by a CO_2 phase provide an essential natural analogue for assessing the safety and viability of the geological storage of anthropogenic CO₂ over millennia timescales^{1,2,5,6}. Here, we show that the dominant subsurface sink of CO₂ in nine natural gas fields from North America, China and Europe is through dissolution (solubility trapping) in the formation water. All fields, whether siliciclastic or carbonate dominated reservoir lithologies, exhibit a reduction in CO₂ relative to ³He, an inert and highly insoluble tracer that correlates with an increase in formation waterderived noble gases. Reservoir CO_2 phase loss, sometimes > 90% of that emplaced, is therefore quantitatively related to formation water involvement in the system. CO₂/³He and $\delta^{13}C(CO_2)$ data for seven gas fields indicate that dissolution in formation water at pH=5-5.8 alone is the major sink for the CO₂ loss. Within two siliciclastic dominated reservoirs some CO2 loss through precipitation as carbonate minerals cannot be ruled out, but may account for a maximum of 18% loss of the emplaced CO2. Long term anthropogenic CO₂ storage models in similar geological systems must consider the potential mobility of CO2 dissolved in water and not geological mineral fixation, which is an insignificant CO₂ trapping mechanism.

Noble gas and CO_2 carbon isotopes are powerful tracers of crustal fluid processes that act on subsurface $CO_2^{5,7-10}$. Within a geological storage site, CO_2 injected as a free CO_2 phase (gas or supercritical) may over time be dissolved in solution (solubility trapping), or locked within carbonate minerals by precipitation (mineral trapping)^{4,11}. By using noble gas and carbon

isotope tracers together to study naturally occurring CO₂ systems, we can uniquely identify and quantify the principal mechanism of the CO₂ phase removal, mineral or solubility trapping, over a time scale not accessible through extant injection studies.

We combine noble gas data from five natural CO₂ reservoirs located within the Colorado Plateau and Rocky Mountain provinces, (McCallum Dome, Sheep Mountain, McElmo Dome, CO., Bravo Dome, NM, and St Johns Dome, AZ.)⁷ with new δ¹³C(CO₂) isotope data (Table 1). Previous work has shown that noble gas patterns in these gas fields are explained by CO₂ gas stripping of the formation water during reservoir filling, followed by partial dissolution of noble gases back into the formation water⁷. We also consider published noble gas and stable isotope information in a further four CO₂-rich natural gas fields (JM-Brown Bassett field (JMBB), Permian Basin, Texas⁵; Kismarja field, Pannonian Basin, Hungary⁸; Jilin field, Jilin Province, Songliao Basin; and Subei Basin field, Jiangsu Province in China^{12,13}).

CO₂/³He ratios within the magmatic range of 1-10x10⁹ have been used to identify a primary magmatic origin of the CO₂ contained within five natural CO₂ reservoirs of the Colorado Plateau and Rocky Mountain Provinces⁷. CO₂/³He within the Subei Basin and JM-Brown Bassett field also indicate a magmatic origin, whilst the CO₂/³He values within the Jilin and Kismarja fields are far higher, suggesting a predominantly crustal origin^{5,8}. All of the reservoirs exhibit local variation in the CO₂ content relative to the inert tracer ³He. As there is not a significant source of ³He within the crust¹⁴, and as ³He is inert and highly insoluble⁹, this variation must be due to changes in the CO₂ component within the reservoirs. While many sources and sinks of CO₂ exist in the subsurface^{4,8,9} we argue later that the CO₂/³He variation is caused by CO₂ loss from the reservoir. The difference between the highest CO₂/³He and lower values can provide a minimum estimate of this CO₂ loss. In the case of Bravo Dome, a reduction of CO₂/³He values from 4.82x10⁹ (BD11) to 2.25x10⁹ (BD02)

indicates a >50% loss of the original CO_2 charge in the portion of the reservoir represented by BD02 (Table 1). McElmo Dome samples exhibit a decrease from 8.5×10^9 (YD-1) to 0.68×10^9 (He-2) suggesting >90% emplaced CO_2 loss in portions of this field.

⁴He is continually produced in the subsurface by the radiogenic decay of U, Th and K¹⁴. ²⁰Ne is introduced into the subsurface as a component of air dissolved in water and, as such, can only enter the reservoir system via interaction with formation water⁹. While there is no *a-priori* reason to expect a correlation between ⁴He and ²⁰Ne, this has been observed in natural gases on a regional scale¹⁵. This correlation is the result of ⁴He accumulating in the formation water¹⁶ which also contains atmosphere derived ²⁰Ne, and subsequent quantitative partitioning of both ⁴He and ²⁰Ne into the reservoir phase^{7,15}. Almost all CO₂ reservoirs for which we have ²⁰Ne and ⁴He concentration data show a local ²⁰Ne correlation with ⁴He (Table 1 and supplementary information). A decrease in CO₂/³He is also correlated with ²⁰Ne in most CO₂ reservoirs (Fig. 1) and with ⁴He in all CO₂ reservoirs (Fig. 2).

While there are various mechanisms to add crustal CO_2 ($CO_2/^3He >> 10^{10}$) to these systems 4,10 there is no plausible mechanism that enables crustal CO_2 to be variably added to these systems while preserving a correlation of $CO_2/^3He$ with the formation water-derived noble gases. Neglecting small amounts of 3He dissolution back into the formation water 7 , changes in $CO_2/^3He$ must therefore be due to CO_2 loss in the subsurface via a mechanism directly proportional to the amount of formation water that has been degassed. CO_2 is soluble and reactive. The most likely subsurface CO_2 phase removal mechanisms are solubility and mineral trapping 4,11 .

Reservoir lithology may exert a significant influence on how changes in CO_2 / 3 He relate to $\delta^{13}C(CO_2)$. The carbonate reservoirs (McElmo, JMBB, and St. Johns Domes) show little variance in $\delta^{13}C(CO_2)$ whilst the silicilastic fields (Jilin field, Subei Basin, Kismarja, Sheep

Mountain, McCallum and Bravo Domes) exhibit a greater range in δ^{13} C(CO₂) (Table 1, Supplementary Fig. S1). We consider Bravo and McElmo Domes as case types for each reservoir lithology.

Emplacement of CO₂ at Bravo Dome is believed to have occurred relatively recently (local volcanic activity dates from 8,000-10,000 years)^{7,17} and the field may still be undergoing active CO₂ recharge¹¹. Decreasing CO₂/³He within Bravo Dome correlates with more negative δ^{13} C(CO₂) (Fig. 3a). Taking the highest CO₂/³He of 4.82 x 10⁹ (BD11) to be the sample that experienced the least CO_2 loss, we calculate the coherent change in CO_2 ³He and δ¹³C(CO₂) predicted for CO₂ dissolution into the formation water at various pH and for CO₂ precipitation as a carbonate (see Methods Summary). The data are not consistent with precipitation as carbonate being a major sink for CO₂ at Bravo Dome (Figure 3a). However, while a significant number of the data points are consistent with CO₂ dissolution into formation water at a pH between 6-7, it is not possible to rule out a degree of CO₂ loss due to precipitation together with CO₂ dissolution at a lower pH (e.g. pH=5). In such a two process model an upper limit to the proportion of CO₂ lost to precipitation of approximately 18%, can be attributed (Fig. 3a). Hence, in all cases the major CO₂ sink is dissolution. In situ precipitation of 18% reservoir CO₂ would generate between 3.2-6.1% by mass of the whole rock, dependent on whether dolomite, calcite or dawsonite precipitation was favoured by the reservoir conditions. Whilst evidence for CO₂ rich formation water interaction within the reservoir has been documented, to date no secondary carbonate has been identified¹⁸. Nevertheless, the volume control of the water suggests that the location of the precipitate, if any, is likely to be within the water leg which was not sampled. Lack of reservoir secondary mineralization cannot at this stage rule out any carbonate precipitation as a minor CO₂ sink.

Similar to Bravo Dome, while many of the Sheep Mountain data can be accounted for by dissolution of CO₂ (at pH=5 in this case), a small component of precipitation cannot be ruled out. Adopting the same approach as Bravo Dome, the remaining Sheep Mountain data require a maximum of 10% precipitation and 20% dissolution of the original CO₂ charge (Table 1; Supplementary Fig. S2). In contrast, while minor data scatter may also be due to some small amount of CO₂ precipitation or dissolution at pH=7-8, almost all the data from the other siliciclastic fields of McCallum Dome, Subei basin, Kismarja and the Jilin field can be described by dissolution in the formation water only, within a narrow pH range of between 5-5.3 (Supplementary Figs S3-6).

McElmo Dome carbonate reservoir data show over an order of magnitude change in $CO_2/^3He$ with invariant $\delta^{13}C(CO_2)$ (Figure 3b). This pattern is repeated in the two other carbonate-dominated fields (Supplementary Figures S7, S8). Invariant $\delta^{13}C(CO_2)$ in these fields allows us to discount a two process model of precipitation and dissolution such as at Bravo Dome (Fig. 3a). All data are consistent with CO_2 dissolution into formation water in the pH range of 5.4-5.8 (Figure 3b, Supplementary Figures 7,8), a value similar to the pH obtained for the siliciclastic reservoirs and to values observed (pH=5.7) in carbonate mineral buffered formation water observed in the recent Frio CO_2 injection studies on CO_2 breakthrough¹⁹.

On a reservoir engineering timescale, the early stages of CO_2 injection can result in a drop in pH and dissolution of carbonate minerals into the formation water^{18,20-22}. Any significant CO_2 contribution to the reservoir CO_2 phase from re-dissolution of carbonates would be 3 He-free and therefore perturb the $CO_2/^3$ He correlation with 4 He and 20 Ne. As there is a clear correlation between $CO_2/^3$ He and 4 He in all fields and 20 Ne within the majority, we conclude that dissolution of carbonate minerals into the formation water cannot have had a major influence on $\delta^{13}C(CO_2)$ values. There is no evidence for any precipitation of CO_2 within the

carbonate dominated reservoirs, requiring that the dominant mechanism of reservoir CO₂ loss, up to 90%, is through dissolution into the formation water.

Even the most conservative model we have presented places an upper limit on the CO₂ removed by precipitation at approximately18%, and then only in some samples, from all natural gas fields investigated in a variety of lithological settings. Precipitation of CO₂ over millenial timescales represents at most only a small subsurface trapping mechanism for CO₂, and then only within siliciclastic lithologies. The dominant mechanism of CO₂ loss from most CO₂ natural gas fields can be accounted for through simple dissolution into the formation groundwater within a narrow pH window (pH=5-5.8). This study underscores that geological carbon storage requires careful investigation of existing geologic and hydrogeologic analogues that have naturally accumulated and stored CO₂ over timescales relevant to anthropogenic CO₂ storage facilities. We further demonstrate a means of testing trapping and storage mechanisms via coupled noble gas and carbon isotope measurements in the context of formation/reservoir water pH evolution.

Methods Summary

Detailed descriptions of the sample collection and analysis procedures can be found in the original references^{5,7,8,21,23}. In our calculations (Figure 3 and Supplemental Figures) we use the highest CO_2 / 3 He ratio measured in each field as a reference point to calculate the correlated reservoir CO_2 / 3 He and $\delta^{13}C(CO_2)$ ratios as the CO_2 phase is removed by either precipitation or dissolution. We assume open system loss. In the case of precipitation there is zero 3 He loss from the CO_2 phase and CO_2 / 3 He changes in proportion to the fraction of the remaining CO_2 phase (f). In the case of dissolution the change in CO_2 / 3 He is calculated following the Rayleigh equation.

Changes in $\delta^{13}C(CO_2)$ are calculated using the Rayleigh fractionation equation expressed as: $\delta^{13}C(CO_2) = \delta^{13}C(CO_2)_0 + \epsilon \ln(f)^{24}$

where $\delta^{13}C(CO_2)_0$ is the original system value, f is the fraction of CO_2 remaining in the reservoir, and ϵ is the carbon isotope fractionation, either for precipitation or for dissolution. Carbon isotope fractionation factors (α) are calculated as a function of temperature for $CO_2(g)$ precipitating to form $CaCO_3(s)$, or dissolving to form either $H_2CO_3(aq)$ or HCO_3^- (aq)²⁵. Since all the fractionations are small the simplification can be made that $\epsilon = 1000 ln(\alpha)^{26}$. For typical reservoir waters of pH range 5-8, the contribution of CO_3^{2-} (aq) is negligible. Hence for CO_2 dissolution, carbon isotope fractionation between the dissolved inorganic carbon (DIC) pool and CO_2 gas used in the Rayleigh fractionation equation can be expressed as:

$$\epsilon^{13}C_{DIC\text{-}CO2(g)} = \ x(\epsilon^{13}C_{H2CO3(aq)\text{-}CO2(g)}) + (1-x)(\epsilon^{13}C_{HCO3(aq)\text{-}CO2(g)})^{24}$$

where x is the proportion of CO_2 gas dissolving to $H_2CO_3(aq)$ at the relevant pH²⁴.

Solubility as a function of temperature and salinity is given by the IUPAC solubility series for CO_2^{27} and by Crovetto et al. and Smith for $He^{28,29}$. The average well depth, reservoir pressure, temperature and salinity are presented in the supplementary information for each reservoir, with the corresponding Henry's Law constants K_{He} , K_{CO2} , and fractionation factor $(1000 ln\alpha)$ for $CO_2(g)$ forming $H_2CO_3(aq)$, $HCO_3^-(aq)$ and $CaCO_3(s)$ (Supplementary Table 1).

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Supplementary Information is linked to the online version of the paper at www.nature.com/nature.

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Author Contributions S.G., C.B. and B.S.L. designed the study, analysed the samples, interpreted the data and wrote the paper. G.H., D.B., Z.D., Z.Z. and G.L.C. assisted with sample analysis and interpretation of the data. S.S., M.S. and M.C. assisted with sample collection and provided comments on the manuscript.

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Table 1: Sample location, Producing formation, major gas species and CO₂ carbon isotopes

				2 4			42
Field & Well	Location	Producing	CO ₂ / ³ He	³He/⁴He	⁴ He (x 10 ⁻⁴)	²⁰ Ne (x 10 ⁻⁸)	δ ¹³ C(CO ₂)
Prove Dome ⁷	Twnshp-Rnge/Lat-Long	Formation	x 10 ⁹	(R/R _a)	cm ³ (STP)cm ⁻³	cm ³ (STP)cm ²	‰
Bravo Dome' BD01	23/19N/34E	Tubb	4.53 (10)	1.670 (8)	0.944 (12)	0.169 (2)	-3.96 (4)
BD02	32/21N/35E	Tubb	2.25 (5)	0.764 (4)	4.15 (5)	0.700 (7)	-4.93 (8)
BD03	36/22N/34E	Tubb	2.41 (5)	0.896 (4)	3.31 (4)	0.521 (5)	-4.89 (19)
BD04	8/20N/34E	Tubb	4.61 (10)	1.611 (8)	0.961 (2)	0.181 (2)	-4.23 (8)
BD05	34/20N/35E	Tubb	2.74 (6)	0.965 (5)	2.70 (4)	0.446 (4)	-4.95 (5)
BD06	26/22N/32E	Tubb	3.94 (8)	1.503 (8)	1.20 (2)	0.202 (2)	-4.55 (11)
BD07	3/19N/33E	Tubb	4.34 (9)	2.104 (11)	0.781 (10)	0.180 (2)	-4.85 (1)
BD08	9/18N/33E	Tubb	3.87 (8)	1.143 (6)	1.61 (2)	0.264 (3)	-3.88 (8)
BD09	17/21N/33E	Tubb	4.22 (9)	1.724 (9)	0.981 (12)	0.180 (2)	-4.44 (11)
BD10	7/22N/34E	Tubb	3.25 (6)	1.104 (6)	1.99 (3)	0.308 (3)	-4.88 (7)
BD11	25/19N/30E	Tubb	4.82 (10)	3.784 (19)	0.391 (5)	0.103 (1)	-3.66 (29)
BD12	27/19N/30E	Tubb	4.74 (10)	3.627 (18)	0.415 (6)	0.240 (2)	-3.94 (17)
BD13 BD14	22/18N/35E 16/18N/34E	Tubb Tubb	3.54 (8) 4.39 (9)	1.318 (7) 1.413 (7)	1.53 (2) 1.15 (2)	0.240 (3) 0.179 (4)	-4.42 (3) -4.04 (2)
BD12b	27/19N/30E	Tubb	4.39 (9)	3.634 (18)	0.413 (6)	0.179 (4)	-4.04 (2) -3.94 (17)
McCallum Dome ⁷	21/19IN/30L	Tubb	4.73 (10)	3.034 (10)	0.413 (0)	0.120 (2)	-3.94 (17)
No. 3 (8-3)	8/9N/78W	Lakota	1.52 (4)	0.354 (7)	12.3 (2)	1.17 (2)	-5.1 (3)
No. 5	3/9N/79W	Lakota	1.04 (3)	0.409 (7)	15.5 (2)	2.71 (3)	-5.2 (1)
No. 36	8/9N/79W	Dakota/Lakota	- (-)	0.448 (8)	1.32 (12)	8.10 (8)	nm
No. 13	2/9N/79W	Lakota/Morrison	0.89(2)	0.393 (7)	18.8 (2)	4.36 (5)	-5.3 (2)
No. 79	4/9N/79W	Dakota/Lakota	1.77 (6)	0.406 (6)	9.16 (21)	2.53 (3)	-5.7 (1)
McElmo Dome ⁷							
MC-1	37.4155, -108.7713	Leadville	5.04 (11)	0.145 (2)	9.58 (8)	0.376 (4)	-4.26 (10)
HE-2	37.5052, -108.9094	Leadville	0.68 (15)	0.148 (1)	70.5 (7)	0.307 (30)	-4.40 (10)
YC-4	37.4529, -108.8583	Leadville	4.96 (11)	0.137 (3)	10.2 (10)	0.573 (6)	-4.41 (10)
SC-9	37.3934, -108.8733	Leadville	3.17 (7)	0.150 (3)	14.8 (14)	0.497 (5)	-4.29 (10)
YB-2	37.4472, -108.8075	Leadville	8.74 (20)	0.125 (1)	6.42 (61)	0.371 (4)	-4.40 (10)
YC-1	37.4529, -108.8583	Leadville	4.07 (9)	0.142 (2)	12.1 (12)	0.423 (5)	-4.34 (10)
HF-1	37.4871 -108.8807	Leadville	2.16 (6)	0.169 (1)	19.3 (26)	0.564 (12)	-4.37 (10)
HD-2 YA-2	37.4572, -108.9008 37.4692, -108.7811	Leadville Leadville	4.28 (10) 3.39 (8)	0.140 (3) 0.138 (3)	11.7 (12) 15.0 (15)	0.128 (2) 0.130 (2)	-4.38 (10) -4.42 (10)
YE-1	37.4818, -108.8123	Leadville	4.16 (9)	0.138 (3)	9.75 (8)	0.130 (2)	-4.42 (10) -4.45 (10)
HA-1	37.5289, -108.8718	Leadville	4.56 (10)	0.173 (3)	11.0 (11)	0.205 (7)	-4.66 (10)
SC-10	37.3934, -108.8733	Leadville	4.37 (10)	0.139 (2)	11.6 (11)	0.413 (5)	-4.27 (10)
HC-2	37.4734, -108.8860	Leadville	4.68 (11)	0.140 (2)	10.7 (10)	0.409 (5)	-4.38 (10)
HB-1	37.5087, -108.8802	Leadville	4.74 (11)	0.148 (3)	9.94 (10)	0.247 (4)	-4.49 (10)
YD-1	37.4619, -108.8224	Leadville	8.50 (20)	0.145 (3)	5.68 (6)	0.366 (5)	-4.46 (10)
JM Brown Basset⁵							
Turk State No. 1A	30.38758, -101.85642		5.92 (47)	0.543 (16)	1.25 (9)	nm	-2.88
Bassett Goode No. 3	30.37852, -101.83068	•	5.55 (43)	0.527 (16)	1.42 (10)	nm	-2.89
Brown Bassett No. 2*	30.34433, -101.7995		5.82 (35)	0.502 (15)	1.33 (7)	nm	-2.90
Mayme K. Martin ETAL 1		•	5.29 (40)	0.372 (11)	1.42 (10)	nm	-2.97
Mitchell 109 No. 2*	30.33329, -101.69826		4.58 (36)	0.400 (12)	1.53 (11)	nm	-2.92
Mitchell 5 No. 1X Mitchell 103 No. 2	30.32352, -101.68429 30.3568, -101.63642		5.61 (43)	0.478 (11) 0.246 (7)	1.40(10) 1.39 (10)	nm	-2.84 -2.70
Mitchell No. 6	30.351, -101.58835		4.20 (33) 3.93 (31)	0.240 (7)	1.51 (11)	nm nm	-2.70
Mitchell No. 3	30.33966, -101.61307	•	4.22 (33)	0.240 (7)	1.39 (10)	nm	-3.06
Mitchell A-11 No. 1	30.30286, -101.57677	•	4.07 (32)	0.272 (8)	1.66 (12)	nm	-2.93
Mitchell No. 12	30.29118, -101.57295		4.24 (130)	0.267 (8)	1.46 (10)	nm	-2.96
Sheep Mountain ⁷	•	· ·	` ,	()	` ,		
8-2-P	2/9-28S/70W	Dakota	2.31 (5)	0.981 (10)	3.13 (3)	1.47 (2)	-5.0 (2)
2-10-O	15/9-27S/70W	Entrada	2.44 (6)	0.984 (12)	2.96 (3)	3.04 (3)	-5.2 (1)
9-26	26/9-27S/70W	Dakota	2.57 (6)	0.934 (14)	2.95 (3)	0.613 (9)	nm
2-9-H	9/9-27S/70W	Dakota	2.44 (6)	0.945 (19)	3.07 (3)	9.77 (10)	nm
3-15-B	15/9-27S/70W	Dakota	2.61 (6)	0.937 (16)	2.90 (3)	1.54 (2)	-5.7 (4)
4-13	00/0 070/70/4/	Dakota	2.17 (5)	0.942 (18)	3.47 (4)	1.11 (2)	nm
4-26-E	26/9-27S/70W	Entrada	2.20 (5)	1.024 (18)	3.15 (3)	0.442 (4)	-4.8 (1)
3-23-D 7-35-L	22/9-27S/70W	Dakota Dakota	2.26 (5)	0.988 (14)	3.17 (3)	0.579 (9)	nm -5.0 (2)
7-35-L 2-35-C	2/9-28S/70W 26/9-27S/70W	Dakota Dakota	2.53 (6) 2.57 (6)	0.916 (14) 0.963 (19)	3.06 (3) 2.87 (3)	0.749 (12) 0.573 (8)	-5.0 (2) nm
1-15-C	15/9-27S/70W	Entrada	2.57 (6)	0.963 (19)	2.71 (3)	6.77 (10)	nm
3-4-O	9/9-27S/70W	Dakota	2.71 (6)	0.937 (10)	2.99 (3)	2.64 (3)	-5.8 (3)
4-14-M	22/9-27S/70W	Dakota	2.65 (6)	0.892 (15)	3.00 (3)	1.11 (1)	nm
5-15-O	22/9-27S/70W	Dakota	2.30 (5)	1.056 (15)	2.92 (3)	4.33 (5)	-5.0 (1)
			` '	(- /	` '	` '	` '

Field & Well	Location Twnshp-Rnge/Lat-Lon	Producing g Formation	CO ₂ / ³ He x 10 ⁹	³He/⁴He (R/R _a)	⁴ He (x 10 ⁻⁴) cm ³ (STP)cm ⁻³	²⁰ Ne (x 10 ⁻⁸) cm ³ (STP)cm ⁻³	δ ¹³ C(CO ₂) ‰
Sheep Mountain ⁷			2.90 (7)				
4-4-P	9/9-27S/70W	Dakota		0.970 (14)	2.52 (2)	1.31 (2)	nm
5-9-A	9/9-27S/70W	Dakota	2.39 (6)	1.006 (18)	2.94 (3)	1.28 (2)	nm
1-1-J	2/9-28S/70W	Dakota	3.61 (8)	0.908 (16)	2.16 (2)	0.878 (12)	-5.2 (1)
1-22-H	22/9-28S/70W	Entrada	2.25 (5)	0.981 (17)	3.22 (3)	0.937 (13)	-4.5 (2)
St. Johns Dome ⁷			` ,	` ,	, ,	` ,	` ,
22-1X	34.4265, -109.2664	Supai	0.098 (2)	0.455 (8)	134 (13)	34.4 (47)	-3.65 (5)
10-22	34.2437, -109.1645	Supai	1.91 (42)	0.394 (8)	9.42 (9)	2.30 (4)	-3.79 (5)
3-1	34.3771, -109.2563	Supai	0.22(3)	0.433 (9)	70.6 (7)	15.1 (21)	-3.85 (5)
Jillin Field ^{12, 13, 23}							
Wan 2		Cretaceous	1.44 (4)	4.91 (6)	1.00 (2)	nm	-3.6
Wan 5		Cretaceous	227 (7)	4.10 (4)	0.0076 (2)	0.0547 (15)	-5.0
Wan 6		Cretaceous	8.32 (3)	4.99 (5)	0.169 (4)	0.230 (6)	-3.8
Wan 8		Cretaceous	nm	4.30 (5)	nm	nm	-3.2
Wan 9		Cretaceous	36.6 (10)	4.08 (4)	0.047 (1)	0.130(3)	-3.8
Subai Basin ^{12, 23}							
Huangqyan 1		Permian	2.17 (7)	3.52 (5)	3.13 (3)	1.47 (2)	-3.6
Sutail 74		Devonian	0.493(14)	3.59 (4)	2.96 (3)	3.04 (3)	-4.1
Su203		Eocene	0.459 (13)	2.61 (3)	2.95 (3)	0.613 (9)	-2.7
Kismarja ^{8,30}							
Kismarja 8		Up. Pannonian	20.2 (5)	1.33 (3)	0.226 (7)	nm	-5.0
Kismarja 79		Up. Pannonian	15.5 (4)	1.38 (3)	0.310 (10)	nm	-4.9
Kismarja 61		Up. Pannonian	27.3 (6)	1.16 (2)	0.205 (6)	nm	-5.1
Kismarja 55		Up. Pannonian	13.3 (3)	1.38 (3)	0.360 (11)	nm	-5.1
Kismarja 56		Up. Pannonian	1090 (3)	1.16 (2)	0.0052 (2)	nm	-6.8
Kismarja 74		Up. Pannonian	65.2 (2)	1.34 (3)	0.078 (3)	nm	-6.4
Kismarja 22		Up. Pannonian	1.52 (1)	1.02 (2)	1.31 (3)	nm	-6.6
nm = not measured 1 σ error shown in bracket						ackets	

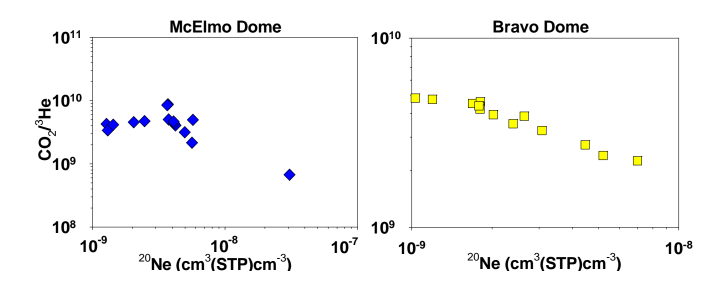
Figure Captions

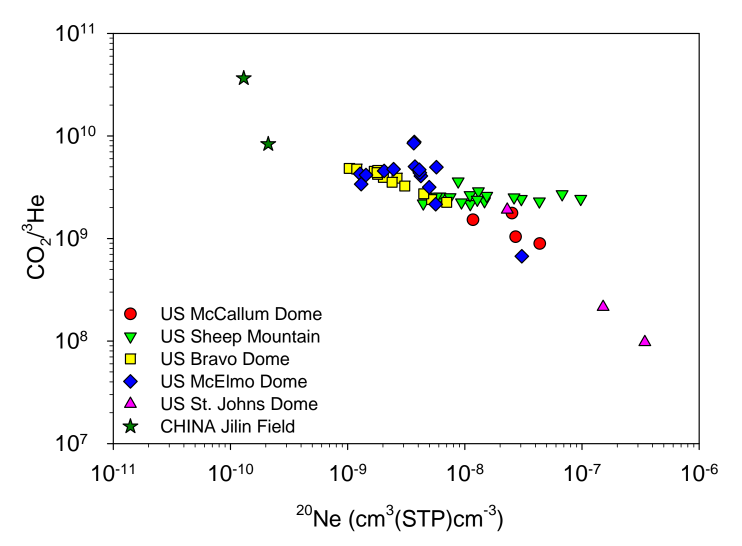
Figure 1. CO₂/³He variation plotted against ²⁰Ne in samples from the 'global' data set of CO₂-rich natural gas fields (see text). There is a general trend in this data set of decreasing CO₂/³He with increasing ²⁰Ne. This trend is most clearly apparent in the siliciclastic case type Bravo Dome data set (inset) but less clear in the carbonate case type reservoir, McElmo Dome (inset). ³He is conservative within the gas phase. Lower CO₂/³He therefore represent subsurface reduction in CO₂ concentration in the emplaced CO₂ phase. Since the only subsurface source of the ²⁰Ne now in the CO₂ phase is the formation water, the CO₂ sink must be linked to the formation water contacted by the gas phase.

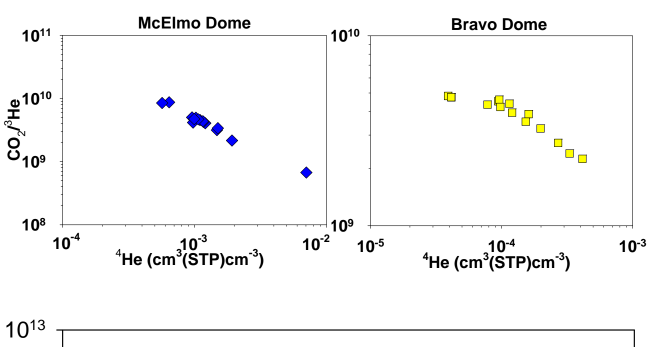
Figure 2. The 'global' data sets of CO₂ gas fields also show a strong correlation between decreasing CO₂/³He and increasing ⁴He concentration. ⁴He accumulates in formation water over time^{7,15,16} and underscores the importance of formation water in controlling the mechanism of subsurface CO₂ removal (Fig.1 and text). We speculate that the formation water ⁴He signature with CO₂/³He is more coherent than the ²⁰Ne (Fig. 1) due to perturbation of ²⁰Ne in ancient formation water through non-water phase interaction⁹ with subsequent ⁴He accumulation providing a homogenous regional scale formation water ⁴He signal^{15,16}. Different CO₂/³He vs. ⁴He gradients will be due to different local formation water ⁴He accumulation rates.

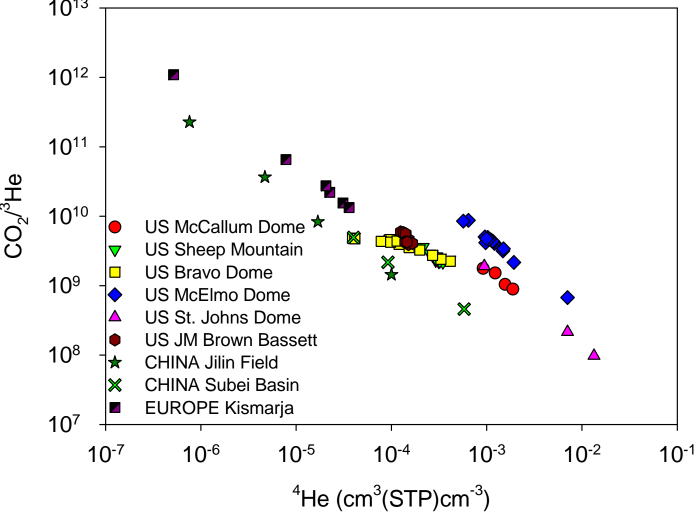
Figure 3. δ^{13} C(CO₂) against CO₂/ 3 He for Bravo Dome (Top) and McElmo Dome (Bottom) Error bars are 1 σ . Top Panel: The solid line shows the predicted trend for carbonate mineral precipitation and the dashed lines show CO₂(g) dissolution trends

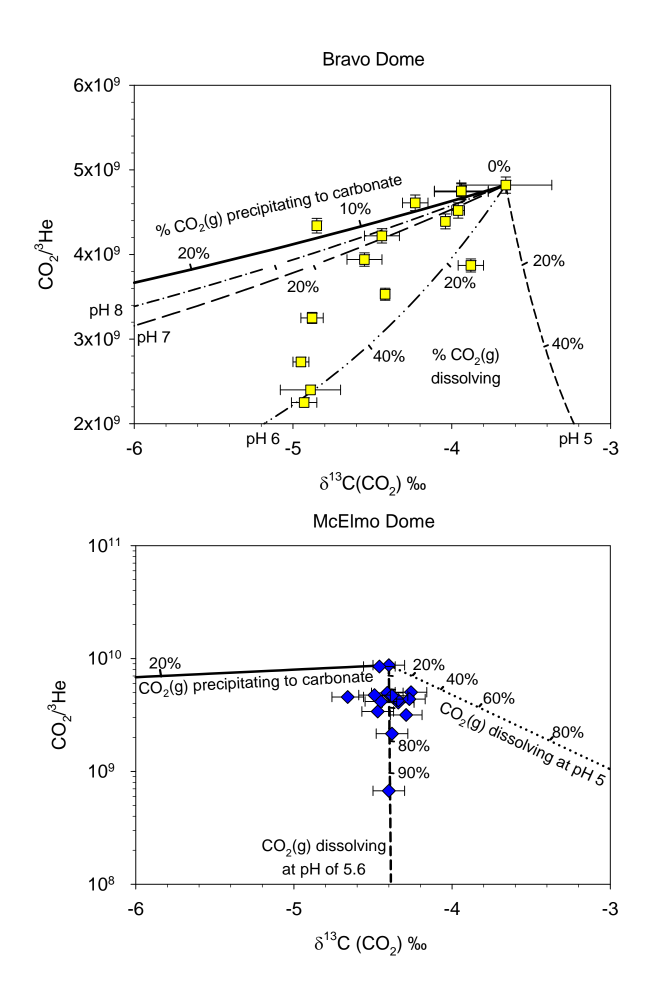
for varying formation water pH (see methods). Bravo Dome data is not consistent with the major CO_2 sink being precipitation of carbonate (see text). Bottom Panel: Invariant $\delta^{13}C(CO_2)$ with over an order of magnitude change in CO_2 / 3 He in McElmo Dome gases cannot be accounted for by precipitation (solid line). Dissolution of reservoir CO_2 into formation water at pH=5.6 would produce the observed results (see text).











Supplementary Information:

Supplementary Table 1 Reservoir Conditions Used in Models

-	Reservoir	Average Well Depth (m)	Pressure (MPa)	Borehole Temperature (K)	TDS (molar)	K _{He} (GPa)	K _{CO2} (GPa)	Fractionation (1000lnα) for CO _{2(g)} forming (‰) [24]		
_	- Neser von							H ₂ CO _{3(aq)}	HCO ₃	CaCO ₃
	Bravo Dome ³⁰	820	8.03	314	1.45	20.1	0.349	-0.846	6.63	8.55
	JM Brown Bassett⁵	2800*	27.4*	373	1.00*	14.0	0.774	-0.864	3.36	4.96
l	McCallum Dome ⁷	1630	16.0	<u>338</u>	0.228	14.1	0.460	-0.854	5.10	6.85
l	McElmo Dome ⁷	2450	24.0	344	0.200	11.9	0.487	-0.856	4.76	6.48
	Sheep Mountain ⁷	1400	13.7	331	0.0137	13.6	0.438	-0.852	5.51	7.29
	St. Johns Dome ³¹	630	6.17	322	0.0720	14.4	0.281	-0.849	6.08	7.92
l	Jilin Field ²¹	840	8.23	333*	1.00*	17.4	0.491	-0. <u>853</u>	5.39	7.16
	Subei Basin ¹²	2251	22.1	357*	1.00*	15.4	0.665	-0.860	4.09	5.75
	Kismarja ²⁹	825	8.08	326*	1.00*	17.7	0.438	-0.850	5.82	7.63

^{*}JM Brown Bassett depth estimated from bottom hole temperature and 30°C/km geothermal gradient. Jilin Field, Subei Basin and Kismarja borehole temperatures estimated from depth and geothermal gradient. JM Brown Bassett, Jilin Field, Subei Basin and Kismarja salinity estimated.

Additional References for Supplementary Table

- 30. Broadhead, R. F. Carbon dioxide in northest New Mexico. *West Texas Geological Society Bulletin* **32**, 5-8 (1993).
- 31. Stevens, S. H., Fox, C., White, T. & Melzer, S. Natural CO₂ analogs for Carbon Sequestration. *Final Report for USDOE* (2006).