

CLAY MINERALS AS THE KEY TO THE SEQUESTRATION OF CARBON IN SOILS

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Abstract -Results from earlier laboratory and field experiments are interrogated for the possibilities of sequestration, or long-term accumulation, of carbon from excess greenhouse gases in the atmosphere. In the laboratory study, samples of three (top) soils dominated by kaolinite and illite together, smectite and allophane were examined for the adsorption and desorption of dissolved organic carbon (DOC). Adsorption and desorption of DOC were carried out on clay fractions extracted physically and after first native organic matter and then iron oxides were removed chemically. Labelled organic material was added to the soils to assess the priming effect of organic carbon (OC). In the field, changes in OC were measured in sandy soils that had been amended by additions of clay for between 3 and 17 years both through incorporation of exogenous clay and delving of *in situ* clay. The laboratory experiments demonstrated that a portion of DOC was held strongly in all soils. The amount of DOC adsorbed depended on clay mineral types, including Fe oxides. Much adsorbed DOC was lost by desorption in water and substantial native OC was lost on priming with new OC. Addition of clay to soils led to increased OC. Therefore, addition of clay to soil may enhance net sequestration of C. OC close to mineral surfaces or within microaggregates is held most strongly. C sequestration may occur in subsoils with unsaturated mineral surfaces. However, incorporation of carbon into macroaggregates from enhanced plant growth might be most effective to remove excess carbon from the atmosphere, albeit over the short-term.

27 **Key Words:** -Adsorption, Desorption, Priming, Clay Amendment, Delving, Dissolved Organic Carbon.

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INTRODUCTION

33 The earth's climate is undergoing change, and the consensus is that its principal cause is the
34 rise in the atmosphere of so-called "greenhouse gases" from anthropogenic activities. In particular,
35 carbon dioxide generated by the combustion of fossil forms of carbon (Le Quéré *et al.*, 2009) and
36 also methane, often generated by agriculture (Smith *et al.*, 2008), are major sources of these gases
37 in the atmosphere. These gaseous forms of carbon are also released into the atmosphere when
38 formerly frozen sources such as permafrost Arctic regions are warmed (Schuur *et al.*, 2011).

39 Many authors in the scientific literature (see, e.g. Stockmann *et al.*, 2013; Minasny *et al.*
40 2017) have proposed that soil organic matter (SOM) can be used as a "managed" sink for
41 atmospheric carbon gases and particularly carbon dioxide, through carbon sequestration. Politicians
42 and others in public life have seen carbon sequestration as a solution to the problems that rising
43 emissions of these gases cause to the climate. Hence there is a strong need to assess the viability of
44 the apparently useful role for SOM in halting or at least diminishing the advance of climate change.

45 The global stock of carbon in the soil as organic matter (SOM) has been estimated as ~ 700
46 Pg to a depth of 30 cm and ~1500 Pg to a depth of 1 m (Batjes, 1996). The SOM to 30 cm depth
47 represents about twice the amount of C in the atmosphere and 3 times that in above-ground
48 vegetation (Powlson *et al.*, 2011). Carbon as soil organic carbon (SOC) is particularly labile.
49 Historically, the reservoir of C held in land has diminished drastically over decades, whereas that
50 held in oceans, which is also labile and of a comparable size, has changed only a little (Lal, 2014). The

51 loss of C from land is attributable to agriculture. Many (e.g. Lal (2014) have concluded that, because
52 C has been lost from their soils, managed agricultural soils have the capacity to replenish the losses
53 and hence sequester C from the atmosphere.

54 The clay fraction of soils, comprising clay minerals (including metal oxides, oxyhydroxides
55 and hydroxides) is their most reactive fraction for forming associations with organic matter
56 (Churchman, 2010, 2018; Sarkar *et al.*, 2018; Singh *et al.*, 2018).

57 Several approaches have been used to investigate the importance of clay minerals in the
58 uptake and retention, hence, potentially, the sequestration of carbon in soils (e.g. Churchman and
59 Velde, 2019). These include correlations of the contents of SOC with properties of soil clays,
60 including their contents and their cation exchange capacities (CECs). Where narrow ranges of soil
61 types have been studied, some good correlations have been obtained, but where the ranges are
62 wider, SOC contents and those of clays or their properties have been poorly correlated, if at all
63 (Churchman and Velde, 2019). In tropical Australia (Spain, 1990), such correlations were poor when
64 soils originated from basalt were included, and a set of 167 soils taken from throughout New
65 Zealand showed no correlation between SOC and clay fraction contents (Percival *et al.*, 2000). The
66 lack of a relationship between SOC and clay contents or properties was explained by the important
67 roles played by compounds of Fe (Spain, 1990; Percival *et al.*, 2000) and Al (Percival *et al.*, 2000),
68 particularly when these were poorly crystalline (Percival *et al.*, 2000).

69 Another approach to the study of mineral-organic interactions in soils include the
70 fractionation of soils, either by particle density, selective chemical dissolutions or physical
71 disaggregation. These studies have confirmed the important roles played by Fe (Eusterhues *et al.*,
72 2003, 2005) in the stabilisation of SOC in many soils, and by Ca^{2+} in some (Oades, 1989). In particular,
73 leached, hence, generally, acid soils involve Fe and Al in the stabilisation of C by bridging while Ca^{2+} is
74 more involved in bridging with SOC in high base status soils (Oades, 1989; Rowley *et al.*, 2018).

75 Studies of incubations of soils have revealed that clay minerals with the highest surface area,
76 namely allophane and smectite in the soils they used, had the strongest effect on the suppression of
77 the decomposition of SOC (Saggar *et al.*, 1996), while Rasmussen *et al.* (2007) found that poorly
78 crystalline Fe oxyhydroxides strongly suppressed SOC decomposition.

79 These three types of approaches, together with others (e.g. Churchman and Velde, 2019),
80 have shown that clay minerals, including metal oxides, oxyhydroxides and hydroxides, are important
81 in the uptake and retention of SOC in soils. Even so, these studies do not provide an assessment of
82 the strength and duration of the binding of SOM by soil minerals. Strong and long-term binding of
83 SOM is required for the sequestration of carbon in soils. “Sequester” is defined as “to hold on to” or
84 “to keep separate” (Powlson *et al.*, 2011). In the chemical or environmental context, sequestration
85 means “the trapping of a chemical in the atmosphere or environment and its isolation in a natural or
86 artificial storage area” (Dictionary.com, 2019). The sequestration of organic C in soils is generally
87 taken to mean the retention of SOC for a “stipulated duration timeframe (usually 100 years)”
88 (Stockmann *et al.*, 2013). The most useful meaning for carbon sequestration for a decrease in
89 greenhouse gases in the atmosphere is given by a net gain of SOC. This represents “new”
90 sequestration of carbon.

91 Several recent experiments (Singh *et al.*, 2016, 2017a; 2017b; 2019; Schapel *et al.*, 2018)
92 have been carried out to help assess or improve the effectiveness of soils for the uptake and
93 retention of OC. Experiments were carried out in the laboratory and also in the field.

94 The laboratory experiments included determinations of the stabilisation of OC by clay
95 minerals as they occur in soils, in contrast to clay minerals from deposits, as determined by Saidy *et al.*
96 *al.* (2012). Clay minerals originating in soils can have quite different properties from “pure” clay
97 minerals from deposits (Churchman, 2010; Churchman *et al.*, 2012; Churchman and Lowe, 2012;
98 Churchman and Velde, 2019).

147 Clay fraction samples taken from the adsorption experiments at various levels of equilibrium
148 concentrations of DOC were first washed with Milli-Q water to remove entrained DOC then
149 equilibrated for 24 h and supernatants were analysed for DOC after filtering. See Singh *et al.* (2016)
150 for details.

151 The percentages of adsorbed C that was desorbed in water from each clay fraction without
152 treatment, after OC removal and after iron removal following OC removal, in relation to the amounts
153 of DOC added in the adsorption experiments, are shown in Table 2. These data are from adsorption
154 in 0.01M Ca(NO₃)₂. There were similar trends for desorption for clay fractions following adsorption
155 from the nitrate solutions differing in calcium or sodium and with two different concentrations (0.01
156 or 0.1M) (Singh *et al.*, 2016).

157 (Insert Table 2 about here)

158 More than 30% of the C adsorbed in Ca nitrate and more than 50% of that adsorbed in Na
159 nitrate was released on simple desorption in water (Singh *et al.*, 2016). The proportion of adsorbed C
160 that was desorbed was greatest for the kaolinite-illite clay, less for the smectite clay and least for the
161 allophanic clay, but even in this latter case, ~ 20% of that adsorbed in Ca nitrate and ~30% of that
162 adsorbed in Na nitrate was lost on desorption in water. In each case, the proportion of adsorbed C
163 that was lost on desorption increased when DOC adsorption had been performed on clays from
164 which Fe-oxide was removed, and diminished when adsorption had occurred on clays from which
165 native C was removed.

166 *Effects of priming*

167 Samples of the whole soils that had been sieved to < 2 mm to remove coarse particles were
168 equilibrated with ¹⁴C – labelled malic acid and also unamended in sterile Milli-Q water. CO₂ from
169 respiration was collected at different time periods in a solution of NaOH and the unconsumed alkali

170 was determined by titration with HCl. See Singh *et al.* (2017a) for details of the experiments and
171 calculations.

172 Results showed that more C was lost (as CO₂) from all soils after priming with malic acid
173 (Singh *et al.*, 2017a). However, the extent of loss on priming depended upon the mineralogy of the
174 soils, with more C lost from the smectite soil than from the soil dominated by illite-kaolinite and the
175 allophanic soil. These two latter soils had similar losses on priming. Decomposition of native C
176 increased by 108% in the smectitic soil upon priming and by 37% in the kaolinitic-illitic soil and 42%
177 in the allophanic soil (Singh *et al.*, 2017a).

178 FIELD EXPERIMENTS AND RESULTS

179 Soils in South Australia that were classified as Luvisols in the World Reference Base (Michéli
180 and Spaargaren, 2012) were studied at four sites where clay amendments had been carried out. All
181 soils were texture contrast. Farming systems in all sites were rotations of cereal and grazed pasture,
182 using minimum or no tillage and stubble retention. At each site, the soil had been amended by both
183 incorporation of clay-rich soils from subsoils within pits dug nearby and also by deep incorporation
184 from *in situ* using mechanical means. Trials were sampled at depth intervals to 50 cm after at least 3
185 years following treatments. See Schapel *et al.* (2018) for detailed descriptions of the sites and
186 processes used for clay amendment.

187 Soils were sampled during the non-growing phase as cores from various depths. They did not
188 contain gravel so were ground to pass a <2 mm sieve. Homogenised samples were analysed for OC
189 by the Walkley-Black wet digestion and bulk densities were also determined gravimetrically for
190 known volumes of soil. See Schapel *et al.* (2018) for details of methods.

191 Schapel *et al.* (2018) found that the stock (or reservoir) of OC was significantly and positively
192 correlated ($R^2 = 0.47$) with the stock (or reservoir) of clay when these values were adjusted to an
193 equivalent soil mass (ESM) from bulk density determinations.

194 Results showed that OC stock (or reservoir) in the surface (0-30 cm) of 14 clay-amended soils
195 increased on average by 5.5 Mg C ha⁻¹ (range -1.6 to 14.2 Mg C ha⁻¹) from unamended sands, with
196 the increases largely brought about by increases in clay content. Clay and OC were concentrated at
197 the surface in clay-spread treatments and at depth in delved treatments (Schapel *et al.*, 2018).

198 DISCUSSION

199 Organic carbon in solution is adsorbed by soils, and especially by their most reactive clay
200 fractions, to various extents, depending upon their clay mineral composition. The uptake of OC
201 correlates with the specific surface area of the soil clay fractions (Table 1). Uptake of OC was
202 increased by the presence of iron oxides, as also demonstrated by Kahle *et al.* (2003, 2004). Some
203 organic matter in natural soils is held particularly strongly, and, notwithstanding that a small fraction
204 was even resistant to peroxidation, the oxidation of OC enables further uptake. Others, e.g., Kaiser
205 and Guggenberger (2003), and Mikutta *et al.* (2005), have also found that some OC is held
206 particularly strongly in soils. The strongly-held portion may be located within small pores (McCarthy
207 *et al.*, 2008).

208 Desorption shows that considerable organic matter was easily lost in water (Table 2). OC was
209 held more strongly, i.e., less was released on desorption, after removal of native OC had occurred
210 prior to its uptake. OC was held less strongly when Fe oxides were removed prior to its uptake.
211 Nonetheless, much adsorbed OC was quite labile. Kahle *et al.* (2004), Mikutta *et al.* (2007), and Saidy
212 *et al.* (2013) found that OC adsorbed on pure clay minerals included a substantial portion that was
213 lost easily on desorption.

214 The results of the priming experiments showed that, for all the soils, addition of extra OC led
215 to an enhanced release of CO₂ due to stimulation of microbial activity, hence respiration. Singh *et al.*
216 (2017a) attributed the greater effect of priming in the smectitic soil than in the allophanic soil or the
217 soil dominated by kaolinite and illite to the higher microbial activity which they measured for the
218 smectitic soil. Even so, priming, showing breakdown of SOC, occurred even in the allophanic soil,

219 where OC was held more strongly against desorption than in the other two soils studied. This
220 confirms the results of Finley *et al.* (2018), who found a high rate of decomposition of native OC in
221 soils comprising the short-range order minerals, allophane and ferrihydrite.

222 Field experiments (Schapel *et al.*, 2018) demonstrated that both direct incorporation of
223 exogenous clay addition and 'delving' of *in-situ* clay addition resulted in an increase in SOC content.
224 Increases in SOC were also observed 8 years after additions of subsoil kaolinitic clay to sandy topsoils
225 by delving in Western Australia (Hall *et al.*, 2010). The top 10 cm of soil contained only 1% clay,
226 which increased to 6% clay on delving. There was a resulting increase of 0.2% in SOC. In South
227 Australia, Bailey and Hughes (2012) found that delving soils in 11 sites increased the mean SOC
228 contents of their A2 horizons from 0.3% to 0.7%. Other clayey material can also enable increases in
229 SOC, and wastes from bauxite processing (85% silt, 11% clay and 4% sand) were added to sandy soils
230 in Western Australia (Harper *et al.*, 2012; Churchman *et al.*, 2014). Over a 19-20 year period,
231 addition of this waste led to increases in SOC in the top 30 cm of soil of between 0.1 to 0.65%,
232 resulting in a significant ($P < 0.01$) increase of 11.6 Mg C ha⁻¹ across the sites with a strong ($r^2 = 0.93$,
233 $P < 0.001$) curvilinear correlation between clay content and SOC for the 0-5 cm layer (Churchman *et*
234 *al.*, 2014). This amounts to an annual uptake of 1.9 Mg CO₂-eq. ha⁻¹ year.

235 *Implications for carbon sequestration in soils*

236 Carbon is turned over - and lost from the soil system- through the biological processes
237 needed for soil functioning in agriculture (e.g. Janzen, 2006). Sequestration of C through long-term
238 storage and the use of soil for growing plants appear to be conflicting aims. For example, Bolan *et al.*
239 (2012) concluded that amending soils with biochar, which is more stable than composts, could
240 enhance soil carbon sequestration (long term carbon storage) but this carbon might not be useful for
241 an immediate improvement of soil health, such as for supplying food to soil microorganisms for
242 keeping the nutrient cycling operational. Following Jensen (2006), the most efficient way of

243 extracting extra carbon as CO₂ from the atmosphere may be by using it more rapidly for increasing
244 plant production, hence flow-through of SOM.

245 Increased adsorption which occurred when native C was first removed from all soils (Singh *et al.*
246 *et al.*, 2016) shows that some C had occupied the most reactive mineral sites. The results from
247 desorption (Singh *et al.*, 2016) show that further C occupied less reactive sites in all soils and was
248 easily removed. The effect of priming (Singh *et al.*, 2017a) shows that simple addition of new C to
249 soils may result in the loss of C that is already present. Overall, these results show that simply adding
250 new C to soils that already contain C does not result in additional C that is held strongly, i.e.
251 sequestered.

252 However, the addition of clays to soils can enable the net uptake of C. Furthermore, the rate
253 of incubation of wheat residues added to a sandy soil was retarded by the addition of the clay
254 fractions used in the studies of sorption of DOC (Singh *et al.*, 2019). The addition of clay produces
255 new surfaces for holding C.

256 According to Churchman and Velde (2019, p. 152) "SOC can be associated with almost any
257 secondary (and also altered primary) minerals, but shows a preference for poorly crystalline oxides
258 and also silicates of Fe and Al". If these are rare, SOC binds to phyllosilicates according to their
259 relative surface reactivities, with smectites the most reactive of these. pH also plays a role in
260 governing reactivities of minerals for SOC. Low pHs favor Fe and Al compounds and higher pHs favor
261 phyllosilicates. The type, hence strength, of these associations also vary with pH. Associations
262 between SOC and minerals tend to be inner-sphere, hence stronger, at low pH and outer-sphere at
263 high pH (Kleber *et al.*, 2015).

264 Various functional groups in organic matter can be bound to minerals in layers. Proteins and
265 molecules containing carboxyl groups often occur adjacent to mineral surfaces while molecules
266 containing hydrophobic groups tend to occur in outer layers (e.g., Kleber *et al.*, 2007). The layering
267 of organic molecules around the clay minerals is onion-like. Functional groups with a strong

268 attraction to minerals are likely to form associations with them early in soil development that are
269 long-lasting. Examples are proteins, which are attracted to the negative charges that characterize
270 many clay minerals, especially 2:1 (Si:Al) phyllosilicates via electrostatic bonds through $-NR_3^+$ groups
271 (where R is H or CH_x and $x= 1-3$), and also carboxyl and O/N alkyl groups, which form covalent bonds
272 with Fe oxides that may be free or else associated with phyllosilicates (Schöning *et al.*, 2005).

273 Organic matter most intimately held on mineral surfaces would be held for longer times
274 than that beyond the surfaces. Even so, associations of organic matter with minerals in soils often
275 occur in 3-dimensional entities known as microaggregates. These are distinguished from
276 macroaggregates by size, although the demarcation between the two may be arbitrary and
277 pragmatic (Totsche *et al.*, 2018). In the literature, microaggregates are defined as being variously in
278 the $< 250 \mu\text{m}$ (Totsche *et al.*, 2018), $53-250 \mu\text{m}$ (Beare *et al.*, 1994; Six *et al.*, 1999; Deneff *et al.*,
279 2004) and/or $2-50 \mu\text{m}$ size ranges (Tisdall and Oades, 1982; Paradelo *et al.*, 2016) while Liefeld and
280 Kögel-Knabner (2003) measured their mean weight diameters in a range of soils as from $11.8 - 15.6$
281 μm . Using transmission electron microscopy of thin slices of soil, Chenu and Plante (2006) and
282 Churchman *et al.* (2010) have found many microaggregates involving mineral and organic
283 associations in the size range of $1-5 \mu\text{m}$.

284 In any case, microaggregates provide the basis of the structure of soils, stabilising them
285 against disruption by agricultural practices and erosion. In particular, SOC is stabilised in
286 microaggregates. In microaggregates, organic matter, often occurs within “shells” of the minerals
287 (Chenu and Plante, 2006; McCarthy *et al.*, 2008; Churchman *et al.*, 2010).

288 Organic matter turnover in microaggregates has been estimated variously in at least four
289 different studies as from $\sim 100 - 1000$ years (Churchman and Velde, 2019). Studies of
290 chronosequences (Totsche *et al.*, 2018) have shown that the accretion of newly formed soil
291 components such as microbial residues or hydrous Fe oxides into microaggregates appears to take
292 place within ~ 200 years. Turnover time is slow even in larger, silt-size aggregates (Virto *et al.*, 2010).

293 In truth, gases such as carbon dioxide are not removed directly from the atmosphere by soils
294 or SOM. Generally, they are taken up by plants for photosynthesis leading to their growth and
295 release into the soils. This occurs during the growth of plants, e.g., through root exudates or after
296 their death, by their decomposition. The extra OC added to soils when clay was added to overcome
297 non-wetting (Churchman *et al.*, 2014; Schapel *et al.*, 2018) came about because addition of clays
298 encouraged plant growth, mainly by holding water on their large hydrophilic surfaces for later
299 release to plant roots. In other experiments, additions of clays to light-textured (sandy) soils have
300 been shown to increase plant production. In the laboratory, Churchman and co-workers (unpub.
301 results, 2001) found that addition of 12.5% by weight of foundry waste containing 35% bentonite to
302 a non-wetting sandy soil from South Australia not only overcame its hydrophobicity but trebled dry
303 matter production. In the field, addition of similar foundry waste to a sandy soil in Thailand (without
304 non-wetting problems) also led to increases of up to three times in the dry matter yield of maize
305 (Soda *et al.*, 2006). Pot trials adding 40 t ha⁻¹ to a degraded Oxisol and a sandy Ultisol from northern
306 Australia showed increases in the biomass production of forage sorghum of 3 and 8 times
307 respectively (Noble *et al.* 2001). C contents were not measured in these various experiments, but the
308 increased plant growth in each case would have led to substantial increase in SOC. Wherever excess
309 clay is available locally, increased retention, if not new sequestration of C can occur in sandy soils
310 where sparse plant growth is the norm.

311 Limited sources of clays occur as industrial wastes, with, e.g. ~ one million tonnes of
312 “bleaching earths”, which are acid-activated bentonites being released as wastes from cooking oil
313 manufacture world-wide each year (Crossley, 2001), as well as some from foundries. However, huge
314 areas of sandy soils are found world-wide (> 900 million hectares, according to FAO/UNESCO, 1995)
315 and their augmentation by clays from these sources offers prospects for only limited overall
316 sequestration of carbon.

317 The prospects for the (new) sequestration of C are constrained by 1) the availability of
318 reactive surfaces, and 2) the possible formation of microaggregates. Most surfaces in soils are
319 mineral rather than organic (Mayer and Xing, 2001), with microorganisms and organic matter
320 occupying only a very small fraction (<1%, according to Kleber *et al.*, 2015) of soil surfaces. Selective
321 spots, variously described as “organo-mineral clusters with rough surfaces” (Vogel *et. al.*, 2014, p. 5)
322 appear to occur for the uptake of OC in soils. Etch pits, micropores and cracks on mineral surfaces
323 are likely to constitute such spots (Churchman and Velde, 2019), which are probably related to those
324 for microorganisms, and include the rhizospheres of plants (Hinsinger *et al.*, 2009). Hence the
325 opportunities for uptake, and, especially, for strong uptake, i.e., sequestration, of C are quite limited.

326 C sequestration takes place in microaggregates but takes a long time, so that their formation
327 does not provide an immediate solution to the removal of newly released greenhouse gases from
328 the atmosphere. Nonetheless, formation of microaggregates takes place within macroaggregates
329 (Oades, 1984; Golchin *et al.*, Six *et al.*, 1999; Balesdent *et al.*, 2000) and the formation of these is
330 favoured by vigorous plant growth (Six, 2004). Although strong plant growth may not sequester C in
331 the long term, its promotion could remove some C from the atmosphere. The removal of C by
332 isolation in larger, macroaggregates is transitory (Balesdent, 1996; Puget *et al.*, 2000; Six *et al.*,
333 2002), but continues as long as the strong plant growth is maintained. The process “buys time” while
334 possible new technologies may become available, as Minasny *et al.* (2017) claimed for the “4 per
335 mille” initiative for increasing soil C world-wide. The C isolated in macroaggregates may be
336 transferred into microaggregates and hence truly sequestered, but only in the very long-term.

337 Soil management has often been proposed as the key to increasing the sequestration of C
338 (Lal, 2004, 2014; Minasny *et al.*, 2017). West and Six (2007) suggest step-wise increases in the
339 capacity of soils to sequester C with distinct changes in soil management. The introduction of no-till
340 farming (NT) is a common management strategy, for example, but considerable evidence suggests

341 that NT may only change the distribution of C within the soil profile (Luo *et al.*, 2010), generally
342 towards the surface of profiles.

343 Even so, the experiments reported here, as well as those cited in discussion, involve surface
344 soils. Studies of organic matter in subsoils, e.g. Lorenz and Lal (2005), Chabbi *et al.* (2009) and
345 Salomé *et al.* (2010) have found several characteristics of organic matter in subsoils that suggest that
346 its functioning and stabilisation differs from that in topsoils. For example, (some) subsoil SOC may be
347 very old (several thousands of years old) (Chabbi *et al.*, 2009) and strongly bound to minerals
348 (Chabbi *et al.* 2009), including in structural units (Salomé *et al.*, 2010). Subsoil SOC appears to have
349 high spatial heterogeneity (Chabbi *et al.*, 2009; Salomé *et al.*, 2010) and may not be subject to a
350 priming effect (Salome *et al.*, 2010). Sequestration of C in subsoils may be brought about by the use
351 of plants/cultivars with deep and thick root systems (Lorenz and Lal, 2005). Nonetheless, new C
352 sequestration in subsoils is also subject to considerations of the degree of saturation of mineral
353 surfaces. If fully saturated, they are not good candidates for net sequestration. Certainly, the
354 subsoils in the texture contrast soils studied by Schapel *et al.* (2018) and also others discussed by
355 Churchman *et al.* (2014) proved to be useful for net uptake of C, hence good candidates for C
356 sequestration.

357 Soils have a limited capacity for OC (Hassink, 1997; Stewart *et al.*, 2008). The limits for C
358 sequestration, rather than for overall, maybe partly transitory, uptake of C, should reflect the
359 capacity of reactive mineral surfaces for carbon. Such limits are likely to be somewhat less than the
360 total capacity of soil for C. Following analyses of several results from field trials of the addition of C
361 to soils, Stewart *et al.* (2007) proposed an “effective stabilization capacity” which defines the
362 “maximum sequestration possible with increasing C input level under a particular management
363 scenario”. In this work, we confirm the validity of Stewart *et al.* (2007)’s “effective stabilization
364 capacity” concept, but through the use of laboratory experiments and those of additions of clays to

365 soils in the field. If, or when, the adsorption capacity of its reactive mineral surfaces is reached, there
366 is little – or no- prospect of new sequestration of C from the atmosphere into a soil.

367 CONCLUSIONS

368 The results of recent laboratory experiments on the interaction of DOC with various southern
369 Australian soil clays and of field experiments on the effects on carbon of additions of clays to a
370 Southern Australian soil, together with the literature, have been interpreted in the context of
371 prospects for the sequestration of carbon in soils. .

372 It has been found that the amount of organic carbon (OC) that can be adsorbed by soils depends on
373 their clay minerals, including Fe oxides. However, much of the OC that can be adsorbed by soils can
374 also be lost easily and substantial OC in soils can be lost when more is added (by priming).

375 Nonetheless, some (native) OC is held strongly in soils. OC close to mineral surfaces or within
376 microaggregates is held most strongly. Even so, only very limited areas of mineral surfaces allow the
377 uptake of OC, and when reactive surfaces are occupied, new C will not be held strongly.

378 It was found that addition of clay provides new surfaces for the uptake and retention of C.

379 Overall, it can be concluded that net new sequestration of C in soils cannot be achieved by simply
380 adding C to unamended (top) soils, although net C sequestration may occur in subsoils. On the other
381 hand, increasing the isolation of OC within macroaggregates by enhancing plant growth, although
382 transitory, may provide a more effective method of decreasing atmospheric C than by its
383 sequestration.

384

385 CONFLICTS OF INTEREST

386 On behalf of all authors, the corresponding author states that there is no conflict of interest.

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599

LIST OF TABLES

600 Table 1. Maximum amounts (mg g^{-1}) of DOC adsorbed in $0.01 \text{ Ca}(\text{NO}_3)_2$ by the clay fractions of soils
601 with various dominant minerals in relation to their contents of C and Fe and their specific surface
602 areas (SSAs) calculated using the BET equation for adsorption by N_2 gas (data from Singh, 2016)

603

604 Table 2. Percent desorption of DOC in water after its adsorption in $0.01 \text{ Ca}(\text{NO}_3)_2$ at various loadings
605 from the clay fractions of soils with different dominant minerals sequentially treated to remove
606 native C and Fe (data from Singh, 2016)

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608 **Table 1. Maximum amounts (mg g^{-1}) of DOC adsorbed in $0.01 \text{ Ca}(\text{NO}_3)_2$ by the clay fractions**
 609 **of soils with various dominant minerals in relation to their contents of C and Fe and their specific**
 610 **surface areas (SSAs) calculated using the BET equation for adsorption by N_2 gas (data from Singh,**
 611 **2016)**

Dominant clay type	Treatment	C (mg kg^{-1})	Total Fe	SSA ($\text{m}^2 \text{g}^{-1}$)	Max. adsorbed (mg g^{-1})	
Kaolinite + Illite	None	15	1.6	55	22.0	
	“	Removal of C	4	-.*	69	29.2
	“	Removal of C + Fe	2	-.*	39	16.9
Smectite	None	23	1.2	74	50.2	
	“	Removal of C	3	-.*	93	77.5
	“	Removal of C + Fe	1.2	-.*	60	36.9
Allophane	None	130	7.5	119	101	
	“	Removal of C	32	-.*	140	123.5
	“	Removal of C + Fe	15	-.*	90	71

612 * Not determined

613

614 **Table 2. Percent desorption of DOC in water after its adsorption in 0.01 Ca(NO₃)₂ at**
 615 **various loadings from the clay fractions of soils with different dominant minerals sequentially**
 616 **treated to remove native C and Fe (data from Singh, 2016)**

Dominant clay type	Treatment	Loading of DOC for adsorption (mg g ⁻¹)					
		25	50	75	100	150	200
		Percentage of adsorbed C that was desorbed in water					
Kaolinite + Illite	None	16	16	20	29	34	34
	“	12	14	17	23	28	28
	“	18	17	23	32	40	40
Smectite	None	6	12	17	23	28	28
	Removal of C	10	10	12	17	23	23
	Removal of C + Fe	12	12	15	22	31	31
Allophane	None	7	11	14	16	24	24
	Removal of C	6	7	7	12	15	15
	Removal of C + Fe	12	12	15	20	27	27

617