



Sedimentary talc in Neoproterozoic carbonate successions

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12 Abstract

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13 Mineralogical, petrographic and sedimentological observations document early 14 diagenetic talc in carbonate-dominated successions deposited on two early Neoproterozoic 15 (~800-700 million years old) platform margins. In the Akademikerbreen Group, Svalbard, talc 16 occurs as nodules that pre-date microspar cements that fill molar tooth structures and primary 17 porosity in stromatolitic carbonates. In the upper Fifteenmile Group of the Ogilvie Mountains, NW Canada, the talc is present as nodules, coated grains, rip-up clasts and massive beds that are 18 19 several meters thick. To gain insight into the chemistry required to form early diagenetic talc, we 20 conducted precipitation experiments at 25°C with low-SO₄ synthetic seawater solutions at varying pH, Mg²⁺ and SiO₂(aq). Our experiments reveal a sharp and reproducible pH boundary 21 (at \sim 8.7) only above which does poorly crystalline Mg-silicate precipitate; increasing Mg²⁺ 22 23 and/or SiO₂(aq) alone is insufficient to produce the material. The strong pH control can be 24 explained by Mg-silica complexing activated by the deprotonation of silicic acid above ~8.6-8.7. 25 FT-IR, TEM and XRD of the synthetic precipitates reveal a talc-like 2:1 trioctahedral structure 26 with short-range stacking order. Hydrothermal experiments simulating burial diagenesis show 27 that dehydration of the precipitate drives a transition to kerolite (hydrated talc) and eventually to 28 talc. This formation pathway imparts extensive layer stacking disorder to the synthetic talc end-29 product that is identical to Neoproterozoic occurrences. Early diagenetic talc in Neoproterozoic

30 carbonate platform successions appears to reflect a unique combination of low Al concentrations 31 (and, by inference, low siliciclastic input), near modern marine salinity and Mg^{2+} , elevated 32 SiO₂(aq), and pH > ~8.7. Because the talc occurs in close association with microbially 33 influenced sediments, we suggest that soluble species requirements were most easily met through 34 microbial influences on pore water chemistry, specifically pH and alkalinity increases driven by 35 anaerobic Fe respiration.

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37 Keywords: Proterozoic; geobiology; mineralogy; geochemistry; carbonate; silica

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39 1. Introduction

40 Talc, Mg₃Si₄O₁₀(OH)₂, is typically interpreted as a high-temperature mineral that forms 41 from hydrothermal alteration or metamorphism of Mg-rich and ultrabasic rocks (Evans and 42 Guggenheim, 1988; Marumo and Hattori, 1999). Lower temperature reactions can also produce 43 talc-bearing assemblages, for example through the weathering of serpentinite deposits (Velde 44 and Meunier, 2008); however, this class of reactions is far less common than its high temperature 45 counterpart. As such, outside of the scattered reports of talc in evaporite or carbonate rich 46 deposits (Bodine, 1983; Braitsch, 1971; Calvo et al., 1999; Friedman, 1965; Millot and Palausi, 47 1959; Noack et al., 1989), talc is rarely reported as a sedimentary mineral. Here we document the 48 unusual occurrence of early diagenetic talc associated with Neoproterozoic (~800-700 Ma) 49 carbonates deposited on two separate platform margins: the Akademikerbreen Group in Svalbard 50 and the Fifteenmile Group in the Ogilvie Mountains of northwestern Canada. The formation of 51 sedimentary talc and its mineralogical precursors requires a specific set of chemical conditions;

52 its presence places tight quantitative constraints on Neoproterozoic ocean chemistry and provides 53 additional insight into the biogeochemistry of marine sediments at that time.

At low temperatures (i.e., less than $\sim 30^{\circ}$ C), the Mg-silicate system is controlled mainly 54 55 by kinetic phenomena (Evans and Guggenheim, 1988; Jones, 1986; Wollast et al., 1968), and so 56 constraints on early diagenetic chemistry are difficult to derive based on thermodynamics alone. 57 To address this problem, we designed a series of experiments to evaluate the effects of changing pH, Mg^{2+} and SiO₂(aq) on the formation of Mg-silicates from low-SO₄ (~2.8 mmol/kg), 58 59 Neoproterozoic-like seawater solutions. When these experiments and companion modeling 60 results are coupled with stratigraphically constrained geochemical data, we are able to posit that 61 the early diagenetic talc in Neoproterozoic carbonate successions reflects a unique combination of low siliciclastic input, near marine salinity and Mg^{2+} , elevated SiO₂(aq), and pH greater than 62 \sim 8.6-8.7. In what follows, we discuss: (1) the sedimentology, geochemistry and mineralogy of 63 64 the Akademikerbreen and Fifteenmile talc occurrences, (2) experimental constraints on Mg-65 silicate formation from modified Neoproterozoic-like seawater solutions, and (3) reports of 66 similar mineral assemblages from other Neoproterozoic successions. Finally, we consider both 67 the specific constraints these data place on the chemistry of waters bathing Neoproterozoic 68 carbonate platforms and why this interval in Earth history may have favored sedimentary talc 69 formation.

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71 2. Geologic Setting

72 Sedimentary talc occurs in Neoproterozoic strata that crop out discontinuously over a 73 distance of more than 140 km in the Coal Creek and Hart River inliers of the Ogilvie Mountains, 74 Yukon (for exact locations see Macdonald and Roots, 2009). The talc is stratigraphically

confined to the lower Callison Lake Dolostone (unit PF2 of the Fifteenmile Group), which consists predominantly of dolomite with interbedded shale (Fig. 1 & 2) (Macdonald and Roots, 2009; Macdonald et al., 2010). A tuff within the lowermost map unit of the Fifteenmile Group provides a maximum age constraint on the talc of 811.51 ± 0.25 Ma; a minimum constraint comes from a 717.43 ± 0.14 Ma quartz-phyric rhyolite flow in the overlying Mount Harper volcanic complex (Macdonald et al., 2010).

81 On a ridge ~10 km to the northwest of Mt. Harper, the lower Callison Lake Dolostone is 82 well exposed and measures 77 m thick (Macdonald and Roots, 2009). Here, these strata rest 83 unconformably on a brecciated surface of PF1 platformal carbonate and consist of a basal 2 m of 84 channelized, sub-rounded quartz gravel conglomerate. These beds are overlain by 17 m of green 85 and red shale and siltstone with dolomite lenses containing microbialite textures. The 86 microbialite is draped with a ~10 cm thick bed of hematite iron formation. Above the varicolored 87 shale and siltstone are an additional 60 m of black shale with laterally discontinuous dolomitic 88 bioherms and abundant microbialite. However, the black shale has a peculiar waxy luster, 89 reminiscent of phosphorite, and XRD analyses indicate that this shale consists almost exclusively 90 of talc. The black talc-bearing shale also contains lenses of redeposited talc rip-up clasts 91 suspended in fine-grained dolomite matrix and abundant black chert nodules. These strata are 92 succeeded gradationally by over 300 m of silicified dolostone (Unit PF3), dominated by 93 stromatolites, microbialaminite, and edgewise conglomerate with cm-sized coated grains, 94 occasional exposure surfaces, and patchy silicification. These facies all suggest deposition in a 95 shallow-water, episodically exposed, marginal marine environment.

96 The talc is stratigraphically confined and laterally persistent along the outcrop belt for 97 over 30 km to the east to Mt. Gibben, and occurs again an additional ~110 km to the east in the

98 Hart River inlier (Abbott, 1997). This lateral extent suggests that talc deposition was at least a 99 basin-wide phenomenon. The Fifteenmile Group has been correlated with the Little Dal Group in 100 the Mackenzie Mountains (Macdonald et al., 2010), which has been interpreted as a marginal 101 marine carbonate bank deposited on a rifted passive margin (Turner and Long, 2008). However, 102 the significant unconformity recently identified at the base of the Callison Lake Dolostone 103 suggests that it may have formed in an additional distinct basin-forming episode (Macdonald and 104 Roots, 2009), perhaps coeval with the Coates Lake Group, which lies stratigraphically above the 105 Little Dal Group. The Coates Lake Group was deposited in narrow, restricted grabens (Jefferson 106 and Parrish, 1989). Although neither evaporitic minerals or pseudomorphs have been identified 107 in the Fifteenmile Group, evaporites are present in both the Little Dal and Coates Lake groups 108 (Jefferson and Parrish, 1989), and we cannot rule out restriction on the platform.

109 In Svalbard, mm to cm scale talc nodules occur within shallow marine carbonates of the 110 Hunnberg Formation, Nordaustlandet, and its lateral equivalents in Spitsbergen, the upper 111 Grusdievbreen and Svanbergfiellet formations (Fig. 2 & 3; Knoll and Swett, 1990). As in 112 Canada, the talc deposits occur in settings ranging from coastal to subtidal environments below 113 storm wave base and can be traced along strike for several hundred kilometers, suggesting a 114 basin-scale phenomenon. Also as observed in Canada, cm-scale hematite layers drape microbial 115 dolomites in the talc-bearing interval. In the Svanbergfjellet Formation, talc forms cm-scale 116 nodules within relatively deep carbonaceous shales and carbonates. More extensive and striking, 117 however, are rounded nodules that formed within molar tooth structures and primary voids in 118 microbialites, in both cases before penecontemporaneous void-filling microspar cements were 119 deposited (Fig. 3; Knoll, 1984, who mistakenly identified the talc nodules as phosphorite).

No radiometric dates closely constrain the age of talc-bearing Svalbard carbonates, but C 121 and Sr isotopic chemostratigraphy and biostratigraphy suggest an age broadly comparable to that 122 of the Ogilvie succession (Halverson et al., 2007; Knoll et al., 1986; Macdonald et al., 2010).

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3. Analytical and experimental methods

125 Our sample suite was selected from the lower Callison Lake Dolostone of the Fifteenmile 126 Group and from throughout the Akademikerbreen Group (Fig. 2; Table 1). The Fifteenmile 127 Group samples, all from outcrop, include dolostone hosting nodular talc, bedded talc deposits, 128 and a small horizon of iron formation. With the exception of some visible surface staining, late 129 stage alteration/oxidation of the samples generally appears minor. Samples from the 130 Akademikerbreen Group are predominantly from the Svanbergfjellet Formation and correlative 131 horizons in the Hunnberg Formation, where nodular talc was identified in the field, but they also 132 include carbonates and shales from units above and below this interval. All Svalbard samples 133 were collected from outcrop and were chosen to capture the range of carbonate lithofacies 134 reported in Knoll and Swett (1990). Again, late stage alteration/oxidation is minor in Svalbard samples, as documented by geochemical screens (e.g., Sr concentration, Mn/Sr, Sr/Ca and δ^{18} O; 135 136 Derry et al., 1989).

137 Mineralogical analyses included X-ray diffraction (XRD) of <2µm (and in some cases, 138 <0.2µm) oriented aggregates from decarbonated samples, and bulk XRD of unfractionated 139 samples. Petrographic analysis and electron microprobe analysis of selected samples were 140 performed using polished, carbon-coated thin sections. Further details of sample preparation and 141 analysis are given in the Supplementary file.

142 Precipitation experiments were performed by addition of SiO₂(aq) to 1 kg batches of synthetic low-SO₄ seawater at various $Mg^{2+}(aq)$ concentrations and pH. Experiments were run in 143 144 water baths at $25 \pm 0.1^{\circ}$ C for a minimum of 4-5 weeks to a maximum of 7 months, depending on 145 conditions. Filtered aqueous samples were collected periodically and analyzed by ICP-AES, and 146 residual solid precipitates were collected at experiment termination, washed and analyzed by 147 XRD, FT-IR and TEM. Selected solid precipitates were extracted and reacted with deionized 148 water at 180°C and 400°C to evaluate mineralogical changes in response to heating/dehydration. 149 Further details of the experimental and analytical procedures are given in the Supplementary file. 150

151 4. Mineralogy and petrography of Neoproterozoic talc

152 4.1 Fifteenmile Group

153 Nodular talc samples from the Callison Lake Dolostone have three components in 154 varying proportions: dolomite, chert, and talc. Detrital siliciclastic input is apparent only in the 155 lower 40 m of varicolored shale and siltstone with interbedded microbial dolomite (Fig. 2). The 156 thin horizon (~10 cm) of iron formation that drapes the microbialite is mineralogically simple, 157 consisting of euhedral hematite grains dispersed in a dolomite matrix, with minor silicification 158 and rare detrital quartz grains. <2µm size fractions reveal a single 7Å kaolin-serpentine group 159 phase with trioctahedral occupancy, consistent with greenalite. The dolomite is present both as 160 primary microbial laminations, coated grains, and as interstitial microspar cement. Talc both 161 drapes and fills laminae within the microbialites and, along with the black chert, forms early 162 diagenetic nodules. Bedded chert is also common. In silicified dolostones of the Callison Lake 163 Dolostone that host nodular talc, carbonate mineralogy consists exclusively of dolomite, with no 164 calcite indicated from powder XRD. Aside from talc, the clay mineralogy of decarbonated 165 samples indicates lesser amounts of either discrete highly crystalline illite and/or mixed layer 166 illite/smectite. The composition of the mixed layer phases is highly consistent from sample to 167 sample, and analyses after ethylene glycol treatment are consistent with R0 (disordered) 168 illite(0.7)/smectite.

169 The bedded talc horizons consist predominantly of black, talc-rich shale. Thin lenses of 170 sub-angular to sub-rounded rip-up clasts of talc supported by very fine-grained dolomite matrix 171 are interbedded with the talc horizons. The interstitial dolomite cements between the reworked 172 talc clasts suggest that at least some of the talc formation predates the formation of these 173 particular cements. The mineralogy of bedded talc samples is also simple; talc overwhelms bulk 174 analyses and clay fractions in abundance. However, two samples of bedded talc revealed 175 subordinate amounts of discrete highly crystalline illite and mixed layer R0 illite(0.7)/smectite 176 (Table 1). In contrast to bulk samples, oriented <2µm size fractions of bedded talc indicate that 177 no mixed layering is apparent. In other words, the talc is present as a discrete mineral phase. No 178 other Mg-silicates were identified in any bulk or <2µm fractions from Fifteenmile Group 179 samples (Fig. 4). Randomly oriented specimens of the <2µm clay fraction further indicate that 180 the talc exhibits a high degree of stacking disorder; it is essentially turbostratic. A number of hkl 181 peaks are significantly modulated, particularly in the 20-25 degree and 30-45 degree range. This 182 type and extent of disordering is present in all of the Fifteenmile talc occurrences.

The carbonate overlying the talc horizons consists of silicified dolomite with microbial lamination, coated grains, and abundant stromatolites. Void-filling cements consist predominantly of dolomite and silica. Decarbonated samples from these strata also yielded talc with lesser amounts of discrete illite and mixed layered illite/smectite. The talc in these samples

187 is distributed throughout the carbonate matrix and not as nodules or other conspicuous
188 sedimentary features otherwise resolvable by petrographic analysis.

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190 4.2 Akademikerbreen Group

191 Akademikerbreen talc occurs as mm- to cm-scale nodules and disseminated particles in 192 the <2µm fraction of the carbonate matrix. Talc nodules fill molar tooth structures, primary void 193 space in microbialites and intergranular space, predating precipitation of penecontemporaneous 194 microspar cement. In general, nodules are petrologically homogeneous; there are no discernable 195 nuclei nor is chemical zoning evident (Fig. 3).

Powder XRD of isolated and cleaned nodules hand picked from Akademikerbreen carbonates shows the same type and extent of layer stacking disorder as in the Fifteenmile talc samples (Fig. 5). That is, the Akademikerbreen talc is also highly turbostratic. XRD patterns again display modulated and/or extinguished *hkl* reflections in the 20-25 and 30-45 degree 2Θ ranges. Aside from talc, the carbonate mineralogy frequently includes ankerite in addition to dolomite and lesser amounts of calcite.

The clay mineralogy of decarbonated Akademikerbreen samples shows more variation among the Mg-silicates. Analyses of decarbonated residues reveal talc in the carbonate matrix in some samples where nodules are not apparent. The talc again shows no evidence of mixed layering or occurrences with other low temperature Mg-silicates such as sepiolite or palygorskite. However, saponite (a Mg-rich trioctahedral smectite) and its burial diagenetic equivalents, mixed layered chlorite/smectite and corrensite, have all been identified (Fig. 4; Table 1; Supplementary file). In addition to Mg-silicates, Akademikerbreen clay fractions contain varying amounts of discrete highly crystalline illite, illite/smectite (most of which corresponds to R1 illite(0.7)/smectite), kaolinite, and, in one sample only, chlorite. In general, the results summarized in Table 1 show that samples containing talc and other associated Mg-silicates consistently lack illite, I/S, and kaolinite. The reverse is also true: aluminous clay mineral assemblages are rarely accompanied by talc.

The Svanbergfjellet, Draken and Backlundtoppen Formations also contain discrete siliciclastic horizons that represent detrital pulses into the carbonate platform (Butterfield et al., 1994; Knoll and Swett, 1990). The clay mineralogy of these horizons is dominated by illite and kaolinite in the <2µm fraction; the <0.2µm fraction is composed almost entirely of discrete illite, with significant defect broadening.</p>

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4.3 Fifteenmile and Akademikerbreen Group Mg-silicate occurrences: Summary

Sedimentology and petrology indicate that talc formed during early diagenesis, perhaps penecontemporaneously, in both the Akademikerbreen and the Fifteenmile Groups. In both environments the clay mineralogy also indicates that talc generally occurs in environments that received low background siliciclastic flux, and where alumino-silicates are present, talc has not been identified.

227 Saponite, chlorite/smectite and corrensite in Akademikerbreen carbonates, all indicate 228 that saponite formed during early diagenesis and was transformed to chlorite-rich mineral species 229 upon burial and late-stage alteration. Because there is little evidence for mafic input to the 230 Akademikerbreen platform, saponite formation suggests that the initial deposition of smectite 231 (originally as a dioctahedral smectite such as montmorillonite), or some other detrital precursor,

initiated a recrystallization/transformation to trioctahedral saponite, likely in response tofavorable aqueous chemistry associated with the platformal carbonates (discussed below).

Samples from the Fifteenmile Group, on the other hand, yield very little decarbonated residue and show little variation in background siliciclastic component; they are dominated by highly crystalline illite (muscovite) and illite/smectite. Together this implies very low background detrital fluxes and negligible input of terrigenous clay beyond detrital muscovite.

The sedimentology, petrography and mineralogy of these deposits are indicative of unusual aqueous chemistry associated with platformal carbonates that facilitated the early diagenetic precipitation of Mg-silicates, whether talc or a precursor phase, from solution. What drove these reactions during Akademikerbreen and Fifteenmile Group deposition?

242

243 5. Constraints from Mg-silicate precipitation experiments

244 To better understand what chemical environment is required to initiate the precipitation 245 of Mg-silicates, we performed a systematic series of experiments interrogating the role of 246 SiO₂(aq) and pH on talc formation. We incubated one liter batches of standard, "modern" 247 synthetic seawater at various SiO₂(aq) concentrations and pH, monitoring the bottles for both 248 precipitation of solid phases and changes in aqueous chemistry. In certain experiments, a visible 249 precipitate formed within the first 48 hours. The precipitate often clouded the solution before 250 flocculating and, eventually, settled to the bottom of the reaction vessel. In other experiments, no 251 precipitate was formed over > 7 months.

When the experimental conditions favored precipitation, XRD analyses on randomly oriented powders showed broad and poorly resolvable *hkl* bands and no 001 reflections (Fig. 5). All XRD analyses showed weak bands at d = 1.55 to 1.52Å, indicating the presence of

trioctahedral layers, and all *hkl* bands were reproducible from experiment to experiment, displaying little variation in their relative intensity and *d*-spacing. In only one case, the experiment conducted at the highest pH, a precipitate gave broad low-angle reflections corresponding to *d*-spacings of 28.7, 13.9 and 10.4 Å. With the exception of the highest pH experiment, the structure of the precipitate is both insoluble and reproducible from experiment to experiment. Perhaps most striking, however, is the observation that the precipitate only formed at a pH of 8.66 and above, implicating pH as a major control on Mg-silicate precipitation.

262 Although XRD data show poorly crystalline products, FT-IR spectra indicate that the 263 experimental precipitates are almost identical to talc. The close correspondence between Mg-O 264 vibrations from the precipitates and lattice vibrations from talc (including the diagnostic 535 cm⁻ 265 ¹ (Mg-O)-OH feature (Farmer, 1974; Zhang et al., 2006)) reflect a similar bonding environment for Mg (Fig. 6). Absorptions at ~ 1020 cm⁻¹ show the same bonding arrangement for SiO₄ 266 267 tetrahedra: a 2:1 layered configuration of tetrahedral silicate layers and octahedral MgO₆ layers (Fig. 6). A trioctahedral occupancy (consistent with XRD peaks at 1.55-1.52 Å), manifested by 268 the OH-stretching vibration at \sim 3676 cm⁻¹, is also evident for both the precipitates and for talc 269 270 standards. However, the lack of an observable basal 001 reflection in oriented XRD patterns 271 suggests that there is little to no stacking order between TOT sheets, perhaps due to variable interlayer/surface hydration (consistent with absorptions at ~1630 and ~3400 cm⁻¹ indicating 272 273 appreciable bound molecular H₂O). Taken together, XRD, FT-IR and TEM data (shown in the 274 Supplemental File) indicate the experimental precipitates are simply a hydrated, disordered form 275 of talc; they are composed of 2-dimensional talc TOT sheets not neatly stacked as in true talc, 276 but organized similar to a pile of bricks loosely cemented with water of hydration.

277 To ascertain the effects of mild heating on poorly crystalline Mg-silicate precipitated 278 from modified seawater solutions, we also performed experiments involving the hydrothermal 279 treatment of poorly crystalline Mg-silicate (the products of precipitation experiments) in the 280 presence of deionized water. After 4 weeks of reaction at 180°C, the poorly crystalline Mg-281 silicate transformed to kerolite (Mg₃Si₄O₁₀(OH)₂ $\cdot n$ H₂O), a distinct mineral phase and hydrated 282 structural analog of talc (Fig. 5). The final product exhibited an apparent d(001) of 9.9 Å and *hkl* 283 peaks that correspond to the kerolite structure discussed in Brindley et al. (1977). Combined 284 XRD, FT-IR and TEM data indicate that simple dehydration -- in this case, in response to 285 temperature increase - drives the transformation from poorly crystalline Mg-silicate to the 286 mineral kerolite. Further heating at elevated temperature or over long timescales in turn causes 287 additional dehydration and experiments conducted at 400°C and 1kbar show that this dehydration 288 drives the transition from kerolite to talc (and minor cristobalite) (Fig. 5). The talc produced 289 from this process exhibits significant turbostratic ordering and powder diffraction data show 290 similar peak modulation to Neoproterozoic talc samples discussed above.

291

292 6. Discussion

293 6.1 Mg-silicate system at low temperature

Present-day seawater is supersaturated with respect to crystalline talc, yet modern and even ancient environments lack authigenic talc (Fig. 7). Like in the case of carbonate precipitation, there must exist a kinetic barrier to talc precipitation; a barrier that we can address through experimental interrogation. Our data and ensuing discussion allow talc precipitation to be reconstructed in an environmentally tuned, step-wise fashion. 299 Previous experimental work has targeted talc precipitation from modern-like seawater, 300 and presents contrasting results to those described herein. Wollast et al. (1968) spiked filtered 301 modern seawater with sodium metasilicate (Na₂SiO₃:9H₂O) at 25°C and precipitated a Mg-302 silicate phase at pH > 8.3. The Mg-silicate was described as "poorly crystalline sepiolite" on the 303 basis of FT-IR and XRD -- a phase not identified in any of our experiments. In addition, the 304 apparent solubility calculated from their experiments is lower than the experiments described 305 here (Fig. 7), implying a lower degree of supersaturation needed for precipitation. Upon closer 306 inspection, however, the Mg-silicate precipitated by Wollast et al. (1968) is not inconsistent with 307 our results. In fact, their FT-IR and XRD analyses also lack the most diagnostic features of 308 sepiolite, including the 12 Å and 7.5 Å diffraction peaks and a Si-O feature in FT-IR attributed to 309 Q₃ bonding of silica tetrahedra in chain silicates (Russell and Fraser, 1994). However, their 310 material also appears to exhibit FT-IR features that ours does not.

311 Aside from the identity of the initial precipitate, there also remains a clear discrepancy in 312 the literature in the aqueous conditions needed to initiate precipitation. To address this, we 313 duplicated some of the experiments conducted by Wollast et al. (1968) using a synthetic 314 "modern" seawater solution at elevated $SiO_2(aq)$. The only difference is the nature of the silica 315 source: TEOS solutions in experiments described here versus sodium metasilicate solutions in 316 Wollast et al. (1968). The results show that Mg-silicate formation does not occur until higher 317 levels of supersaturation than those reported by Wollast et al. (1968) and that the initial 318 precipitate is a known kerolite precursor. We argue that the difference in silica source may well 319 be critical. We have taken precautions to ensure that $SiO_2(aq)$ added to seawater solutions was 320 equilibrated in the monomeric form prior to the addition of MgCl₂ (Dietzel, 2000; Iler, 1979). In 321 contrast, sodium metasilicate solutions are known to polymerize rapidly and form colloidal

322 material upon dilution (Iler, 1979), perhaps in response to decreasing silica levels and/or rapid 323 changes in pH associated with their preparation. Thus, it is possible that the addition of metasilicate solutions to seawater solutions co-precipitated silica with Mg^{2+} in a way that may 324 325 not be truly representative of natural surface environments. This is a complication avoided in 326 our experiments through the use of TEOS as a $SiO_2(aq)$ source.

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6.2 Mg-silicate formation: pH as a master variable

To assess what special combination of Mg^{2+} , $SiO_2(aq)$ and pH results in Mg-silicate 329 precipitation, we ran experiments varying Mg^{2+} concentrations (by 10 times over modern 330 331 seawater) and at variable pH (Fig. 7). As discussed above, raising SiO₂(aq) alone was 332 insufficient to drive precipitation. In fact, our experiments identify pH as the primary control on 333 Mg-silicate formation. At a pH_f of 8.0, we observed no precipitation over the course of the experiment. Within error, $SiO_2(aq)$ levels remained unchanged for > 5 months with no sign of 334 precipitation. In contrast, the same Mg^{2+} and $SiO_2(aq)$ at elevated pH resulted in the rapid 335 336 precipitation of Mg-silicate on the timescale of days, suggesting that despite exceedingly high Mg²⁺ levels, pH appears to be the primary control on Mg-silicate formation from seawater-337 338 derived solutions.

339 Solution chemistry at experiment termination tells a similar story. Relationships between 340 $SiO_2(aq)$ loss and pH_f reveal a sharp speciation boundary for SiO₂(aq) at ~8.6-8.7 (Fig. 8). 341 Below this threshold, SiO₂(aq) is unchanged within analytical error (compared to "blank" 342 experiments conducted in the absence of MgCl₂). At SiO₂(aq) concentrations between 343 amorphous silica and quartz saturation, the dominant silica species in seawater is monomeric silica; $H_4SiO_4^{0}$. As pH is increased above ~8.5, $H_4SiO_4^{0}$ deprotonates, favoring $H_3SiO_4^{-}$ (Fig. 9). 344

345 As a result, at these higher pH values, the concentration of a Mg-silica complex (MgH₃SiO₄⁺) 346 increases sharply. The MgH₃SiO₄⁺ species alone is probably not representative of all possible 347 aqueous Mg-silica complexes in solution, especially given the complexity of silica speciation 348 revealed by NMR (Felmy et al., 2001). Nevertheless, the stability constant for this species 349 (derived from actual potentiometric measurements of SiO₂(aq) and Mg-bearing solutions at 350 various pH and high salinity) shows that regardless of actual species distribution, the pH dependence of Mg-silica complexing is well represented by MgH₃SiO₄⁺ (Santschi and Schindler, 351 352 1974).

353 It is possible that at low pH, the rate of crystallization is too slow to be observed. 354 However, the idea of a minimum pH for Mg-silicate formation is well supported by previous 355 synthesis experiments. For example, Siffert and Wey (1962), using dilute MgCl₂ solutions and 356 monomeric SiO₂(aq), precipitated "true" sepiolite at pH >8.5. At pH >9, talc and stevensite 357 formed. Abtahi (1985) and La Iglesia (1978) obtained similar results. The same trends have been 358 observed by workers synthesizing "Mg-silicate hydrate" for industrial applications, with some 359 citing a canonical pH for "Mg-silicate scale" formation of ~8.5 (Brew and Glasser, 2005a; Brew 360 and Glasser, 2005b; Delacaillerie et al., 1995; Mizutani et al., 1990; Packter, 1986; Strese and 361 Hofmann, 1941; Takahashi et al., 1994; Wei and Chen, 2006).

 $MgH_3SiO_4^+$ and similar species are, in turn, the solution precursors to Mg-silicate precipitates (Packter, 1986). Theoretical treatments of layer silicate nucleation from solution require that an excess surface energy barrier be overcome by a critical supersaturation of a preexisting solution complex (Carrado et al., 2006). The critical supersaturation barrier to initiate Mg-silicate formation, then, must occur close to pH 8.6-8.7. Below this point, speciation calculations show that aqueous Mg-silicate complexes, while present, are unlikely to be 368 concentrated enough to initiate the nucleation of a silicate phase; deprontonated silica species 369 would be kept to scant amounts in solution, preventing appreciable interaction with Mg^{2+} . This 370 strong kinetic control on Mg-silicate precipitation from seawater is reminiscent of the more 371 familiar problem of CaCO₃ precipitation from modern surface seawater. For both systems, 372 despite a clear thermodynamic driving force, a critical level of supersaturation is required. For 373 Mg-silicates, deprotonation of $H_4SiO_4^0$ at elevated pH appears to be the "switch" that enables 374 Mg-silica complexing, in turn driving Mg-silicate nucleation and precipitation.

375

376 6.3 Burial diagenesis and turbostratic stacking disorder in talc

377 The sedimentary record of authigenic Mg-silicates can be related to solution chemistry 378 only with knowledge of thermal transformations possible during burial diagenesis. Our 379 hydrothermal experiments run with poorly crystalline Mg-silicates precipitated from seawater 380 show that upon dehydration, the material readily transforms to kerolite. Previous work has 381 shown that with continued dehydration, either with time or at increased temperature, kerolite, in 382 turn, transforms to crystalline disordered talc (Mitsuda and Taguchi, 1977). Our heating 383 experiments with kerolite confirm these observations and show that the talc formed by 384 dehydrating kerolite exhibits severe layer stacking disorder identical to the disorder observed in 385 Akademikerbreen and Fifteenmile samples (Fig. 5).

Stacking disorder in talc is not uncommon. In fact, it results from an inherently weak electrostatic charge between 2:1 layers (Giese, 1975). The disorder observed in our samples is best matched by ordered stacking of 2:1 layers by a/3 along one of the pseudo-hexagonal axes, with random displacements of $\pm b/3$ along another (Gualtieri, 1999). It is not clear why this type of disordering results, but as others have noted, it may well be controlled by the pathway of

391 crystallization (in this case, dehydration) rather than solid state transformation (Baronnet, 1992; 392 Kogure and Kameda, 2008). Specifically, the 2:1 layer dehydration mechanism of the low 393 temperature precipitate, which initiates stacking to form kerolite and, in turn, talc, may well exert 394 the most control on the type of disorder that results from this pathway. Nevertheless, this poorly 395 understood feature is present in all Neoproterozoic samples, consistent with low temperature 396 precipitation and subsequent dehydration during burial.

397 Although there are no direct temperature constraints on the samples described herein, 398 Neoproterozoic strata in the Yukon are relatively low-grade for rocks of their age. Regionally, 399 condont alteration indices and vitrinite reflectance from the overlying Paleozoic strata suggest 400 these rocks have not seen more than 140 °C (Van Kooten et al., 1997). A similar thermal history 401 of the Svalbard samples is suggested by paleomagnetic studies in which demagnetization 402 experiments removed a Caledonian thermochemical remnant magnetization at 150 °C (Maloof et 403 al., 2004). Moreover, the orange-brown color of preserved organic matter in the 404 Akademikerbreen Group indicates that the degree of graphitization and maximum burial 405 temperatures were modest. In reconciling these constraints with experimental data discussed 406 above, it is important to note that we have used elevated temperature (up to 400°C) purely to 407 observe the dehydration of poorly crystalline Mg-silicate to kerolite and eventually to talc on 408 convenient laboratory timescales. However, this transition is known to occur in response to 409 lower temperature and longer reaction time (Mitsuda and Taguchi, 1977). Indeed, if the kinetics 410 of the Mg-silicate dehydration reaction were to behave in a similar fashion to other burial 411 diagenetic reactions involving clay minerals, the dehydration of poorly crystalline Mg-silicate to talc could, for example, take as little as 10^5 years at ~80°C (Velde, 1985). 412

414 6.4 Mg-silicate formation on Neoproterozoic carbonate platforms

Taken together, experimental data indicate that low temperature talc formation in association with carbonates requires: (1) elevated $SiO_2(aq)$, between quartz and amorphous silica saturation; (2) sufficient (at least "modern") Mg^{2+} ; (3) near marine salinity; (4) low Al (i.e., a low background detrital flux, see below) and (5) elevated pH of at least ~8.7.

419 One key requirement for talc precipitation that we have yet to discuss is low Al, requiring 420 that detrital fluxes were low during deposition of the talc-bearing intervals in the Akademikerbreen and Fifteenmile groups. As Al^{3+} increases in concentration, talc, a nominally 421 422 Al-free mineral, is no longer stable, being replaced instead by Al-bearing smectites, chlorites, 423 palygorskite or other minerals depending on local pore water chemistry and sediment supply 424 (Jones, 1986; Jones and Galan, 1988; Weaver and Beck, 1977). Low detrital fluxes have been episodic features of carbonate platform environments throughout recorded Earth history, and 425 426 don't place a unique chemical constraint beyond that discussed. Thus, our mineralogical 427 observations of low siliciclastic flux in talc-bearing facies of the Akademikerbreen and 428 Fifteenmile successions helps to explain the distribution of early diagenetic talc in space but not 429 in time.

The requirement of elevated $SiO_2(aq)$ concentration was also readily met in Neoproterozoic seawater. Indeed, prior to the evolution of silica skeletons in sponges, radiolarians and diatoms, Precambrian seawater should have carried higher silica concentrations relative to modern levels (Maliva et al., 2005). Consistent with this expectation, early diagenetic chert in Neoproterozoic carbonate successions, largely from peritidal or possibly mildly evaporitic environments, provides evidence for local pore water enrichment in $SiO_2(aq)$ on shallow water platforms (Maliva et al., 2005). The Akademikerbreen Group abounds with

evidence for early diagenetic silicification (e.g., Fairchild et al. 1991; Knoll, 1984; Knoll and
Swett, 1990). Fifteenmile Group samples are also heavily silicified, but record a more complex
history of silicification (Mustard and Donaldson, 1990).

440 The Akademikerbreen and Fifteenmile Groups were likely deposited on marginal to 441 isolated marine platforms (Knoll and Swett, 1990; Macdonald et al., 2010). Near marine salinity (\sim 35‰) and Mg²⁺ concentrations are consistent with these settings, yet the salinity and Mg²⁺ 442 443 content of Proterozoic seawater is poorly constrained. Oolites in the Akademikerbreen Group 444 provide petrographic and geochemical (high Sr) evidence for an aragonite sea at the time of their deposition, suggesting that contemporaneous seawater may have had Mg²⁺ concentrations 445 446 broadly similar to those observed today (Swett and Knoll, 1989). Although no evaporite minerals 447 or their associated pseudomorphs have been identified in either succession, episodic restriction in 448 the Akademikerbreen and Fifteenmile Groups cannot be ruled out. On the other hand, seawater 449 evaporation should decrease pH (e.g., McCaffrey et al., 1987), which is inconsistent with the 450 stability of talc.

451 Talc formation by meteoric mixing faces a similar constraint, as freshwater influx would 452 tend to drive pH below marine values. Additionally, salinity would further decrease in response 453 to post-depositional meteoric influx. Higher water activity (or lower salinity) favors the 454 precipitation of Mg-silicates of higher hydration, namely sepiolite. Lower water activity (or near 455 normal marine salinity), on the other hand, favors kerolite and/or other 2:1 silicates, and 456 synthesis work reinforces these constraints (Jones and Galan, 1988; Siffert and Wey, 1962; 457 Stoessell and Hay, 1978). Mg-silicate occurrences in peri-marine and lacustrine environments 458 also conform to this trend, so well, in fact, that Mg-silicate assemblages have been used to track salinity fluctuations and the influence of meteoric water in a number of modern and ancient 459

settings (Calvo et al., 1999; Jones, 1986; Stoessell and Hay, 1978; Weaver and Beck, 1977;
Webster and Jones, 1994).

Of all the requirements for early diagenetic talc formation, elevated pH (≥ -8.7) is the 462 463 most problematic, given the difficulty of increasing seawater pH, which is largely buffered by 464 the carbon cycle, to the required level. The more realistic alternative is to increase pH in pore 465 waters via microbial processes. Elevated pH can be achieved through certain microbial reactions that remineralize organic matter, a set of processes that need not have decreased salinity, Mg^{2+} 466 467 and/or SiO₂(aq) levels. Both the Akademikerbreen and Fifteenmile talc deposits are intimately 468 associated with microbial facies: bedded talc drapes occur with microbialaminites and 469 stromatolites in Fifteenmile samples, and both localities host talc nodules within microbialites.

470 Anaerobic respiration is a particularly attractive mechanism for increasing pH. In a 471 Neoproterozoic world with low pO₂ and no macrometazoan bioturbation, anoxia would have 472 been a common feature of early diagenetic carbonate sediments. Given the additional likelihood 473 that nitrate levels were also low (Fennel et al., 2005), ferric iron and sulfate would have been the most energetically favorable oxidants available for organic carbon remineralization, with Fe³⁺ 474 favored over SO_4^{2-} (Canfield et al., 2005). Under conditions where the reactive Fe flux was high, 475 476 (perhaps ultimately sourced from hydrothermal Fe fluxes rather than detrital input in these 477 sediment-starved environments) dissimilatory Fe reduction would have been particularly 478 effective at increasing pH through the following reaction:

479
$$2CH_2O + 8Fe(OH)_3 \rightarrow 2CO_2 + 8Fe^{2+} + 6H_2O + 16OH^{-1}$$

When the reactive Fe pool was kept low, either by exhaustion of reactive Fe³⁺ or by pyrite precipitation (Johnston et al., 2010; Lyons and Severmann, 2006), sulfate reduction would have contributed to a pH increase in solution:

483
$$2CH_2O + SO_4^{2-} \rightarrow 2CO_2 + H_2S + 2OH^{-1}$$

484 Fe-speciation and isotopic analyses of other early Neoproterozoic successions indicate that these 485 two metabolic pathways strongly influenced marine chemistry in both basinal and platform 486 settings, with organic carbon fluxes and variations in basinal Fe and S cycles controlling the 487 balance between them (Canfield et al., 2008; Johnston et al., 2010; Johnston et al., 2009). But 488 regardless of the interplay between different types of anaerobic respiration, both pathways would 489 have resulted in a pH increase without changes in other conditions needed for talc precipitation. 490 Pore waters in microbially-dominated sediments would have been especially susceptible to pH 491 increase driven by anaerobic respiration, where higher sediment to pore-water ratios favored 492 more effective changes in chemistry

493 In addition to the Svalbard and Canadian occurrences described here, sedimentary (oolitic) 494 talc has been reported in association with stromatolitic dolomites in an Infracambrian 495 (Neoproterzoic) succession of the Volta Basin of West Africa (Millot and Palausi, 1959). The 496 authors argue from petrographic analyses that talc formation preceded dolomitization and 497 silicification. Noack et al. (1989) also reported onlitic talc in the Schisto-Calcaire Group of the 498 West Congo Basin. Overlying the Upper Diamictite Formation, Schisto-Calcaire carbonates have 499 strontium isotope values of 0.7073 (Frimmel et al., 2006; Poidevin, 2007), consistent with a 500 Cryogenian age (716-635 Ma). The sedimentology of the Schisto-Calcaire Group has been 501 interpreted to reflect alternately shallow marine and lagoonal conditions, separated from outer 502 subtidal regions of the platform by stromatolitic reefs (Trompette and Boudzoumou, 1988). 503 Noack et al. (1989) interpret the talc to have formed by transformation from stevensite or 504 sepiolite on the basis of thermodynamic considerations. This interpretation is not incompatible 505 with ours; stevensite precipitation is generally only observed from synthesis studies at high pH

and near marine salinity (La Iglesia, 1978; Siffert and Wey, 1962). We have not conclusively identified stevensite in our experiments, but have identified a corrensite-like phase forming with tale at high pH. Corrensite formation at low temperature requires the initial formation of a stevensite-like smectite into which brucite-like layers are precipitated (Reynolds, 1988).

510 The apparent concentration of low temperature talc deposits in mid-Neoproterozic 511 carbonate platforms may find explanation in the superposition of Neoproterozoic-specific 512 environmental conditions on the broader set of requirements (low detrital influx, high silica, low 513 oxygen, sufficient Mg, efficient sulfide scavenging (Ben-Yaakov, 1973)) likely to have 514 characterized Precambrian carbonate platforms in general. Fe speciation chemistry indicates that 515 iron played a more important role in organic remineralization after 800 million years ago than it 516 did earlier in the Proterozoic Eon (Canfield et al., 2008; Johnston et al., 2010). Also, the 517 hypothesized shift from a Mesoproterozoic biosphere with warm, high pCO_2 atmosphere with 518 high seawater DIC to a more glacial-prone later Neoproterozoic world with lower pCO₂ and DIC 519 would further have eased chemical resistance to increases in pore-water pH in Neoproterozoic 520 carbonate platforms (Kah and Bartley, 2004). During deposition of the Akademikerbreen and 521 Fifteenmile Groups, anaerobic respiration in microbial sediment receiving little terrigenous input 522 provided a fortuitous combination of chemistry and depositional conditions to increase pH and 523 precipitate authigenic Mg-silicate at near marine salinity. The subsequent burial of this material 524 would have driven a thermal transformation to kerolite and eventually to turbostratically-525 disordered talc.

Talc is not the only product to result from unusual pore water chemistry on Neoproterozoic platforms. As discussed above, the identification of saponite, an Al-bearing trioctahedral smectite, suggests additional diagenetic reactions. In the absence of kinetic

529 constraints, but instead using thermodynamic considerations based on field data, Weaver and 530 Beck (1977) suggest modern seawater need only increased SiO₂(aq) and/or pH to drive a 531 diagenetic conversion from detrital montmorillonite to saponite. Thus, the same chemical 532 conditions that led to talc precipitation were also capable of driving smectite recrystallization. 533 Bristow et al. (2009) identified saponite (and corrensite) in Member 2 of the Ediacaran 534 Doushantuo Formation and suggested a lacustrine depositional environment. Our data, however, 535 show that early diagenetic modification of Neoproterozoic seawater is also capable of producing 536 saponite. Thus, in the absence of independent sedimentological evidence of non-marine 537 deposition, saponite cannot be taken as evidence for lacustrine deposition in Neoproterozoic 538 successions. Indeed, the common occurrence of saponite in modern lacustrine settings rather than 539 marine may simply reflect decreased seawater silica levels after the radiation of siliceous 540 plankton in addition to lower modern marine pH. This evolutionary innovation alone would have 541 been sufficient to leave modern saline lakes among the few environments able to concentrate and 542 drive saponite formation from a dioctahedral precursor.

543

544 **7. Conclusions**

Sedimentology, petrography and mineralogy all indicate that sedimentary talc reflects distinct aqueous chemistry recorded on two separate Neoproterozoic carbonate platform margins. Precipitation experiments with modified "Neoproterozoic-like" seawater show that elevated SiO₂(aq), normal marine salinity and Mg²⁺ levels, and low Al³⁺ all favor talc formation. However, the switch that enables talc nucleation is elevated pH. Above pH 8.6-8.7, the deprotonation of H₄SiO₄⁰ initiates Mg-silica complexation and nucleation, which leads to the subsequent formation of a kerolite precursor composed of hydrated 2:1 layers with little to no stacking order. The progressive dehydration of this material, driven by low water activity in solution or by burialdiagenesis, leads to the formation of kerolite and eventually turbostratic talc.

554 Our analyses, together with other reports of sedimentary talc through this interval, 555 suggest that the later Neoproterozoic Era seawater unusually favorable for early diagenetic talc 556 precipitation. Although we argue that talc precipitation was occurring within the sediment, 557 contemporaneous seawater must have been specially poised, only requiring reasonable changes 558 to chemistry within the pore-waters. Following from this, we suggest that anaerobic respiration 559 may have provided the alkalinity pump that tipped the scales toward talc formation in soft 560 sediments. The degree to which sedimentary processes altered overlying seawater, neither of 561 which are known uniquely, hinders direct interpretation of paleo-seawater compositions and pH. 562 Further, whether the Neoproterozoic was the first and last time talc formed with marine 563 carbonates at low temperature is unclear, but its identification of multiple continents speaks to 564 the state of the Cryogenian oceans. This study does elucidate that insight into seawater and pore-565 water chemistry is clearly recorded by early diagenetic Mg-silicates. An improved understanding 566 of the kinetic and thermodynamic controls behind their formation will continue to exploit these 567 minerals as recorders of early diagenetic chemistry in modern and ancient marine and lacustrine 568 settings.

569

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- 768

769 Figure captions

Figure 1. Bedded and nodular talc interbedded with microbialite in the Callison Lake Dolostone,

771 Ogilvie Mountains, Yukon.

772

Figure 2. Stratigraphic sections of the Fifteenmile and Akademikerbreen talc-bearing deposits.

Figure 3. Void-filling talc nodules (in outcrop (A) and thin section (B) under cross-polarized
light) formed in dolomitic stromatolite of the Hunnberg Fm, Nordaustlandet, Svalbard. The talc
in (B) (orange-brown) appears penecontemporaneous with void-filling microspar cement.

778

Figure 4. XRD patterns of oriented aggregates of the <2µm fraction extracted from talc-bearing rocks. Black lines indicate scans acquired in the air-dried (Ca-saturated) state and red lines indicate scans acquired in the EG-solvated state. (A) Talc from unit PF2 (U. Fifteenmile Group; F927-58m) (B) Talc, saponite, and chlorite/smectite from the Hunnberg Fm (K2016). (C) Saponite and illite/smectite from the Hunnberg Fm (HU1246). (D) Trioctahedral low-charge corrensite from the Svanbergfjellet Fm (86-G-3).

785

Figure 5. Representative powder XRD patterns of randomly-oriented experimental precipitates formed by (A) low temperature precipitation from modified seawater, (B) heating a low temperature precipitate at 180°C for 4 weeks, (C) heating the precipitate from (B) at 400°C for 4 days. For comparison, a cleaned talc nodule from the Svanbergfjellet Fm is shown (D), indicating extensive layer stacking disorder. "C" = cristobalite; "D" = dolomite.

791

Figure 6. FT-IR spectra of experimental Mg-silicate precipitates (black) in comparison to talc
collected from the Svanbergfjellet Fm (red). (A) Lattice vibration region, (B) Si-O stretching
region, (C) H₂O and CO₃ region, (D) OH stretching region.

795

Figure 7. Solubility diagram showing the equilibrium solubilities of crystalline talc (Jones,
1986), kerolite and sepiolite (Jones and Galan, 1988), "amorphous sepiolite" from Wollast et al.

(1968) and the equilibrium boundary between montmorillonite and saponite from Weaver and Beck (1978). Data points represent *apparent solubilities* calculated from solutions collected upon experiment termination. Experiments that have resulted in Mg-silicate precipitate form an apparent boundary approximately parallel to kerolite and/or sepioilite, but these data do not represent true equilibrium conditions with respect to either of these crystalline phases. Green point represents experiment at elevated Mg and $pH_f = 8.0$ where no precipitate formed.

804

Figure 8. SiO₂(aq) loss as a function of final pH showing the sharp pH control on Mg-silicate precipitation. Process blank experiments conducted with no MgCl₂(aq) indicate that one experiment (pH_f ~9.2) has resulted in SiO₂(aq) loss that cannot be explained by experimental artifacts alone and suggests the formation of a precipitate that was unable to be extracted for analysis.

810

Figure 9. Thermodynamic calculations of $SiO_2(aq)$ species distribution in "Neoproterozoic-like" seawater as a function of pH. The sharp pH control on the MgH₃SiO₄⁺ species represents the effect of pH on potential nucleation of Mg-silicate from seawater at elevated SiO₂(aq). Initial SiO₂(aq) = 60 mg/kg and initial SO_{4T} = 2.8 mmol/kg. Charge balance was satisfied by readjustment of CO₂-HCO₃-CO₃ equilibria in an open system with respect to CO₂(g). All mineral precipitation was suppressed. Figure 1.





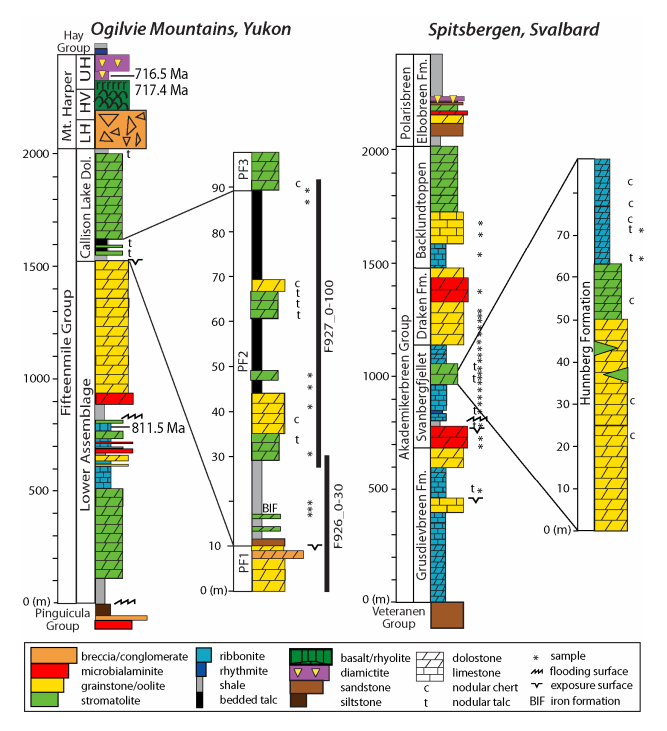


Figure 3.

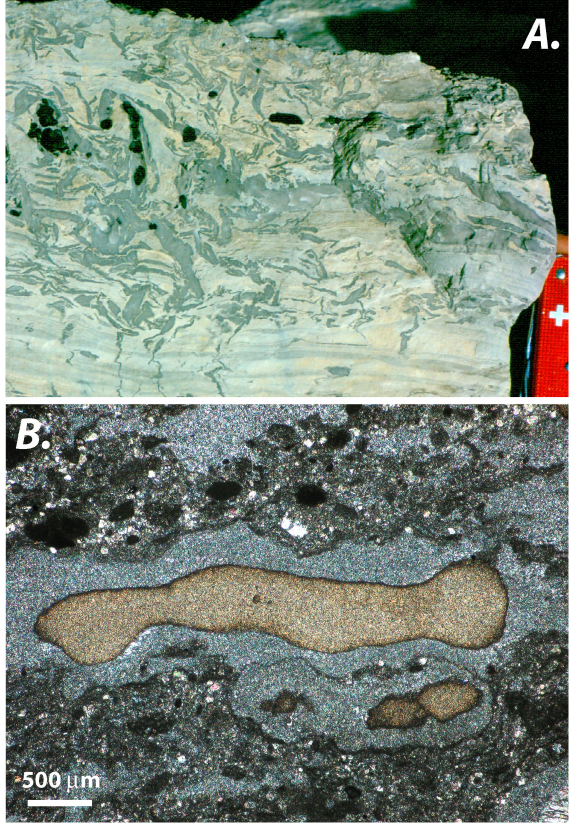
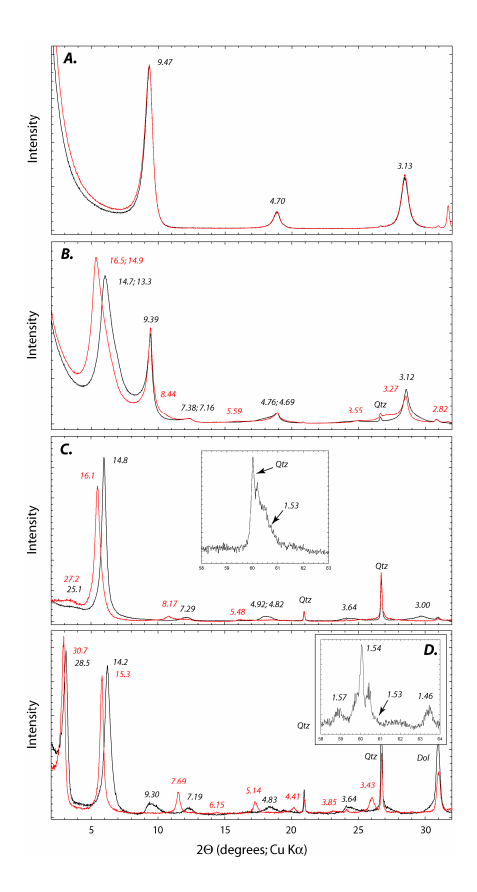
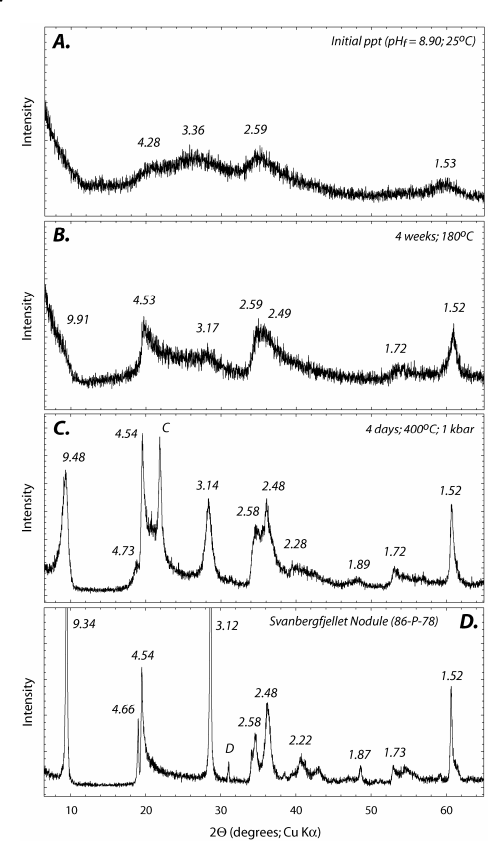
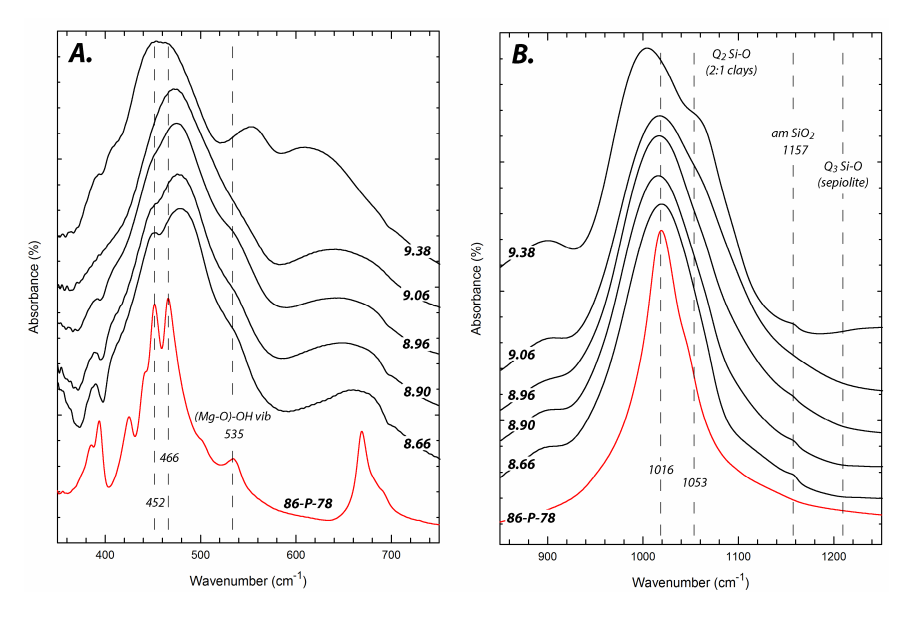


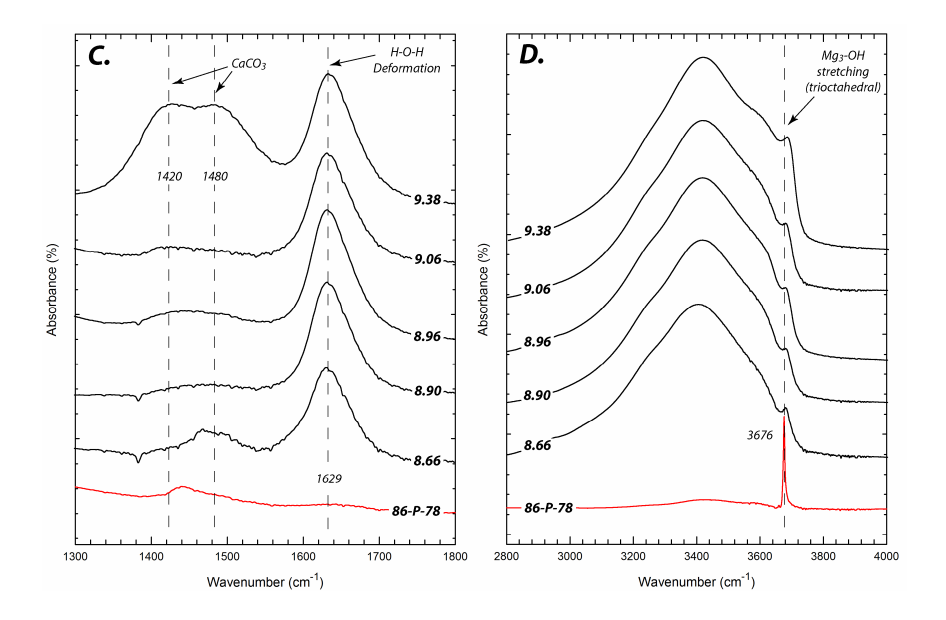
Figure 4.



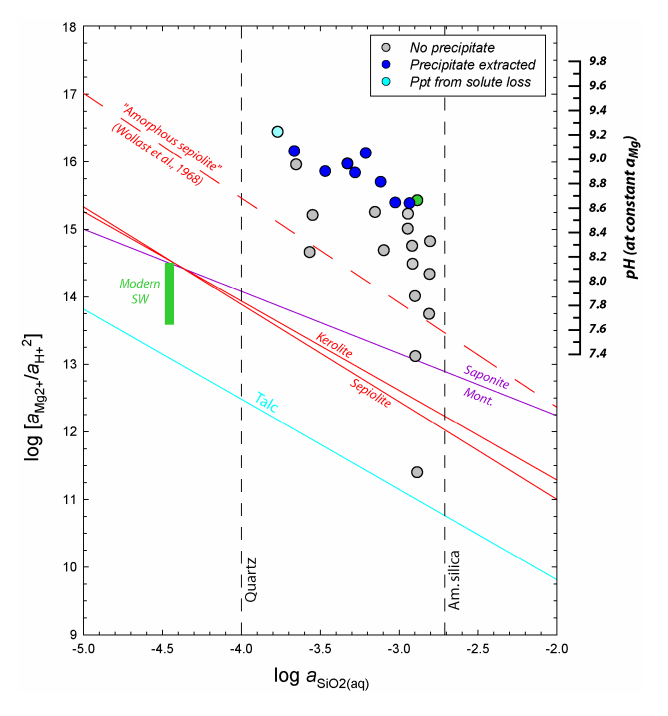




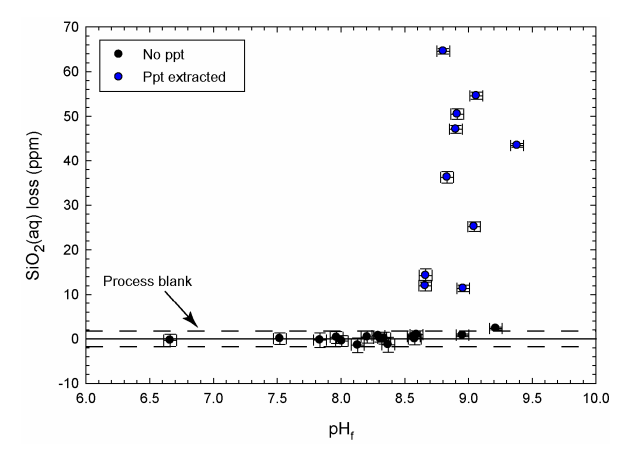




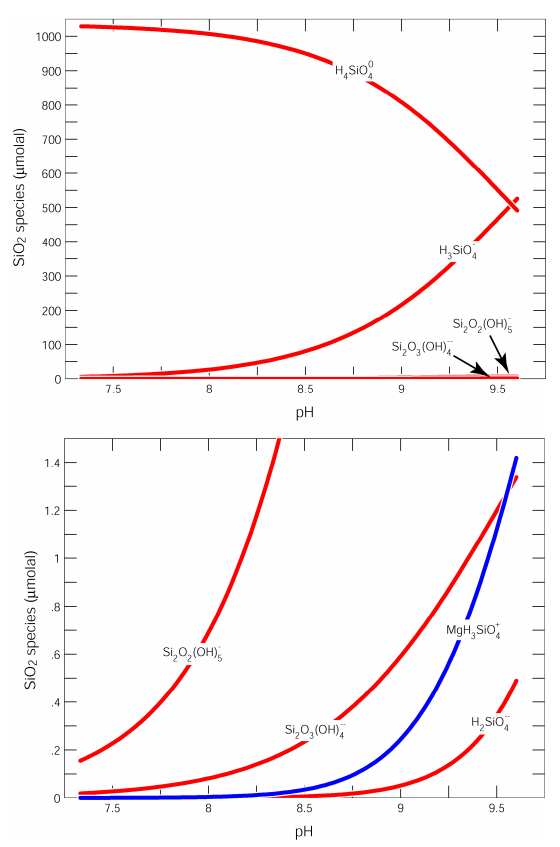












Supplemental Data File (Tosca et al., Sedimentary talc in Neoproterozoic carbonate <u>successions</u>)

3

4 Mineralogical and geochemical analyses

5 Mineralogical analyses were performed on samples that were first cut on all sides to 6 expose fresh surfaces. Samples were processed with a jaw crusher and then by hand in an agate 7 mortar and pestle. To aid carbonate dissolution, some samples were pre-ground in an agate swing 8 mill for <1-2 minutes at 200 rpm. Powders were again hand ground in an agate mortar and pestle, 9 and dissolution was performed by step-wise addition of 0.3 mol/kg acetic acid. Solutions were 10 monitored for effervescence to ensure that pH values did not drop below 4, minimizing Mg-11 silicate structural modification or dissolution (Moore and Reynolds Jr., 1997; Yebra-Rodriguez 12 et al., 2003). Samples were repeatedly rinsed and re-suspended in de-ionized water with sodium 13 phosphate as a dispersal agent and sonicated. The $<2\mu m$ fraction was then separated by gravity 14 settling and transferred to a clean glass slide in preparation for XRD using a filter-peel technique 15 (Moore and Reynolds Jr., 1997). Samples were analyzed as air-dried oriented aggregates and 16 again after ethylene glycol solvation by exposure to ethylene glycol vapor overnight at 60°C.

17 X-ray diffraction of clay separates and bulk samples was performed with a Bruker D-8 18 Advance powder diffractometer with a Cu K α radiation at 40 kV and 30mA and also with a 19 Siemens D5000 diffractometer at 30 kV and 20mA. Divergent slit sizes of 0.2 mm and receiving 20 and anti-scatter slit sizes of 0.6 and 1.0 mm, respectively, were used. For oriented aggregates, 21 samples were analyzed at intervals of 0.03 degrees with 4-8 seconds counting per step. Randomly oriented samples were analyzed from 3-65 degrees at 4 seconds per 0.02° step, with 22 23 $<2\mu$ m size fractions randomly oriented and analyzed from 50-65 degrees at 30 seconds per step 24 to resolve weak 060 peaks that constrain octahedral occupancy. Electron microprobe analysis 25 was performed on polished and carbon-coated thin sections with a Cameca SX-100 electron 26 microprobe equipped with 5 wavelength dispersive spectrometers, capable of analyzing elements 27 from B to U.

28

29 *Experimental procedure*

30 The source of aqueous silica used in the experiments was a 99.9% reagent-grade 31 anhydrous tetraethoxysilane (or TEOS) solution, which, upon contact with water, rapidly hydrolyzes, producing SiO₂(aq) and a small amount of residual ethanol. Although the hydrolysis of organo-silica complexes is rapid, there is a well-documented lag period during which the initially introduced SiO₂(aq) de-polymerizes and reaches equilibrium with the surrounding solution in mostly monomeric (i.e., H₄SiO₄) form (Dietzel, 2000; Iler, 1979). With this in mind, all SiO₂(aq) introduced in the experiment was given enough time to de-polymerize and equilibrate in the solution before Mg was introduced as MgCl₂(aq).

Aqueous samples (1 mL) were periodically extracted from the experiments and immediately filtered with 0.2 µm syringe-driven nylon filters, acidified and stored for analysis. Solution samples were analyzed for major element chemistry using a Varian Vista Pro simultaneous ICP-AES. Calibration was performed with eight standard solutions bracketing expected sample concentrations. In addition, two external quality control standards (matrix matched) were periodically analyzed with samples to determine external precision (0.44% relative standard deviation for Mg and Si, and 0.69% for Ca) and accuracy (0.57-1.03%).

45 At experiment termination, remaining solution was filtered through a 0.2 µm nylon filter 46 membrane to extract any solid precipitate that formed over the course of the incubation period. 47 Solid precipitates were dried overnight at 50° C and prepared as oriented aggregates and 48 randomly oriented samples for XRD analysis. Solid samples were also analyzed by FT-IR using 49 a Bruker IFS 66v infrared spectrometer. IR measurements were collected on KBr pellets with a sample:KBr ratio of 1:300. Measurements were collected from 350-5000 cm⁻¹ at 1 cm⁻¹ 50 51 resolution using a DTGS detector with a KBr window and beamsplitter. Selected samples were 52 prepared for TEM analysis by dispersion in ethanol and deposition on Cu grids.

In addition to experiments run at 25°C, selected solid precipitates were loaded in Teflon vessels with deionized water at a solid:water ratio of 1:75. These experiments were run in hydrothermal bombs at $180 \pm 1^{\circ}$ C for 2-4 weeks to simulate the effects of burial and heating on seawater precipitates over the course of burial diagenesis. Higher temperature experiments were run with precipitate and deionized water (at a solid:water ratio of 1:75) sealed in Au tubes and loaded into a cold-seal hydrothermal apparatus. Experiments were run at 3-4 days at 400°C and 1 kbar hydrostatic pressure.

60

61 Thermodynamic speciation calculations

62 Thermodynamic speciation calculations for SiO₂(aq)-bearing seawater were performed 63 with Geochemists Workbench using a Pitzer-based method for ion activity coefficient 64 calculation based on models developed by Harvie et al. (1984) and Marion and Farren (1999) for 65 major seawater components. For silica speciation in solution, we included H₄SiO₄ dissociation 66 constants from Hershey and Millero (1986) and Pitzer coefficients for SiO_2 species calculated by 67 Felmy et al. (2001). We also included aqueous silica complexes from Felmy et al. (2001) and 68 Santschi and Schindler (1974). Speciation calculations discussed in the text were conducted as 69 follows: initial low-SO₄ (2.8 mmol/kg) Al- and Fe-free seawater compositions were equilibrated 70 with 60 mg/kg SiO2(aq) and a CO₂-containing atmosphere at a pCO₂ to yield an initial pH of 7.25 (log fCO₂ = -1.0). CO₂ fugacity was then decreased while maintaining equilibrium with 71 72 seawater by pH adjustment until the final seawater pH reached ~9.7 at a log fCO₂ = -4.0.

73

74 Details of clay mineral identification

Saponite was identified by a strong basal 001 reflection in air dried oriented aggregates which shifted to 16.1 Å after ethylene glycol treatment. XRD analyses of randomly oriented $<2\mu$ m powders revealed a 060 peak at 1.529Å, corresponding to trioctahedral occupancy. The saponite was found only in samples from the Hunnberg Formation. One sample dominated entirely by finely laminated dolomicrite and abundant molar tooth structures yielded almost pure saponite (with a small amount of R1 illite(0.5)/smectite) after decarbonation.

81 Corrensite, a mixed layered regularly ordered chlorite(0.5)/smectite was identified in one 82 sample from the Svanbergfjellet Formation (86-G-3). An intense superstructure 001* reflection 83 in both the air dried and ethylene glycol-solvated states, along with a rational series of 00*l* 84 reflections upon ethylene glycol-solvation, were sufficient to unambiguously identify corrensite 85 in this sample. The corrensite is of the trioctahedral low-charge variety, indicating that the 86 chlorite component is mixed with trioctahedral saponite.

Figure 4B in the manuscript shows an XRD pattern of the $<2\mu$ m decarbonated residue from sample K2016 (Hunnberg Fm). Talc can be clearly identified from the reflection at 9.37Å. However, the broad peak centered at ~6 degrees 2 Θ contains two reflections: 14.7 Å and 13.3 Å in the air-dried state, and at 16.5 Å and 14.9 Å in the EG-solvated state. From high angle XRD scans of random powders, an 060 peak is present at 1.528 Å, which, combined with reflections at 14.7 and 16.5 Å in air-dried and EG-solvated, respectively, can be identified as saponite. The 00*l* 93 reflections of the additional phase are all well-matched by R1 chlorite(0.75)/smectite. Calculated 94 X-ray diffraction patterns of this phase (using the program NEWMOD; (Reynolds Jr. and 95 Reynolds, 1996)) correspond precisely with observed reflections throughout the 00*l* series. Such 96 a mixed-layer phase is relatively uncommon in natural samples; corrensite is by far the most 97 common chlorite/smectite species, but other forms of chlorite/smectite mixed layering have been 98 reported in association with carbonate and/or evaporite deposits (e.g., (Hillier, 1993)).

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131	Figure 1. Comparison of oriented <2mm fraction (EG-solvated) of sample K2016 from the
132	Hunnberg Formation, Nordaustlandet, Svalbard (A) with (B) a simulated one dimensional X-ray
133	diffraction pattern (calculated by NEWMOD) for a physical mixture of: 67% talc, 27% tri-
134	octahedral smectite (saponite), and 7% R0 tri-tri chlorite 0.7/tri-smectite.
135	
136	Figure 2. Transmission electron micrograph of poorly crystalline Mg-silicate precipitated at a
137	pH of 8.901 (with selected area electron diffraction pattern shown in inset). Photomicrographs of
138	experimental precipitates show a crinkly morphology at the nanometer scale, and in some
139	regions, the development of hexagonally-shaped particles. Some regions of the precipitate
140	exhibit observable patterns in selected area electron diffraction implying crystalline order at the
141	nanometer scale. The smearing of diffracted spots into weak circular patterns is consistent with
142	low stacking order in the z direction.
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Expt ID	рН _f	[SiO ₂] _i (ppm)	[Mg] _i (ppm)	Composition	Precipitate
MgSi_1	6.661	69.70	1182	SW (2.8 mmol/kg SO ₄)	
MgSi_2	7.521	68.80	1181	SW (2.8 mmol/kg SO ₄)	
MgSi_3	7.966	69.88	1178	SW (2.8 mmol/kg SO ₄)	
MgSi_4	8.339	69.17	1178	SW (2.8 mmol/kg SO ₄)	
MgSi_5	8.661	69.39	1180	SW (2.8 mmol/kg SO ₄)	PCMS
MgSi_6	8.901	69.39	1180	SW (2.8 mmol/kg SO ₄)	PCMS
MgSi_7	8.307	45.53	1165	SW (2.8 mmol/kg SO ₄)	
MgSi_8	8.593	42.87	1159	SW (2.8 mmol/kg SO ₄)	
MgSi_9	8.959	43.04	1167	SW (2.8 mmol/kg SO ₄)	PCMS
MgSi_10	9.382	43.81	1157	SW (2.8 mmol/kg SO ₄)	Corr; pc-Talc
MgSi_11	8.294	16.07	1165	SW (2.8 mmol/kg SO ₄)	
MgSi_12	8.570	17.25	1165	SW (2.8 mmol/kg SO ₄)	
MgSi_13	8.953	15.70	1165	SW (2.8 mmol/kg SO ₄)	
MgSi_14	9.215	15.50	1165	SW (2.8 mmol/kg SO ₄)	
MgSi_15	9.062	69.90	1165	SW (2.8 mmol/kg SO ₄)	PCMS
MgSi_16	7.835	84.70	1174	SW (2.8 mmol/kg SO ₄)	
MgSi_17	8.131	85.06	1171	SW (2.8 mmol/kg SO ₄)	
MgSi_18	8.371	85.65	1168	SW (2.8 mmol/kg SO ₄)	
MgSi_19	8.665	85.14	1195	SW (2.8 mmol/kg SO ₄)	PCMS
MgSi_20	8.833	85.14	1200	SW (2.8 mmol/kg SO ₄)	PCMS
MgSi_21	8.912	85.14	1175	SW (2.8 mmol/kg SO ₄)	PCMS
MgSi_22	8.207	68.92	1164	SW (28 mmol/kg SO ₄)	
MgSi_23	8.579	67.75	1165	SW (28 mmol/kg SO ₄)	
MgSi_24	9.044	68.34	1164	SW (28 mmol/kg SO ₄)	PCMS
MgSi_25	8.007	62.94	12900	SW (2.8 mmol/kg SO ₄)	
MgSi_26	8.803	64.88	12900	SW (2.8 mmol/kg SO ₄)	PCMS

Table 1. Experimental conditions

PCMS: Poorly crystalline Mg-silicate **pc-Talc:** Poorly crystalline talc **Corr:** Corrensite (trioctahedral)

Figure 1.

