# 1 Unusual mixed silica-carbonate deposits from magmatic-

# 2 hydrothermal hot springs, Savo, Solomon Islands

3 D.J.Smith<sup>1</sup>\*, G.R.T. Jenkin<sup>1</sup>, M.G.Petterson<sup>1</sup>, J. Naden<sup>2</sup>, S. Fielder<sup>1</sup>, T. Toba<sup>3</sup>, S.R.N.

4 Chenery<sup>2</sup>

5 1. University of Leicester, Leicester, LE1 7RH, UK

- 6 2. British Geological Survey, Keyworth, Nottingham, NG12 5GG, UK
- 7 3. Geology Division, Ministry for Energy, Mines, and Rural Electrification, Honiara,

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8 Solomon Islands
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9 (\*corresponding author djs40@le.ac.uk)

## 10 Abstract

11 The volcanic island of Savo, Solomon Islands, hosts an active hydrothermal system 12 discharging unusual alkaline (pH 7–8) sulphate-rich, chloride-poor fluid, with variable 13 admixtures of Ca-Mg-HCO<sub>3</sub><sup>-</sup> rich fluid. Hot springs and their outflow streams precipitate a 14 variety of deposits, including travertine, silica sinter and unusual mixed silica-carbonate 15 rocks. Travertine fabrics are dominated by ray-crystal calcite, associated with rapid abiotic 16 precipitation from a supersaturated solution. Sinter is produced by evaporation of thermal 17 waters, and downstream samples contain preserved traces of micro-organisms, which 18 potentially acted as templates for precipitation. Trace element chemistry of sinters and 19 travertines includes anomalously high levels of Te, indicating a magmatic origin for a 20 component fluid in the hydrothermal system. Springs are close to or at saturation with both 21 calcite and amorphous silica. Increased contributions from the Ca-Mg-HCO<sub>3</sub><sup>-</sup> end-member 22 favours calcite formation; this fluid is of low temperature origin, and as such is favoured by 23 high rainfall. Mixed samples show cyclical changes between silica and carbonate 24 precipitation, potentially as a result of seasonal variation in rainfall.

25 Supplementary material: Spreadsheet of water and deposit chemical analyses is available at

26 www.geolsoc.org.uk/SUP00000

# 27 [INTRODUCTION]

28 Travertine (CaCO<sub>3</sub>) and siliceous sinter (using the terminology of Renaut & Jones, 2003) are

29 common features around hot springs and the streams they feed. However, mixed silica-

30 carbonate deposits are rare, with only a few recognised worldwide, most notably at Ohaaki,

31 Ngatamariki, and Waikite, New Zealand (Jones *et al.* 1996; Campbell *et al.* 2002; Jones &

32 Renaut 2003a). On Savo (Solomon Islands), a volcano with an active magmatic-

33 hydrothermal system (Smith et al. 2010), highly unusual mixed deposits, accompanied by

34 separate deposits of sinter and travertine are commonly found at springs and thermal streams.

35 Sinter is most commonly associated with discharge of near-neutral, chloride-rich thermal

36 waters (Hedenquist *et al.* 2000) that have reacted with host rocks at temperatures in excess of

37 175°C (Fournier & Rowe 1966; Uysal *et al.* 2011), or more rarely with acid sulphate waters

38 (Rodgers *et al.* 2004) and acid sulphate-chloride waters (Schinteie *et al.* 2007). However, the

39 fluids discharged at Savo are alkaline sulphate fluids with very low chloride (Smith *et al.* 

40 2010). Thus, the alkaline sulphate springs on Savo represent an environment of sinter

41 formation not previously described. It is important to understand the range of environments

42 and fluids that form sinters – and the nature of the sinters that different environments produce

43 – if deposits in the geological record are to be interpreted properly. Furthermore, as the study

44 of surface hydrothermal deposits provides information on the origin of life and mineral-

45 microbe interactions (e.g. Farmer 2000), it is important to understand the range of chemical

46 and physical environments in which they form.

47 In some cases sinter can be the surface manifestation of a mineralised system (Vikre 2007)

48 and therefore an indicator of gold-bearing palaeo-hydrothermal and hydrothermal systems

49 (Hedenquist *et al.* 2000; Uysal *et al.* 2011). As Savo occurs in a region prized for epithermal

50 gold deposits (e.g. Ladolam, Lihir, Papua New Guinea; Carman 2003) might hydrothermal

51 deposits, like those on Savo, be a useful indicator of potential mineralisation?

52 This paper describes the morphology and mineralogy of travertine, sinter and mixed silica–

53 carbonate deposits on Savo, their chemistry and the waters from which they precipitate. The

54 aims of the study are: 1) to identify the mechanisms which precipitate travertine, sinter and

55 mixed deposits; 2) to determine the processes behind changes between carbonate-dominant,

56 silica-dominant and mixed deposit precipitation; 3) to determine the significance of trace

57 element concentrations in the precipitates.

# 58 Background

59 Savo is a recently active volcano in the central Solomon Islands (Fig. 1), dominated by sodic

60 trachyte and mugearite rocks (Smith *et al.* 2009). Eruptive activity (last eruption 19<sup>th</sup> century)

61 has been dominated by dome formation and subsequent collapse to pyroclastic debris currents

62 (Petterson *et al.* 2003). At present, an active hydrothermal system manifests at the surface in

a series of hot springs and fumaroles. Smith *et al.* (2010) discussed the chemistry of the

64 various springs in detail. In brief, high temperature (>80°C) springs are divided into two main

classes – (1) acid sulphate hot springs (pH 2–7) of steam heated origin, and (2) more

voluminous alkaline sulphate springs (pH 7–8), that are dilute, chloride-poor (< 50 mg/l), and

67 sulphate- (> 600 mg/l) and silica-rich (> 250 mg/l). Table 1 summarises the chemistry of

alkaline sulphate type springs. There are a number of lower temperature  $(25-60^{\circ}C)$ 

69 bicarbonate-sulphate springs around the island that discharge fluids with (relative to the

alkaline sulphate springs) lower silica and sulphate concentrations, and higher bicarbonate,

71 Mg and Ca contents (Table 1).

72 Smith et al. (2010) interpreted the hot springs at Savo as the output of a magmatic-

73 hydrothermal system that has been subject to considerable dilution by meteoric-derived

vaters. Weather stations on Guadalcanal record monthly average rainfalls of ~25 cm in the

75 November–April wet season and ~10 cm in the April–November dry season (Solomon

76 Islands Meteorological Service, www.met.gov.sb).The hot spring compositions indicate that

the subsurface of the volcano can be considered an open system, with considerable mixing

78 between end-member fluids leading to intermediate water types. In particular, alkaline

sulphate hot springs of the Crater Wall/ Poghorovorughala area (Table 1) show enhanced

80 bicarbonate, Mg and Ca contents relative to the Eastern/Rembokola springs, indicating

81 mixing of a lower temperature Ca, Mg-rich water into a hotter, hydrothermal silica-rich

82 water. Water-rock equilibration temperature for the high temperature, silica-contributing end-

83 member was estimated to be 260°C (Smith *et al.* 2010).

# 84 Distribution, morphology and mineralogy

#### 85 Rembokola Valley

86 The Rembokola stream, on the east side of the island (Fig. 1), is fed by hot springs on the

87 upper flanks of the volcano. These springs discharge small volumes (<0.5 litres per second)

88 of fluid, and have produced terraced sinter deposits (Fig. 2A). Individual benches are no more

than a few square centimetres in area, thus in the terminology of Fouke *et al.* (2000) are

- 90 considered micro-terracettes. The sinter is highly porous and friable opal-A (X-ray
- 91 amorphous silica, using a Philips PW 1716 X-ray diffractometer; Fig. 2B) with small
- 92 amounts of anhydrite. Small silica-encrusted filaments (<5 µm diameter) are visible in places,
- 93 particularly within pore spaces (Fig. 2C).

94 Downstream, in an area of vigorous hot springs discharging directly into the stream, sinter

95 grows above the water surface on detritus, usually developing into small (1–2 cm) pointed

96 columns (Fig. 3A). SEM analysis of the spicular sinter shows that it is opal-A, often with

97 anhydrite crystals on the top surface (Fig. 3B). This spiculose sinter is similar in appearance

98 to that from Ngatamariki stream, New Zealand, as described by Campbell *et al.* (2002).

99 Downstream of the springs sinter coats and cements sediment and leaf litter in the stream

100 channel, and forms crusts in the stream channel and banks where accumulations are thicker

101 (Fig. 4A). It is typically finely laminated (layers <1 mm; Fig. 4B). Some of the layers are

102 non-porous opal-A (Fig. 4C), others a mixture of non-porous opal-A and hollow, opal-A

103 encrusted filaments up to 5 μm in diameter and 100 μm in length (Fig. 4D, E), aligned

104 orthogonally to the layers. This laminated sinter facies occurs from within a few metres of the

105 springs, to the mouth of the stream.

106 In some locations along the Rembokola stream, mixed silica–carbonate terraces occur above

107 the current water level (upper surface approximately 30 cm above water level at time of

108 observation in dry season, April–November; Fig. 5A). It is unclear from the field relations as

109 to whether the mixed deposits represent high-water level deposits (i.e. wet season, Nov-

110 April) or older deposits on the margins of the stream channel that have been eroded away

111 from the active channel. The exposed surfaces of the mixed show signs of weathering,

112 erosion and lichen colonisation, suggesting that they represent the remnants of older deposits,

113 rather than recent or latest wet season deposits.

114 The mixed deposits consist of alternating layers of opal-A and calcite (XRD). Individual

115 layers are up to 10 mm thick, and unconformities are common (Fig. 5B, D). Carbonate layers

116 are formed from ray-crystals of calcite (Folk *et al.* 1985; Chafetz & Guidry 1999) organised

117 into near-vertical fans (Fig. 5C). Silica layers are mainly non-porous, but in cavities where a

three dimensional view is possible, small filaments are visible (Fig. 5E). Porosity and internal

119 structure appears to have been infilled/ overprinted by continued silica precipitation (Jones &

- 120 Renaut 2003a), again suggesting that the mixed deposits are older than the layered sinters
- 121 (which still preserve abundant filament structures and porosity).

### 122 Poghorovorughala Valley

Deposits form around alkaline sulphate hot springs and in the base of the stream. Deposits are
carbonate-dominated (aragonite and calcite, with possible minor dolomite indicated by XRD)
with opal-A. Distinct depositional facies can be observed (Fig. 6A):

- 126 1) Lobate deposits form adjacent to alkaline sulphate springs, in areas frequently splashed
- 127 and bathed by thermal waters. They typically have smooth, rounded upper surfaces of
- 128 carbonate (microcrystalline aragonite or calcite) with opal-A and minor anhydrite (Fig.
- 129 6A–D). Individual lobes are finely laminated in cross section (Fig. 6B). The cores/ bases
- 130 of the concentric layers often contain portions of the substrate (typically kaolinite,
- 131 formed by steam heated alteration of volcaniclastic host sediments; Fig. 6B). Trigonal
- 132 prisms of calcite are visible in SEM images, typically in sheltered areas between lobes
- 133 (Fig. 6E, G–H). Pyrite and some manganese oxide precipitate on the underside of the
- 134 lobes (i.e. slightly submerged or at the contact with the hot spring water; Fig. 6F).
- 2) Spicular deposits form slightly further from the springs, typically in areas splashed and
  bathed infrequently. The physical appearance is identical to the spiculose sinter near the
  Rembokola springs (Fig. 3), although contains much more carbonate. Spicular growths
  were observed developing on a lobate travertine substrate (Fig. 6A).
- 3) Interlayered silica–carbonate deposits occur in the discharge channels of springs and in
  the stream. The ~3 m high Mound Spring (Fig. 7A) is constructed of layered precipitates
  (based on surface exposure) with micro-terracetted surface texture (Fig. 7B). The layers
  are 5–50 mm thick, and generally pale in colour, although some dark layers do occur
  (Fig. 7C). Dark layers tend to be dominated by opal-A (Fig. 7D); whereas the pale layers
- 144 are composed of ~1 mm long calcite ray-crystals organised into fans that diverge
- 145 upwards (Fig. 7F; mineralogy confirmed with XRD). Minor anhydrite is present, mostly
- 146 within the carbonate dominated layers (Fig. 7G).

## 147 Tanginakulu Valley

148 Travertine occurs in the stream channel along the entire length of the Tanginakulu stream. In 149 relatively flat areas, travertine coats and cements stream detritus; whereas greater thicknesses 150 of layered travertine develop at rapids and waterfalls (Fig. 8A). Layers are finer than those 151 observed in the Poghorovorughala layered travertine (generally <5 mm thick), but the calcite 152 has a similar morphology of elongate calcite ray-crystals in upwards-diverging fans (Fig. 8B– 153 C).

# 154 Analytical methods

155 Bulk samples were powdered in a steel jaw crusher and agate planetary mill. Other samples

- 156 were obtained by drilling out individual layers. At the British Geological Survey (BGS),
- 157 Keyworth, for each sample 0.1 g powder was dissolved in a mixture of 2 ml  $H_2O$  and 2 ml
- 158 aqua regia. HF acid was added to samples to dissolve any silica (1 ml for sinters, and mixed
- samples; 0.2 ml for travertines). Resulting mixtures were heated to 40°C for one hour to
- 160 encourage complete dissolution, then left to dry overnight at 120°C. Dried samples were
- 161 redissolved in a mixture of 1.2 ml  $H_2O$ , 0.4 ml HCl and 0.9 ml HNO<sub>3</sub>, and warmed to 40°C
- 162 for 30 minutes. Following complete dissolution,  $10 \text{ ml H}_2\text{O}$  and  $2.5 \text{ ml H}_2\text{O}_2$  was added,
- 163 followed by a further dilution with 10 ml  $H_2O$ . All  $H_2O$  was 18.2 M $\Omega$  quality. Samples were
- analysed at BGS using an Agilent 7500 series ICP-MS in helium gas collision cell mode to
- 165 minimise spectral interferences. The instrument was calibrated with a series of ISO17025
- 166 traceable multi-element solutions (SPEX CertiPrep<sup>TM</sup>). A series of quality control standards
- 167 independent of the calibration solutions were analysed throughout each run. Precision and
- 168 bias of these chemical solutions was better than +/-10% and typically better than +/-5%. Four
- analyses of BCR-2 reference material (values taken from GeoReM http://georem.mpch-
- 170 mainz.gwdg.de) typically yielded precision and bias better than +/- 15% with notable
- 171 exceptions being Ni bias -36% and Cu bias -18%.
- 172 A subset of bulk powders was analysed at Acme Analytical Laboratories, Canada. Samples
- 173 were crushed and powdered as above, and analysed by ICP-MS following aqua regia
- 174 digestion. Precision and accuracy were estimated by duplicate analysis of laboratory standard
- 175 DS7; precision  $(2\sigma)$  was <5% for all species except Zn, Se, Ag (<15%); Cu (43%) and Au
- 176 (63%). The low reproducibility of Cu and Au indicates a nugget effect with standard DS7.
- 177 Similar values were obtained for repeat analysis (n = 4) of a Savo carbonate. The accuracy
- 178 (mean measured DS7 vs. accepted value) was better than +5% for Mn, V, Zn; -5% for Ag,
- 179 Pb, Se; 10% for As, Au, Ba, Cd; +15% for Sr, Cu; -12% for Sb; and +19% for Te. The full
- 180 dataset is included in the supplementary data.
- 181 Water samples were collected and analysed as per Smith *et al.* (2010).

# 182 **Results**

#### 183 Streams

- 184 The Rembokola stream is fed by alkaline sulphate hot springs in the Toakomata area (the area
- 185 immediately around sample locations a–c, Fig. 1), and has a similar chemistry, with high Na,
- 186 Ca, Si, K, and  $SO_4^{2-}$ , and low Cl<sup>-</sup> (Table 2). Arsenic occurs in concentrations of 60–70 µg/l;
- 187 for comparison, typical seawater concentrations are only 1 µg/l (Cabon & Cabon 2000). The
- 188 chemistry shows no abrupt downstream changes, reflecting the fact that there are no
- 189 tributaries. There are, however, gradual changes to the stream chemistry including a
- 190 downstream decrease in temperature, DIC, Mn, and Si; and an increase in Cl<sup>-</sup>, and pH (Table
- 191 2, Fig. 9).
- 192 The Tanginakulu stream is fed by low discharge warm springs of bicarbonate-sulphate type,
- and is relatively high in Mg (Table 3). Temperature, DIC, Ca, and  $SO_4^{2-}$  all decrease
- 194 downstream, whereas pH increases (Fig. 10). Unlike the Rembokola stream, there are no
- 195 consistent changes in the concentrations of conservative elements (e.g. Cl<sup>-</sup>).
- 196 Sinters, travertines and mixed deposits
- 197 The Rembokola sinters analysed in this study were dominated by opal-A, with more
- 198 anhydrite-rich mineralogy in spicular facies. Layered sinter samples have elevated Al, Na and
- 199 Ti compared to travertines (Table 4), reflecting a greater component of detrital clastic
- 200 material (e.g. feldspar and magnetite, abundant in the volcanic host rocks) entrained within
- 201 the sinter layers. The near-neutral to alkaline pH waters do not transport significant
- 202 concentrations of Al or Ti, and Na is conserved in the aqueous phase and not readily
- 203 precipitated. Sulphur is below detection limits in layered sinters, but relatively high
- 204 (2.24 wt%) in the spicular sinters, confirming the presence of anhydrite in this facies. Sinter
- samples analysed by ICP-MS following aqua regia digestion (Table 5) show Te present in
- 206 concentrations of 40  $\mu$ g/kg, at least 800 times higher than in spring water (<0.05  $\mu$ g/l), and ~8
- 207 times greater than average crustal abundances (Wedepohl 1995).
- 208 In the interlayered mixed, silica–carbonate deposit from Rembokola (SV482), Sr is high due
- 209 to calcite contents (Table 4). Carbonate layers show significant arsenic enrichments
- 210 compared to sinters and silica layers. Tellurium (Table 5) is higher in the bulk-analysed
- 211 mixed sample than the sinters from the same stream, at concentrations >50 times higher than
- average crustal abundance (Wedepohl 1995).

- 213 All bulk Rembokola samples analysed in this study for elements related to epithermal
- 214 mineralisation have low but significant concentrations of Au, Ag, Hg and Cu, as well as the
- 215 Te previously discussed (Table 5).
- 216 The hot spring deposits of the Poghorovorughala area also have mixed mineralogy. Similar to
- 217 the Rembokola sinters, Te contents of the deposits are high (250–380  $\mu$ g/kg; Table 5) and
- spring waters are below detection limits ( $<0.05 \mu g/l$ ). Lobate mixed silica–carbonate samples
- show elevated Al and Ti in a similar fashion to the layered sinters of Rembokola (Table 4),
- most likely representing detrital or substrate material entrained within the sample (e.g. Fig.6B).
- 222 The travertine deposits of Tanginakulu are dominated by calcite. Similar to the mixed
- samples from Rembokola and Poghorovorughala, Te occurs in very high concentrations
- 224 (410 μg/kg; Table 5). Tanginakulu travertines contain notably high Fe concentrations; springs
- in the area deposit distinctly red sediments (sludge) upon discharge, probably indicative of
- reduced waters being oxidised upon exiting at the surface.

# 227 Discussion

- 228 Travertines and travertine-forming waters
- 229 Tanginakulu stream is dominated by relatively cool Ca–Mg–HCO<sub>3</sub><sup>-</sup> type water compositions.
- 230 The magnesium contents indicate a low temperature (<100°C) origin for these waters, as at
- higher temperatures Mg is readily removed by formation of minerals such as chlorite
- 232 (Giggenbach 1988). Stream waters are moderately supersaturated with respect to calcite (Fig.
- 233 10), with  $\log Q/K = 0.5$  to 1, where Q is the ion activity product and K is the equilibrium
- constant of the calcite-forming reaction, as calculated with SOLVEQ (Reed 1982; 1998). The
- 235 Tanginakulu spring analysed in this study has values of  $\log Q/K = -0.3$ .
- Travertine precipitation from stream and spring waters initially enriched with calcium and
  bicarbonate is typically driven by CO<sub>2</sub> removal (Pentecost 2003):
- 238  $Ca^{2+} + 2HCO_3^{-} = H_2O + CaCO_3 + CO_2$  1
- Removal of carbon dioxide can be biotic (photosynthesis), or abiotic (degassing). The latter
  mechanism is the predominant process in most streams and springs, and is particularly
  effective where water is turbulent (Pentecost 2003). There is a strong association with
  travertine deposition and thicker deposits in areas of waterfalls and rapids on Savo. Carbon

243 dioxide loss to the atmosphere is therefore the most likely cause of calcite supersaturation and244 precipitation.

Examination of travertines (Fig. 8) shows that they are composed of layers of calcite ray-

crystals. Calcite is the dominant CaCO<sub>3</sub> polymorph at temperatures <40°C (Jones *et al.* 

247 1996), and so its predominance over aragonite in these deposits is unsurprising. Ray-crystal

248 layers are common in travertine, and are typically abiotic in origin, formed by rapid

precipitation of calcite from supersaturated solutions (Folk *et al.* 1985; Chafetz & Guidry1999).

As well as causing carbonate precipitation,  $CO_2$  loss is an important mechanism for increasing water pH (Chafetz *et al.* 1991; Fouke *et al.* 2000):

253 
$$HCO_3^- + H^+ = H_2O + CO_2$$
 2

Figure 10 shows the changes in DIC and pH in the downstream direction of Tanginakulu stream. In particular, there is a rapid drop in DIC and corresponding increase in pH after discharge from the warm spring into the stream proper. Combined  $CO_2$  loss and travertine precipitation is capable of producing the relationships displayed in figure 10.

258 Tellurium is notably enriched in carbonate bearing samples (Table 5), despite being below 259 detection limit ( $<0.05 \mu g/l$ ) in all water samples in this study. Arsenic also shows high 260 concentrations in carbonates, particularly those within the mixed sample from Rembokola. 261 Arsenic is associated with the higher temperature component: its concentration is higher in the Rembokola springs and stream than in the Ca-Mg-HCO<sub>3</sub><sup>-</sup> enriched springs (Tables 2 and 262 263 3). Tellurium would also be expected to be associated with high temperature fluids, given that 264 it can be transported in a magmatic vapour phase (Cooke et al. 1996). High Te concentrations in the carbonates may reflect tellurate substitution for  $CO_3^{2-}$ , in a similar mechanism to 265 sulphate substitution (Takano et al. 1980). Elevated arsenic in carbonates likely reflects 266 267 adsorption or co-precipitation of arsenate with calcite (Alexandratos et al. 2007). Thus, high 268 concentrations of arsenic and tellurium occur in carbonate deposits – despite these deposits 269 being associated with lower temperature fluids -as carbonate minerals are suitable hosts for 270 As and Te. It appears that in the case of Rembokola, opal-A is less capable of hosting As and 271 Te, despite being associated with the high temperature fluid end-member.

#### 272 Sinters

273 Sinter is deposited within the Rembokola valley. The stream is a relatively closed system:

there are few major springs feeding the stream other than those of the upstream thermal area,

and there are no major tributaries into the stream. In the downstream direction, evaporation

276 causes a decrease in temperature, and increases in conservative elements such as B, Li, and

277 Cl<sup>-</sup>. Simple calculations indicate that approximately 10% of the original mass of water is lost

278 over <1 km to produce the observed Cl<sup>-</sup> and Li increases. Mn and Si decrease by mineral

279 precipitation, whereas  $CO_2$  loss leads to pH increase (Eqn. 2) and DIC decrease (Fig. 9 and 280 Table 2).

281 The combined effects of evaporation, cooling and CO<sub>2</sub> loss is that amorphous silica (opal-A)

282 becomes increasingly supersaturated (increasing log Q/K) downstream (Fig. 9). Calcite is

also supersaturated (decreasing downstream, due to decreasing temperature and retrograde

calcite solubility; Rimstidt 1997) but the DIC contents of the waters are low (<50 mg/l

HCO<sub>3</sub><sup>-</sup> eqv.), and it is likely that any precipitated carbonate is masked by greater volumes of silica.

287 Silica precipitates near hot springs in two distinct facies – as terraced deposits on the steep 288 slopes in the upper Rembokola valley, and as spicules on subaerially exposed substrate in the 289 relatively flat-lying thermal area. The spiculose sinter described in this study is 290 morphologically similar to the silica-carbonate "meringue" deposits of the Pavlova Spring, Ngatamariki, New Zealand (Campbell et al. 2002). These authors concluded that the Pavlova 291 292 deposits formed by evaporation of hot spring-derived water from subaerially exposed 293 surfaces, typically upon partially submerged detritus (principally leaf litter). Meniscoid and 294 capillary creep ("wicking"; Hinman & Lindstrom 1996; Campbell et al. 2002), as well as

sporadic bathing and splashing in the case of the Savo deposits, replenish fluids. Individual

spicules reach a maximum height (~2 cm) above which the wicking process can no longer

replenish moisture in sufficient quantity to allow further growth (Campbell *et al.* 2002).

298 Further evidence of evaporation as a precipitation mechanism is the presence of anhydrite on

the upper surfaces of the spicules (Fig. 3). At spring temperatures and below, anhydrite is

300 undersaturated (Fig. 9). The only effective mechanism for precipitating anhydrite is

301 evaporation.

302 The upstream terraced sinter may also be precipitated through evaporation, but in the case of 303 the deposits on the steep slopes, wicking is less important than direct evaporation from the surface. Water and dissolved silica is supplied constantly by the spring fluids bathing the
discharge apron, whilst never submerging it entirely. Terrace-type constructions are common
in both travertine and siliceous sinter deposits, and occur where precipitation is from sheet
flow (Guidry & Chafetz 2003). The stair-step morphology of the micro-terracettes is

308 produced by random perturbations in deposition, perhaps produced by debris or microbial

309 mats (Chafetz & Folk 1984; Guidry & Chafetz 2003) that eventually reorganise into linear or

310 curvilinear ridges (Hammer *et al.* 2007).

311 Evaporation and cooling of the hot spring fluids as they flow downstream leads to an increase

312 in the saturation index of amorphous silica (Fig. 9) and sinter precipitation (Rimstidt & Cole

313 1983). Around hot springs, sinter only forms upon exposed or periodically bathed surfaces,

but in the stream channel downstream of the hot springs, sinter is deposited upon wholly

315 submerged surfaces. The sinters also contain filaments or tubules preserved by opal-A,

316 typically aligned and orthogonal to the growth laminations of the sinter (Fig. 4). The

317 orientation may be a result of filaments aligning with flow direction in the stream (Jones *et* 

318 *al.* 2003). Such filamentous structures are commonplace in siliceous sinters, and are the result

319 of the enclosure and partial preservation of microbes (Jones & Renaut 2003a, b; Jones *et al.* 

2003; Lynne & Campbell 2003; Konhauser et al. 2004; Fernandez-Turiel et al. 2005; Jones et

*al.* 2005). Filaments were not observed in the spicular facies, and only rarely in the terraced

322 sinter (Fig. 2); as the samples were not preserved with a fixative solution, the presence and

323 abundance of microbes in these facies is unknown.

324 Thermal waters may be colonised by a range of microorganisms, including cyanobacteria,

325 bacteria and fungi; however, low preservation fidelity of the organisms following

326 silicification (replacement and/or encasement with silica, during or shortly after life) often

327 makes taxonomic identification difficult (Jones *et al.* 2003). The fossils preserved in the

328 Rembokola stream sinters are simple, non-branching filaments, approximately  $100 \,\mu m$  in

length and 5  $\mu$ m in diameter, although silica cementation means that the diameter of the

preserved filament may be significantly larger than that of the living organism, (Jones *et al.* 

331 2003). *Phormidium* cyanobacteria are common in thermal areas, and have an appropriate

morphology (Pentecost 2003) but the lack of more complex features preserved in the

333 Rembokola sinters preclude definitive classification.

334 Some noteworthy aspects of the analysed sinter chemistry is the low but significant As

concentrations, and 2–20 mg/kg Cu (Table 4, 5). Despite the Cu and Fe contents of the

336 samples, no sulphide minerals (pyrite, chalcopyrite) were observed under SEM; in fact, with

- the exception of anhydrite in the spicular and terraced sinter and a few clasts of detrital
- 338 material (trachytic volcaniclastics with elevated Al, Na and Ti; Table 4), no minerals other
- than opal-A were observed. Accessory elements can be bound into the structure of opal-A
- 340 without requiring distinct mineral phases (Jones & Renaut 2003a). ICP-MS analysis of a
- 341 subset of the sinter samples (Table 5) show that Te is lower than in carbonate samples,
- 342 possibly reflecting more effective Te scavenging from fluids by carbonate than amorphous
- 343 silica, as discussed above for the travertine samples.

## 344 Lobate and spicular mixed deposits

345 Mixed silica-carbonate + anhydrite spicules grow on the periphery of Poghorovorughala hot 346 springs, upon infrequently splashed and bathed surfaces (Fig. 6A). The increased proportion 347 of opal-A and anhydrite in these samples indicates that they precipitate from more highly 348 evaporated spring waters, as anhydrite and amorphous silica are marginally saturated and 349 undersaturated, respectively, in the spring waters. The spicules here are morphologically 350 similar to those of the Rembokola area, albeit with more carbonate. Mineralogy is closer to 351 the Pavlova Spring deposits (Campbell et al. 2002), with both carbonate and silica phases, 352 and the spicules at Poghorovorughala are interpreted to form in the same way – by wicking of 353 hydrothermal fluids from infrequently bathed and splashed surfaces, resulting in evaporative 354 precipitation of sinter/travertine.

355 The lobate silica–carbonate deposits surrounding Poghorovorughala hot springs (Fig. 6)

356 contain carbonates, with anhydrite and opal-A. In these deposits, CO<sub>2</sub> loss causes carbonate

357 precipitation, and evaporation precipitates anhydrite and silica, similar to the spiculose facies.

- 358 For the most part, deposits are microcrystalline to amorphous, with the exception of well-
- developed trigonal prisms of calcite in sheltered areas between lobes (Fig. 6H). At
- 360 precipitation temperatures above 40°C, aragonite is the expected polymorph of CaCO<sub>3</sub>, with
- 361 some exceptions. For example, Jones *et al.* (1996) described calcite deposited from Waikite

362 Hot Springs, New Zealand, where water temperatures are >90°C. The near-spring deposits at

- 363 Poghorovorughala contain both calcite and aragonite, and water temperatures are >90°C;
- however, as the deposits are formed in splashed and bathed areas, rather than submerged, it is
- 365 possible that there is precipitation both above *and* below the 40°C boundary temperature.
- 366 Without real-time observations of precipitation it is difficult to determine whether calcite is
- 367 precipitating at an unusually high temperature.

#### 368 Interlayered mixed deposits

369 Interlayered mixed silica-carbonate deposits are found above present stream levels in the 370 mid- to upper reaches of the Rembokola. The mixed deposits are clearly older than the silica 371 sinters, as they are above the present day stream level, and have indurated and weathered 372 upper surfaces (Fig. 5A). The silica layers include filaments in void spaces, similar to those 373 observed in the stream sinters (Fig. 4E; Fig. 5E). The silica layers tend to be more massive 374 than in the silica-only sinters, with fewer preserved filaments and lower porosity, perhaps as a 375 function of age. Over time, diagenetic transformation and continued silica precipitation leads 376 to the destruction of primary depositional fabrics (Jones & Renaut 2003a). As discussed

377 above, silica layers tend to have a higher clastic content than carbonate layers (Fig. 5F).

378 Carbonate layers in the Rembokola mixed deposit consist of ray-crystal calcite (Fig. 5C),

379 similar to the travertines at Tanginakulu. Similar precipitation mechanisms are envisaged –

380 CO<sub>2</sub> degassing in an area of turbulent flow leads to calcite supersaturation and precipitation.

381 SV482 in particular shows enrichments of Te and As, both considered pathfinder elements

382 for epithermal Au deposits (White & Hedenquist 1995), with Te in particular associated with

383 alkaline-related epithermal deposits (Jensen & Barton 2000). The increased concentration of

384 Te and As in SV482 relative to Rembokola sinters is potentially a result of the combination

385 of more effective scavenging of these elements by carbonates. SV482 also has higher As and

386 Te than travertine from Tanginakulu, potentially reflecting increased concentration of these

387 elements in the waters forming mixed deposits versus those that only form travertine.

388 Interlayered mixed silica–carbonate deposits are also found surrounding the Mound Spring at

389Poghorovorughala (Fig. 7). Layers of opal-A contain filaments similar to the sinters

390 elsewhere on the island (Fig. 7D; Fig. 4D). The lack of alignment in the filaments is most

391 likely a result of the low flow rate on the Mound Spring's discharge apron relative to the

392 Rembokola stream.

Although elsewhere calcite and silica can be found in the same deposits, (Jones *et al.* 1996;
Campbell *et al.* 2002), the situation is rare, as the two minerals are associated with different

fluid types (in terms of origin and chemistry) in most geothermal areas (Canet *et al.* 2005).

396 The interlayered silica–carbonate deposits show that the Rembokola stream and

397 Poghorovorughala Mound Spring have historically alternated between travertine and sinter

398 formation. Carbonate precipitation is from waters with a higher contribution from low-

399 temperature fluid (e.g. Tanginakulu bicarbonate-sulphate spring), and silica precipitation is

400 from waters dominated by the higher temperature fluids. The periodic switching between the

401 two situations reflects changes in the degree of mixing between the two end-member fluids at

402 source. If DIC contents are too low, then calcite precipitation is masked by silica

403 precipitation, or simply prohibited by the lack of sufficient supersaturation.

- 404 Comparison between the Rembokola and Poghorovorughala springs shows that fluid mixing
  405 already occurs (Smith *et al.* 2010); for example, Mg contents are far higher than would be
- 406 expected for waters at the temperatures recorded (Giggenbach 1988). The Poghorovorughala

springs have a higher contribution from the Ca-Mg-HCO<sub>3</sub><sup>-</sup> end-member compared to the

- 408 Rembokola springs. Poghorovorughala spring waters are supersaturated with a number of
- 409 mineral phases at discharge temperature, most notably with calcite (log Q/K  $\approx$  1.2), and
- 410 aragonite (log Q/K  $\approx$  1.1), and saturated with anhydrite (log Q/K  $\approx$  0.1). The waters are
- 411 under-saturated with respect to amorphous silica (log Q/K  $\approx -0.2$ ) although should saturate
- 412 upon cooling to approximately 60°C (calculated with SOLVEQ; Reed 1982; Reed 1998). At
- 413 present, the Rembokola waters precipitate only opal-A (and minor anhydrite) whereas the
- 414 Poghorovorughala springs precipitate carbonates, opal-A and anhydrite.
- 415 What causes the periodic changeover between carbonate and silica precipitation in 416 interlayered mixed samples is unknown. Three principal mechanisms can be suggested: 1) 417 episodic changes in relative contribution of magmatic fluids to the hydrothermal system 418 (Boichu et al. 2008); 2) changes in the hydrology and plumbing of the system causing 419 varying contributions of both components (e.g. Leeman et al. 2005); 3) seasonal changes in 420 rainfall causing variation in the low temperature components (e.g. López et al. 2006). All 421 three models may operate to produce periodic changes in the hydrothermal system at Savo; 422 indeed, López et al. (2006) point out that there may be relationships between the different 423 mechanisms, such as atmospheric pressure affecting rainfall, degassing rate and seismic 424 tremor (López et al. 2006; Neuberg 2000). Irrespective of the mechanism driving periodic 425 switchover between carbonate and silica in interlayered mixed samples, the importance of 426 high meteoric water input to the hydrothermal system of Savo is underlined, as it is the 427 mixing of the meteoric-derived low temperature end-member that drives hot springs and 428 streams from silica to carbonate precipitation.

# 429 Conclusions

407

Hydrothermal discharges at Savo produce a range of deposits, including travertine, sinter and
unusual mixed silica–carbonate rocks. Previous work has shown that there are multiple fluid

432 types within the hydrothermal system, including a silica-rich end-member associated with

433 high temperature water-rock-gas interaction, and a Ca-Mg-HCO<sub>3</sub><sup>-</sup> end-member derived by

434 low temperature water-rock-gas interaction. The streams and springs discussed in this study

435 can be classified according to which component dominates: Rembokola is dominated by the

436 high-temperature end-member, Tanginakulu by the low-temperature end-member, and

437 Poghorovorughala springs are mixed.

The different water chemistries of the springs and their outflow streams give rise to different

439 surface deposits. Waters dominated by the Ca–Mg–HCO<sub>3</sub><sup>-</sup> end-member precipitate

440 travertine, those dominated by the high-temperature end-member form silica sinter, and the

441 Poghorovorughala springs form intimately mixed silica–carbonate lobes and spicules.

442 Unusual interlayered mixed silica-carbonate deposits occur in the Rembokola stream system 443 and surrounding the Mound Spring at Poghorovorughala. The carbonate layers are similar to 444 the travertine found in the Tanginakulu area, and the silica layers similar to the sinters 445 currently forming in the Rembokola stream. The samples may reflect meteorological 446 changes, possibly seasonal, with high rainfall leading to increased contributions from a low 447 temperature, carbonate-forming, fluid end-member. Periodic changes in magma degassing, 448 and reorganisations of the hydrothermal plumbing system by seismic activity may also 449 account for the periodic variations in spring discharges and their products.

The Savo deposits show that sinter and travertine can be deposited from the same spring with relatively little change in water chemistry required, only a change in the relative contributions

452 of two end-member fluids. As a result, travertines – normally associated with fluids

453 peripheral to a hydrothermal system – can carry chemical signatures (e.g. enriched Te) more

454 usually associated with sinter-forming, higher temperature, 'hypogene' fluids fed by

455 magmatic volatiles. In particular, the carbonate layers of the mixed silica–carbonate deposits

456 record high As and Te concentrations. This reflects the increased availability of these

457 magmatic-hydrothermal components in the high temperature end-member fluid, combined

458 with adsorption of arsenate and tellurate species by carbonate minerals.

459 In magmatic-hydrothermal systems with considerable meteoric water input, cooler-water

460 deposits, i.e. travertine, are favoured. As such, travertine, and in particular, mixed silica-

461 carbonate deposits can carry trace elements indicative of the underlying magmatic-

462 hydrothermal system, pointing to potential economic mineralisation at depth.

# 463 Acknowledgements

- 464 DJS was funded by the Natural Environment Research Council (UK) and British Geological
- 465 Survey University Funding Initiative as part of PhD studentship NER/S/A/2004/12339. SF
- 466 received funding from the Society of Economic Geologists Student Research Grants (Hugh
- 467 E. McKinstry Award). The authors would like to thank H. Taylor, K. Green and RA. Shaw
- 468 (BGS); W. Satokana, G. Albert, A. Ramo, D. Billy and S. Basi (Solomon Islands Geology
- 469 Division). JN and SRNC publish with the permission of the Executive Director, British
- 470 Geological Survey (NERC). This paper is dedicated to the late Watson Satokana without
- 471 him this project would not have been possible.

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- 591

Figures



Fig. 1: A) Map of the south west Pacific. Solomon Islands shown in box. B) Map of the Solomon Islands, showing location of Savo. C) Map of Savo showing location of major streams, thermal areas, and water samples discussed in this study. Pogho. = Poghorovorughala.



Fig. 2: Rembokola terraced sinter. A) Terraced sinter (east of point a, Fig. 1). Deposit is composed of micro-terracettes of opal-A. B) Photograph of the interior of the sinter, showing highly porous opal-A. C) BSE image of broken surface showing silica-encrusted filament in cavity.



Fig. 3: Rembokola spicular sinter (points a and b, Fig. 1). A) Opal-A sinter growing on leaf litter in Rembokola stream, at Toakomata hot spring area. B) Back scattered electron (BSE) image of Rembokola spicular sinter. Upper surface is to the right in the field of view, and contains more anhydrite (white) than the lower areas, which is dominated by opal-A (grey).



Fig. 4: Silica sinter in the Rembokola valley (point e, Fig. 1). A) Rembokola stream channel is lined with sinter (covered with algae in this view). The surrounding gorge floor is clastic material cemented by silica sinter. B) Cross section view through a silica sinter crust on sediment substrate. C) BSE image of broken surface showing laminations in sinter; lower layer is massive and low porosity opal-A, upper layer is more porous contains elongate filaments. D) BSE image of elongate hollow filaments in sinter. E) BSE image of filaments within a larger void space.



Fig. 5: Interlayered mixed silica–carbonate sinter, Rembokola valley (point d, Fig. 1). A) Terrace of mixed sinter approximately 30 cm above current stream water level. B) Cross section through sinter showing layers of calcite (dark) and opal-A (pale). C) Thin section through carbonate layers showing fans of ray-crystals (cross polarised light). D) BSE image showing calcite layer onlapping onto older silica and carbonate layers. E) BSE image of silica layer, with filaments visible in void space. F) BSE image from thin section showing silica layer with detrital material (white = magnetite, grey = feldspar, derived from local volcaniclastics). Carbonate layers (top and bottom of image) contain fewer clasts of foreign material.



Fig. 6: Lobate deposits, Poghorovorughala. A) Lobate deposits surround the spring (visible as dark cavity immediately right of lobate label) and discharge channel; spicules occur on the periphery. B) Cross section through lobes, showing concentric laminations. Centre of lobe is kaolinite-dominated, representing the substrate of the steam-heated ground on which the deposits form. C) Upper surface of lobe. D) BSE image of underside of lobe (submerged portion). E) BSE image of rounded lobes of carbonate developing on subaerially exposed / splashed portion. F) BSE image of pyrite on surface of carbonate, opal-A and minor anhydrite in submerged portion. G) BSE image of carbonate and opal-A, with occasional anhydrite crystals, on splashed area of deposit. H) Detail view of calcite, showing trigonal crystal form.



Fig. 7: Interlayered mixed silica-carbonate deposits, Poghorovorughala. A). View of Mound Spring, a 3 m high interlayered deposit of . A hot spring discharges from the summit of the mound. B) Microterracette texture on surface of the Mound Spring. C) Cross section through mixed layers of Mound Spring deposit. D) Dark layer is a mixture of tubes/ filaments of opal-A, and crystals of calcite, shown in BSE image. E) BSE image of carbonate fans from pale layers of mixed deposit. F) Detail view of calcite ray-crystals, showing minor anhydrite.



Fig. 8: Tanginakulu travertine deposits. A) Travertine deposited at small stream rapids. B & C) Cross section photograph and BSE image through travertine showing laminations of carbonate and fans of elongate calcite ray-crystals.



Fig. 9: Changes in temperature, pH, dissolved Cl and Si concentrations, and saturation index of important minerals in the Rembokola stream. Representative alkaline sulphate springs (or maximum and minimum values in the case of a range) shown for comparison. Error bars are  $\pm 1\sigma$ ; not shown when within point size.



Fig. 10: Changes in temperature, pH, dissolved Cl, DIC (as  $HCO_3^-$  eqv.), and saturation index of important minerals in the Tanginakulu stream. SV422 is a warm spring in the upper reaches of the stream (Table 1). Error bars are  $\pm 1\sigma$ ; not shown when within point size.

# Tables

Sample Type	Pogho. hot spring Alkaline sulphate	Rembokola hot spring Alkaline sulphate	Tanginakulu spring SV422 Bicarbonate sulphate	
T (°C)	100	100	47	
рН	7.5	7.5	6.7	
Al (µg/l)	11	7	2	
As (µg/l)	<2	50	19	
В	2.15	8.78	0.20	
Ba (µg/l)	40.9	55.8	44.3	
Ca	239	96	204	
Fe	0.04	< 0.10	3.03	
Κ	16.8	28.5	5.9	
Li (µg/l)	290	1684	55	
Mg	12.0	4.4	98.5	
Mn	0.71	0.14	0.561	
Na	81	216	48.3	
$SiO_2$	256	374	157	
Sr	3.25	2.74	1.49	
DIC	86	38	513	
$\mathrm{SO_4}^{2-}$	669	627	294	
Cl	4.4	46.7	7.6	

#### Table 1: Water chemistry from selected springs on Savo.

All units in mg/l unless noted otherwise. DIC as  $HCO_3^-$  equivalent. Original data reported in Smith et al. (2010).

Table 2: Water chemistr	v data for	Rembokola	stream	samples.
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Sample Distance	а	b	с	d	e	f	g	h	i	j	k
(km)	0.00	0.01	0.11	0.30	0.61	0.86	1.13	1.48	1.73	1.98	2.21
T (°C)	72	75	52	45	40	38	35	35	34	33	31
pН	7.9	8.1	8.5	8.7	8.8	8.6	8.5	8.6	8.6	8.5	8.5
DIC	72	62	49	44	42	47	53	47	44	47	49
Al (µg/l)	7	6	3	bdl	2	3	bdl	2	2	4	9
As (µg/l)	68	65	66	69	70	70	73	69	72	73	69
Ca	169	152	152	153	152	153	155	154	154	153	154
Fe	0.04	0.03	0.03	0.03	bdl	bdl	bdl	bdl	bdl	0.03	0.03
Κ	25.6	26.2	28.1	28.5	28.3	28.5	29	29	28.7	28.5	28.3
Mg	8.8	7.7	8	8	8	8.1	8.3	8.2	8.2	8.2	8.2
Mn	0.39	0.33	0.31	0.22	0.15	0.15	0.14	0.08	0.07	0.07	0.08
Na	174.8	184.1	198.1	200.6	200.4	203.3	202.7	203.9	203.6	202.4	200.9
Si	160	164	172	165	154	150	148	142	141	136	135
$SO_4^{2-}$	684	668	696	710	711	719	715	716	717	713	713
Cl	38	38.9	43	42.9	42.8	44	43.4	44	43.8	43	43.9
CBE (%)	5	5	4	3	4	4	3	4	4	4	5

All values in mg/l unless noted otherwise. bdl = below detection limits; DIC = dissolved inorganic carbon as mg/l  $HCO_3^-$  eqv.; CBE = charge balance error. Full trace element analyses available in Supplementary Data.

Sample	1	m	n	0	
Distance (km)	0.00	0.24	0.56	1.00	
T (°C)	35.4	32	29.9	28	
pН	8.1	8.1	8.4	8.5	
DIC	295	232	199	163	
Al (µg/l)	13	bdl	3	bdl	
As (µg/l)	7	8	10	11	
Ca	166	113	88	82	
Fe	0.06	0.02	bdl	bdl	
K	5.8	5.7	6.0	5.2	
Mg	76.9	75.2	79.5	54.1	
Mn	0.18	0.01	bdl	bdl	
Na	40.9	40.0	43.0	33.7	
Si	60	60	62	53	
$SO_4^{2-}$	286	268	266	190	
Cl	7.5	6.8	7.6	8.0	
CBE (%)	24	17	19	19	

Table 3: Water chemistry data for Tanginakulu stream samples.

CBE (%)24171919All values in mg/l unless noted otherwise. bdl = below detection limits;DIC = dissolved inorganic carbon as mg/l HCO<sub>3</sub><sup>-</sup> eqv.; CBE = chargebalance error. High CBE may be a result of carbonate speciation (i.e. $CO_3^{-2}$ > HCO<sub>3</sub>), or unanalysed HS<sup>-</sup>. Full trace element analysesavailable in Supplementary Data.

Stream Maj	Мар	Sample	Preparation	Sample	Al	As	Ca	Cu	Fe	K	Li	Mg	Mn	Na	Rb	S	Sr	Ti	V
Stream	Ref.	Туре	reparation	Number	mg/kg	mg/kg	wt %	mg/kg	wt%	mg/kg	mg/kg	mg/kg							
			Detecti	on Limits:	196	0.2	0.08	0.5	23	28	0.4	74	8	286	0.08	0.31	14	4	0.1
Rem.	a	Terraced sinter	Bulk	497	-	1.7	0.2	-	44	195	6.6	174	67	619	1.89	-	47	-	0.5
Rem.	b	Spiculose sinter	Bulk	486	3511	40.9	6.2	2.3	1413	1363	18.7	11526	1118	3455	7.69	2.24	1373	70	2.9
Rem.	c	Layered sinter	Opal-A Layer	484 a	9051	5.1	1.8	7.4	4570	1285	3.9	2478	3338	4119	3.30	-	278	171	11.3
Rem.	c	Layered sinter	Opal-A Layer	484 b	6982	4.7	1.6	7.5	3330	1083	3.5	1779	2573	3310	2.79	-	264	133	8.3
Rem.	d	Layered MSC	Carbonate Layer	482 a	326	357	42.1	5.8	2162	278	7.0	2005	9488	748	0.61	1.71	4598	12	0.6
Rem.	d	Layered MSC	Opal-A Layer	482 b	1219	17.8	2.0	10.0	2579	360	2.3	3137	2148	526	1.81	-	246	64	3.7
Rem.	d	Layered MSC	Opal-A Layer	482 c	2457	18.1	3.9	15.6	3504	626	3.2	4076	3471	807	1.86	-	588	94	7.0
Rem.	d	Layered MSC	Carbonate Layer	482 d	523	254	34.5	7.2	1225	245	10.5	2009	6172	617	0.51	1.06	3210	19	0.7
Rem.	d	Layered MSC	Carbonate Layer	482 e	2719	378	37.4	20.1	4442	681	31.6	2779	6273	1786	1.78	1.00	3227	152	8.4
Rem.	g	Layered sinter	Opal-A Layer	475 a	1413	1.8	1.0	3.9	928	336	2.8	1928	2552	671	1.37	-	156	47	2.8
Rem.	g	Layered sinter	Opal-A Layer	475 b	7226	1.9	1.1	9.1	4865	1262	3.8	2303	3118	3299	3.80	-	220	181	14.7
Tan.	1	Travertine	Carbonate Layer	425 a	-	15.3	42.6	1.2	9293	42	0.6	722	384	506	0.14	-	3280	8	0.5
Tan.	1	Travertine	Carbonate Layer	425 b	-	80.4	35.5	2.6	40402	123	0.5	1426	582	393	0.55	-	3060	-	0.3
Tan.	1	Travertine	Carbonate Layer	425 c	-	7.7	19.8	-	2383	59	-	631	346	342	-	-	2849	-	-
Tan.	m	Travertine	Carbonate Layer	434 a	-	8.9	17.7	2.4	1554	65	1.6	6070	451	-	0.20	-	1307	8	0.5
Tan.	m	Travertine	Carbonate Layer	434 b	-	10.4	19.5	3.2	2498	93	1.0	6063	754	-	0.17	-	676	9	0.8
Tan.	m	Travertine	Carbonate Layer	434 c	-	12.1	25.0	2.4	3436	141	1.5	8253	550	395	0.26	-	1223	78	1.0
Pog.		Lobate MSC	Bulk	501	3184	0.4	23.9	0.6	1079	1294	1.7	123	938	893	1.56	0.62	3677	247	2.4
Pog.		Lobate MSC	Bulk	502	32890	0.7	21.7	7.7	2024	6002	10.4	158	312	15584	9.23	0.63	3975	492	14.3
Pog.		Layered MSC	Carbonate Layer	505 a	-	-	26.4	-	332	254	3.5	82	758	543	0.36	0.33	5764	43	0.3
Pog.		Layered MSC	Carbonate Layer	505 b	-	_	41.2	130	603	93	1.1	1429	14021	380	0.23	1.29	2587	11	_
Pog.		Layered MSC	Carbonate Layer	505 c	-	-	30.8	0.6	171	-	0.8	81	893	314	-	-	5080	-	-
Pog.		Spiculose sinter	Bulk	506	431	-	17.4	2.2	1067	307	3.0	8392	1853	688	1.18	3.48	2326	6	0.4

Rem. = Rembokola; Tan. = Tanginakulu; Pog. = Poghorovorughala. MSC = Mixed silica carbonate. Dash in cell indicates analyte below detection limits.

#### Table 4: Whole rock sinter and travertine chemistry.

Stream Map Sample Sample Ag As Au Bi Cd Cu Hg Mo Pb Sb Se Te Zn

		Detection	n Limits:	2	0.1	0.2	0.02	0.01	0.01	5	0.01	0.01	0.02	0.1	20	0.1
Rem.	d	Layered MSC S	V482	3	303	1.9	-	0.11	4.19	14	0.11	0.13	0.03	0.2	300	3.6
Rem.	e	Layered sinter S	V479	9	3.1	1.3	0.02	0.02	18.23	25	0.84	0.72	0.03	0.2	40	18.2
Rem.	g	Layered sinter S	V475	6	2.1	2.9	-	0.04	10.18	17	0.28	0.34	0.02	0.4	40	6.9
Tan.	1	Travertine S	V425	-	54	-	-	0.05	0.37	_	0.04	0.03	0.09	0.2	410	21.0
Pog.		Layered MSC S	V505	3	0.7	_	-	-	0.67	9	0.09	1.25	-	-	250	5.6
Pog.		Layered MSC S	V514	-	0.6	_	_	-	0.15	_	0.08	0.08	_	0.1	380	0.3

Ref. Type Number µg/kg mg/kg µg/kg mg/kg mg/kg

Rem. = Rembokola; Tan. = Tanginakulu; Pog. = Poghorovorughala. MSC = Mixed silica carbonate. Dash in cell indicates analyte below detection limits.

Table 5: Whole rock sinter and travertine chemistry – elements of significance to mineral exploration