1	Dolomitization of the Lower Ordovician Watts Bight Formation of the St Georges
2	Group, Western Newfoundland: evidence of hydrothermal fluid alteration
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19	Reference
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29	Abstract
30	The Watts Bight Formation in western Newfoundland consists of a succession of shallow
31	water carbonates and has been extensively dolomitized. These dolomites occur as both
32	replacements and cements and are associated with complex changes in the rock porosity
33	and permeability. Early replacement micritic dolomites (D1) are finely crystalline and
34	indicate that dolomitisation began during early stages of diagenesis. The calculated $\delta^{18}O$
35	of the earliest (D1) dolomitizing fluids (-6.4 to -9.5‰ VSMOW) fall between the
36	estimated $\delta^{18}$ O values of the Tremadoc seawater and meteoric waters and suggest mixing-
37	zone dolomitization. A second phase of coarsely crystalline (up to $400\mu m$ ) dolomite (D2)
38	replaces D1 dolomite and early calcite and is associated with enhancement in porosity
39	and permeability through the development of intercrystalline pores. A late-stage saddle
40	dolomite (D3) and late burial calcite cements significantly occluded the pores in some
41	horizons. Petrography, fluid inclusions and geochemistry show that D2 and D3 dolomites
42	formed from warm (65 to $125^{\circ}$ C) saline (10 to 25 eq. wt% NaCl + CaCl <sub>2</sub> ) hydrothermal
43	fluids. The calculated $\delta^{18}O_{fluid}$ of D2 ranges from -4.5 to 3.6 $\%$ VSMOW and for D3
44	dolomites calculated $\delta^{18}O_{fluid}$ ranges from 1.4 to 8.4 $\%$ VSMOW, suggesting an influx of
45	basinal brines. The occurrence of high porosity associated with D2, combined with the
46	laterally sealing tight limestone beds, presence of favourable source rocks and thermal

- 47 maturation, may suggest that the Watts Bight carbonates are possible potential
- 48 hydrocarbon reservoirs and suitable targets for future hydrocarbon exploration in Western
- 49 Newfoundland.
- 50

## 51 Introduction

52 Hydrothermal dolomite (HTD) hydrocarbon reservoirs have been recognised as an 53 important exploration target and are major hydrocarbon producers in North America (e.g., 54 Davies and Smith 2006). This is because HTD reservoirs typically retain porosity and 55 permeability much better than precursor limestones. Recently, dolostones including 56 hydrothermal dolomites in the lower Paleozoic rocks of eastern Canada and the United 57 States have been the focus of a number of studies (e.g., Cooper et al. 2001; Lavoie et al. 58 2005; Lavoie and Chi 2006; Smith 2006; Azmy et al. 2008). A number of factors point 59 towards the hydrocarbon prospectivity of the Lower Ordovician St. George Group, 60 Western Newfoundland, including the proximity of mature sources rocks (organic rich 61 shales in the Green Point Formation) and impermeable cap rocks (Table Point Formation) 62 and the successful drilling of the Port au Port #1 well in 1995 (Fowler et al. 1995; Cooper 63 et al. 2001). In early 2007 extended testing from the reservoir in the upper part of the St. 64 George Group, by Canadian Imperial Venture Corporation, resulted in the production of high API (50-56°) oil (315 barrels per day) and significant volume of natural gas ( $10^6$  ft<sup>3</sup>/ 65 66 day).

The current study is focused on the Watts Bight Formation in the St. George Group, which is of particular interest for hydrocarbon exploration as it is extensively dolomitized and associated with significant porosity development (up to 30%; Cooper et al. 2001). The main objectives of this study are: (1) to identify and characterize (petrographically and geochemically) the dolomitization phases in the Watts Bight Formation; and (2) to investigate the origin and nature of the dolomitizing fluids of each phase to understand the diagenetic controls on porosity development and distribution. The results build upon

74 previously published data from the overlying Aguathuna and Boats Harbour Formations

75 (Azmy et al. 2008, in press) and will contribute to predictive models of the porosity

76 distribution in offshore and deeply buried onland counterparts

77

## 78 Geological setting

79 The lower Paleozoic shelf rocks form part of the outer domain of the Humber zone in 80 Western Newfoundland, the westernmost of five tectono-stratigraphic zones in the 81 Canadian Appalachians (Fig. 1; Williams 1979). The evolution of these rocks during the 82 lower Paleozoic has been discussed by several authors and summarized by Cooper et al. 83 (2001). Significant rifting, associated with the opening of the Iapetus Ocean, began 84 during the late Proterozoic and early Cambrian (Waldron and van Staal 2001; van Staal 85 2005) and is first recorded in Western Newfoundland by late Proterozoic, fault-bounded, 86 terrestrial clastics and volcanics of the lower Labrador Group. These deposits were 87 separated by a 90 million year hiatus from unconformably overlying clastic rocks of the 88 late early Cambrian, upper Labrador Group (James et al. 1989). These mixed clastic-89 carbonate sediments were overlain by a thick (1.5 km) middle Cambrian to lower 90 Ordovician carbonate platform succession, deposited on a passive margin to the south of 91 Laurentia. The middle to late Cambrian Port au Port Group is a narrow, high-energy 92 carbonate platform which evolved into a wide, low-energy carbonate platform 93 represented by the early to middle Ordovician St. George Group (James et al. 1989). A 94 regional unconformity, the St. George Unconformity, marks the top of the St. George 95 Group and a shift from a passive margin to a foreland basin (Knight et al. 1991). The St. 96 George Group is overlain by shelf carbonates of the middle Ordovician Table Head

97 Formation (Stenzel et al. 1990), which were deposited in the foreland basin where they 98 are sporadically overlain by deep water carbonates and shales of the Table Cove 99 Formation (Stenzel et al. 1990). 100 The Ordovician St. George Group forms a sequence of subtidal and peritidal limestones 101 and dolostones and is widely exposed in western Newfoundland. The succession is 102 subdivided, in ascending order, into the Watts Bight, Boat Harbour, Catoche and 103 Aguathauna Formations (Fig. 2; Knight and James 1987) and represents at least two 104 sequences of Tremadoc and Arenigian age, termed megacycles by Knight and James 105 (1987). Each of these megacycles consists of a thin lower peritidal layer, a thick middle 106 subtidal layer and a thick upper peritidal layer (Knight and James 1987). 107 The Watts Bight Formation overlies the largely dolomitized peritidal carbonates of the 108 upper Cambrian to lower Ordovician Berry Head Formation of the Port au Port Group 109 (Knight et al. 2008). It has been mapped throughout western Newfoundland (Knight and 110 James 1987) and is extensively dolomitized in many areas (Knight et al. 2008). On the 111 western shore of Isthmus Bay (Fig. 1), Port au Port Peninsula, a continuous ~69m section 112 of the formation (Fig. 2), described in detail by Knight et al. (2008), consists of two 113 sequences of low-energy bioturbated and mound carbonates overlain by higher energy 114 facies associations of mounds and grainstones. Knight et al. (2008) considered that these 115 sequences represented transgression and drowning of a shallow shelf, followed by 116 gradual accretion into the tidal zone. 117 At Isthmus Bay, dolomitization of the Watts Bight Formation is extensive particularly in 118 the lower and middle members where sucrosic dolostones replace both bioturbated and

119 thrombolitic limestone (Knight et al. 2008). These dolostones are dark-grey, fine to

120 medium grained and commonly have a strong condensate smell. Porosity is locally

121 developed in these dolostones, particularly is a section of dolomitized grainstones from

the top of the middle member and in a thrombolitic dolostone ~12m above the base of theformation.

124

## 125 Methodology

126 The Isthmus Bay section was sampled approximately at intervals of ~  $2m (\pm 0.5m)$ , with 127 each sample selected to represent individual beds and/or any carbonate veins in the Watts 128 Bight Formation. A total of forty three (43) samples were collected. In addition, five 129 samples were collected from the underlying Berry Head Formation of the Port au Port 130 Group. Samples were analysed following the procedure outlined in detail by Azmy et al. 131 (2008). In total 48 stained (Alizarin Red S-potassium ferricyanide; Dickson 1966) 132 conventional thin sections were examined using transmitted light microscopy and 133 cathodoluminescence (CL). Permeability was measured in selected dolomite samples 134 using a computer controlled steady state permeameter (Temco Model MP-401). 135 Microthermometric fluid-inclusion analyses were performed using a Linkam THMSG600 136 heating-freezing stage. Calibration with precision of  $\pm 0.2^{\circ}$ C at  $-56.6^{\circ}$ C and  $\pm 1^{\circ}$ C at 137  $300^{\circ}$ C was conducted using synthetic H<sub>2</sub>O and CO<sub>2</sub> fluid inclusion standards. Salinities 138 were calculated using a program by Chi and Ni (2006) for the system of H<sub>2</sub>O-NaCl-CaCl<sub>2</sub> 139 A polished mirror-image slab of each thin section was also prepared for microsampling. 140 For C and O isotope analyses approximately 220µg of powder sample was reacted in an 141 inert atmosphere with ultrapure concentrated (100%) orthophosphoric acid at 70°C in a 142 Thermo Finnigan Gasbench II and the produced CO2 was automatically delivered to the

source of a Thermo Finnegan DELTA V plus isotope ratio mass spectrometer in a stream

- 144 of helium, where the gas was ionized and measured for isotope ratios. Precision of the
- 145 data is better than  $\pm 0.1\%$  for both  $\delta^{18}$ O and  $\delta^{13}$ C ratios. For major and trace element
- 146 analyses ~ 4mg of sample powder was digested in 2.5% (v/v) pure  $H_3PO_4$  acid for 70-80
- 147 minutes and analysed for Ca, Mg, Sr, Fe and Mn (Coleman et al. 1989) using a
- 148 HP4500plus inductively coupled plasma mass spectrometer (ICP-MS). The relative
- 149 uncertainties of these measurements are less than 5%.
- 150 Based on petrographic examination representative samples of dolomite were selected for
- 151 Sr-isotope analysis at the at the Institut für Geologie, Mineralogie und Geophysik, Ruhr
- 152 Universität, Bochum, Germany, following the procedures outlines in Azmy et al. (2008).
- 153 The <sup>87</sup>Sr/<sup>86</sup>Sr measurements were normalized to NBS 987 values bracketing the samples
- 154 (0.710247) and corrected for deviation from value stated by McArthur (1994).
- 155

#### 156 **Results**

#### 157 Petrography

- 158 Based on petrography and CL, three distinct dolomitization events have been recognized.
- 159 In addition, late-stage calcite diagenetic cementation phases and other diagenetic features
- 160 (i.e. stylolites, sulphides and silicification) were described.
- 161 *Micritic algae, limemud and early calcite cements*
- 162 Micritic limemud (C1) was generally observed in limestone and dolomitic limestone
- 163 samples. Limestones consist of peloidal and intraclastic micrite aggregates, with a
- 164 micritic matrix or very fine-grained limemuds. Rarely skeletal grains (brachiopods and
- 165 crinoids) are preserved, these are invariably replaced by coarser grained (100 to 300µm)

166	anhedral Fe-poor calcite, which may represent some degree of recrystallisation. In
167	addition a calcite cement has been recorded in some of the interparticular pores (C2),
168	which predates any dolomitization and therefore is interpreted to be relatively early.
169	First dolomite
170	The first dolomite (D1) recognized in the samples is characterized by a fine-grained (< 25
171	to 50µm) crystals replacing C1 in dolomitic limestones (Fig. 3a). D1 commonly replaced
172	sedimentary laminations and peloidal and intraclastic micrite aggregates (Fig 3a) and
173	therefore retains the sedimentary structures and gross textures. This may be due to the
174	preferential dolomitisation of lime muds during early dolomitisation (Murray and Lucia,
175	1967). D1 consists of tightly packed anhedral to subhedral rhombs with no visible
176	intercrystalline porosity.
177	Second dolomite
178	A later phase of dolomites (D2) postdates D1, and replaces C1/C2 calcite and D1
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189	later diagenetic dolomite and calcite. Under CL, D2b dolomite displays a dull red
190	luminescence, and commonly shows compositional zoning (Fig. 3e-f). When viewed
191	under ultraviolet (UV) light, D2b rhombs showed yellow/dull luminescence zoning. D2a
192	and D2b dolomite rhombs are also commonly coated by bituminous organic material.
193	Third dolomite
194	The latest void-filling dolomite (D3) occurs in previously uncemented pores and/or
195	fractures. The D3 dolomite is relatively rare and consists of dolomite rhombs, which are
196	euhedral to subhedral and range in size from 100 to $400\mu m$ . It usually displays undulose
197	extinction (Fig. 3d) and commonly has non-planar boundaries, features typical of saddle
198	dolomite (Warren, 2001). D3 dolomite displays a uniform red fluorescence under CL.
199	Other diagenetic features
200	D2 dolomite is both crosscut and bounded by high amplitude microstylolites, which
201	postdate the early burial diagenesis. A translucent, coarse blocky calcite cement (C3) is
202	common and fills pores and fractures still open after D3 precipitation (Fig. 3c). C3 calcite
203	ranges in size from 50 to $400\mu m$ and both Fe-poor and Fe-rich calcite has been recorded.
204	Under CL, it shows a uniform bright yellow luminescence. In addition rare pore-filling
205	sphalerite and micro-quartz have been recorded.
206	
207	Fluid inclusion analysis
208	Fluid inclusions were examined in D2 and D3 dolomite and C3 calcite and
209	microthermometric data is summarized in Table 1. The fluid inclusions in dolomite (D2

- and D3) were hosted in clusters in the core of dolomite crystals or in discrete zones
- 211 within dolomite rhombs, and inclusions were commonly elongate in the direction of

212	growth (Fig. 4a). Inclusions in C3 calcite were found in clusters in the core of crystals
213	and can be very large with respect to their host crystals (up to $50\mu m$ ; Fig. 4b). Therefore
214	inclusions in dolomite and calcite are considered primary in origin and represent samples
215	of fluid trapped during growth (Goldstein 2003).
216	D2 dolomite
217	Fluid inclusions in D2a and D2b dolomite are biphase and range in size from 2 to $20\mu m$ .
218	Homogenization temperatures (T <sub>h</sub> ), representing the minimum estimates of entrapment
219	temperatures, ranged from 44.6 to 108.6°C (average of $81.8^{\circ}C \pm 14.2^{\circ}C$ ; Figs. 5 and 6),
220	with no significant variations recorded between inclusions hosted in the cores and rims of
221	D2b crystals. The initial melting temperatures (T <sub>i</sub> ) ranged from -55.2 to -51.3 $^{\circ}$ C (Table
222	1), corresponding to the eutectic temperature for the $H_2O$ -NaCl-CaCl <sub>2</sub> ±MgCl <sub>2</sub> system
223	(Shepherd et al. 1985). Hydrohalite melting ( $T_m$ (hydrohalite)) occurred between -39.1°C
224	and -31.9°C yielding $X_{NaCl}$ values of between 0.14-0.28. Ice melting temperatures
225	(T <sub>m</sub> (ice)) ranged from -25 to -6.2°C, giving a wide range of fluid salinities (10.1 to 23.3
226	eq. wt% NaCl + CaCl <sub>2</sub> ; Fig. 6).
227	D3 dolomite
228	The fluid inclusions in D3 dolomite are biphase and range in size from 2 to $10\mu m$ .
229	Inclusions homogenize (T <sub>h</sub> ) to the liquid phase between 91.4 and 130.1 $^{\circ}C$ (114 $^{\circ}C$ $\pm$
230	13.3°C; Figs. 5 and 6). Initial ice melting temperatures (T <sub>i</sub> ) of -52 to -50.2°C indicate the

- 231 presence of CaCl<sub>2</sub>  $\pm$  MgCl<sub>2</sub>. T<sub>m</sub>(hydrohalite) of -31.5 to -30.2 was used to calculate X<sub>NaCl</sub>
- of 0.29 to 0.34.  $T_m$ (ice) ranged from -14.7 to -21.3°C, with calculated salinities from 17.8
- 233 to 21.7 eq. wt% NaCl + CaCl<sub>2</sub>.
- 234 *C3 calcite*

235	Fluid inclusions in C3 calcite display a wide range of size (2 to $50\mu$ m) and degree of fill
236	(0.6 to 0.95) and a large number of inclusions show evidence of post-entrapment leaking
237	and/or stretching (e.g. microfractures at edge of inclusions: Fig. 4b). Care was taken only
238	to record microthermometric data from inclusions which displayed small $T_{\rm h}$ variations
239	within an individual inclusion group (e.g. a cluster), usually less than 10°C, suggesting
240	that post-entrapment effects are not important (Goldstein and Reynolds 1994). $T_h$ values
241	range from 68 to 126.7°C (average of 94.1 $\pm$ 17.3°C; Fig. 5). $T_i$ ranged from -55.2 to -
242	51°C (close to the eutectic temperature of the H <sub>2</sub> O-NaCl-CaCl <sub>2</sub> $\pm$ MgCl <sub>2</sub> ). Hydrohalite
243	melting occurred between -26.1 °C and -34.6 °C yielding $X_{NaCl}$ values of between 0.22-
244	0.54. $T_m$ (ice) occurred between -14.7 and -20.1°C corresponding to salinities ranging
245	between 17.9 to 21 eq. wt% NaCl + CaCl <sub>2</sub> .

#### 247 Carbon and oxygen isotopes

248 The carbon and oxygen isotopic compositions of calcite and dolomite analysed are

summarized in Figure 7 and Table 2. Because C1 and C2 were inseparable by

250 conventional microdrilling, they are reported indistinctively and great care has to be taken

in interpreting the data.

252 Calcites

- 253 Fifteen analyses of C1/C2 calcite from the Watts Bight Formation yielded  $\delta^{13}$ C values
- from -0.3 to -2.2‰ and  $\delta^{18}$ O values of -7.4 to -8.9‰. The C3 cement was analysed from
- four samples and gave  $\delta^{13}$ C values of -1.5 to -5.1‰ and  $\delta^{18}$ O values of -7 to -8.5‰.

256 Dolomites

- 257 Five samples of D1 were collected from dolomitic limestones and gave  $\delta^{13}$ C values of -1
- to -1.8‰ and  $\delta^{18}$ O values of -5.5 to -7.4‰. Fourteen D2a and twelve D2b dolomite
- samples yield very similar carbon and oxygen isotope values (Fig. 7, Table 2) with  $\delta^{13}$ C
- values of -0.6 to -1.9‰ and  $\delta^{18}$ O values of -5.5 to -11.4‰. Four analyses of D3 dolomite
- 261 gave  $\delta^{13}$ C and  $\delta^{18}$ O values ranging from -1 to -2.1‰, and -5.6 to -9.6‰, respectively.
- 262

#### 263 Strontium isotopes

- A single sample of D1 dolomite yielded an  ${}^{87}$ Sr/ ${}^{86}$ Sr ratio of 0.709283 ± 0.000007 (Fig 8;
- Appendix A). The range of  ${}^{87}$ Sr/ ${}^{86}$ Sr for D2 dolomite is between 0.709106 ± 0.000007
- 266 and  $0.708953 \pm 0.000007$  and the D3 dolomite yielded values ( $0.709075 \pm 0.000007$  to
- 267  $0.7091 \pm 0.000007$ ) (Fig. 8; Appendix A).
- 268

#### 269 Major and trace element geochemistry

- 270 The distribution of major and trace elements in calcite (C1/C2, C3) and dolomite (D1,
- 271 D2, D3) from the Watts Bight Formation is summarized in Table 2. C1/C2 calcite is
- 272 characterized by Sr concentrations of  $302 \pm 54$  ppm and Mn and Fe values of  $451 \pm$
- 273 685ppm and  $65 \pm 20$ ppm, respectively. Dolomites from the Watts Bight Formation had
- 274 Ca concentrations ranging from 70.9 to 54.2%. Compared to C1/C2 calcite, Sr
- concentrations were depleted (generally <200ppm) in later dolomite and carbonate phases
- 276 (D1, D2a, D2b, D3, C3). D1 dolomites were relatively enriched in Fe (1662 ± 251ppm)
- and Mn (269  $\pm$  226ppm) with respect to C1/C2 calcite. In contrast Fe and Mn
- 278 concentrations were lower and relatively constant between D2a and D2b, with Fe values
- of 765  $\pm$  249 for D2a and 537  $\pm$  258 for D2b and Mn values of 63  $\pm$  14ppm for D2a and

69 ± 20ppm for D2b. For D3 dolomites Fe values range from 357 to 876 ppm and Mn
values range from 59 to 219.

282

#### 283 **Discussion**

In the following section, petrographic and geochemical observations and data are presented in order to support our contention that hydrothermal fluids (e.g., fluids having temperature at least 5°C higher than ambient formation temperature; White 1957; Smith and Davies 2006) circulated in the Watts Bight Formation and are responsible for most of the dolomite observed in the unit.

289

#### 290 **Dolomite petrography**

291 Dolomite 1 (D1) is fabric retentive dolomicrite, which commonly replaces early micrite 292 and micritic peloids and intraclasts (C1) and it appears dull to non luminescent under 293 luminoscope. It predates all other dolomite generations and stylolites and was most likely 294 formed during the early diagenetic history prior to significant burial and compaction. 295 Dolomite 2 (D2) includes 2 subphases, D2a and D2b, and is crosscut by and associated 296 with microstylolites, which is consistent with dolomitization likely during shallow to 297 intermediate burial (e.g., Azmy et al., 2008). D2 dolomites are also often coated by a 298 bituminous material which suggests that hydrocarbons migrated in pore space after D2 299 dolomitization. 300 Petrographic studies of the Watts Bight dolomites have shown that early diagenetic 301 dolomitization (D1) led to a low porosity end-product (< 1%), whereas significant 302 intercrystalline porosity (up to 7%) resulted from D2b dolomitization (Appendix A), as

303	has been recorded from other successions in the St. George Group (Azmy et al. 2008, in
304	press). This may indicate that early shallow dolomitization was extensive and occurred
305	under open-system conditions with the circulation of a large volume of dolomite-
306	saturated diagenetic fluids, so that no open pores were preserved. On the contrary, the
307	higher intercrystalline porosity and non-stoichiometric nature of D2b dolomites is
308	consistent with dolomitization of limestones with a much lower fluid/rock ratio and a
309	limited local supply of $Mg^{2+}$ and $CO_3^{2-}$ ions, as dolomite has a lower molar volume than
310	calcite or aragonite (Warren 2000). The lower porosity associated with D2a dolomite may
311	reflect the replacement of D1 dolomite, where no volume reduction would be expected.
312	A large number of intercrystalline pores in D2 dolomite have been occluded by D3
313	dolomite and C3 calcite (Fig. 4c, e, f), reducing the effective porosities of D2. However
314	Knight et al. (2008) recognized two intervals at Isthmus Bay with increased
315	intercrystalline and vuggy porosity. The upper porous interval is a dolomitized,
316	crossbedded grainstone at the top of the middle member (porosity of 4 to 5%; Appendix
317	A) and the lower porous interval is a thrombolitic boundstone dolomite ~12m above the
318	base of the formation (porosity of ~7%; Appendix A). Permeability measurements from
319	selected dolomites (Appendix A) show that permeability in the upper porous interval is
320	low (<2 mD) and is similar to permeability in other non-porous dolomites. However the
321	lower porous interval is characterized by much higher permeability (10 to 16 mD),
322	indicating a higher degree of interconnectivity between intercrystalline pores in D2b
323	dolomite from this horizon. Cooper et al. (2001) reported similar high porosity and high
324	permeability horizons associated with hydrothermal dolomites in the Watts Bight
325	Formation from a series of exploration wells around Port au Port peninsula. Although

there is no evidence that these porous horizons are laterally continuous this demonstrates
that significant and preserved porosity can be associated with dolomites throughout the
Watts Bight Formation.

329

#### 330 Fluid inclusion data

331 There is an increase in the average homogenization temperature  $(T_h)$  of primary fluid 332 inclusions from D2 dolomite ( $81.8 \pm 14.2^{\circ}$ C) to D3 dolomite ( $114 \pm 13.3^{\circ}$ C). This is 333 consistent with an increase in burial depth during dolomitization and/or hydrothermal 334 fluid influxes resulting in high temperature dolomitization. Maximum burial temperatures 335 of 75°C have been estimated from conodont alteration indices (CAI), acritarch alteration 336 indices (AAI) and graptolite reflectance data from the Lower Ordovician rocks of the Port 337 au Port peninsula (Nowlan and Barnes 1987; Williams et al. 1998). Although the 338 interpretation of this data is not straightforward (Williams et al. 1998) it suggests that 339 most of the dolomitization in the Watts Bight Formation likely proceeded from fluids at 340 temperature significantly higher than formation temperature at the time of alteration and, 341 by commonly accepted definition, can be designated as hydrothermal. In the absence of 342 local igneous intrusions, the most likely source of these hydrothermal fluids is from the 343 upwelling of basinal fluids from depth along extensional to transtensional faults and 344 fractures (Davies and Smith 2006). Basin modelling of Cambrian and Ordovician 345 successions in western Newfoundland have predicted that fault footwalls became the foci 346 for dolomitizing fluids (Cooper et al. 2001) and this is similar to Ordovician 347 hydrothermal dolomites elsewhere in eastern Canada (Lavoie et al. 2005; Lavoie and Chi 348 in press) and the northeastern United States (Smith 2006). Fluid inclusions in C3 calcite

349 are characterized by homogenization temperatures that are higher than maximum burial 350 temperatures (94.1  $\pm$  17.3) but lower than T<sub>h</sub> from D3 dolomite, the late carbonate 351 cementation is likely associated with slighter cooler hydrothermal fluid influxes after 352 dolomitization of the Watts Bight Formation, possibly after a relative tectonic uplift. 353 Primary inclusions in D2 dolomite have a wide range of salinities (9.5 to 25.6 eq. wt%) 354  $NaCl + CaCl_2$ ; Fig. 6) over a narrow temperature range, indicative isothermal mixing 355 between two end member fluids of contrasting salinities (Shepherd et al. 1985). The low 356 salinity fluid may represent residual meteoric fluids, seawater or low salinity groundwater 357 in interstitial pore space. The high salinity fluid most likely represents basinal brines 358 which circulated through the basin along faults and fractures. In the absence of significant 359 volume of evaporite deposits in the Cambrian and Ordovician rocks of western 360 Newfoundland, the origin of these high salinity brines is largely unknown as is common 361 in structurally-controlled hydrothermal dolomite (Davies and Smith 2006). No discernible 362 relationship between T<sub>h</sub> values and calculated salinities was recorded in D3 or C3 hosted 363 inclusions.

364

#### 365 Carbon and oxygen isotope data

366 The  $\delta^{13}$ C data from C1/C2 calcite (Fig.7) show a narrow range of values (-0.3 to -2.2‰).

367 These values are within the range of  $\delta^{13}$ C values for the best preserved carbonates

368 precipitated from Tremadoc seawater (0.0 to -2.5%; Shields et al. 2003) and indicate that

- 369 C1/C2 calcite preserves its carbon isotope signature. Carbon isotope signatures from D1,
- 370 D2 and D3 dolomite overlap those of C1 calcite (-0.6 to -2.1‰), reflecting the low CO<sub>2</sub>
- 371 content of the dolomitizing waters in sedimentary basins (Land 1992), as such, the early

to late dolomitization of the Watts Bight Formation occurred in a largely rock-buffered,closed diagenetic system.

The  $\delta^{18}$ O of D1, D2 and D3 dolomites likely reflect the oxygen isotopic composition of 374 375 the dolomitizing fluids since dolomitization involves large volumes of waters to provide the Mg needed for the process. Therefore, the  $\delta^{18}$ O of dolomites can be used to calculate 376 the isotopic signature of the dolomitizing fluid ( $\delta^{18}O_{\text{fluid}}$ ) and constrain the origin of these 377 fluids. The  $\delta^{18}O_{\text{fluid}}$  can be estimated using the equation of Land (1983), providing that 378 379 the temperature of dolomitization can be obtained from the fluid inclusion 380 homogenization temperatures. The dolomiciritic grain size and fabric retention of D1 381 suggest formation at near surface temperature (<50°C) likely around 25°C (e.g., Budd, 1997; Warren, 2000; Azmy et al., 2001; 2008). Therefore,  $\delta^{18}O_{D1}$  of -5.5 to -7.4% may 382 suggest a parent fluid having  $\delta^{18}O_{\text{fluid}}$  of -6.4 to -9.5% VSMOW (Vienna Standard Mean 383 384 Ocean Water) (Fig. 9). The micritic lime mudstones and other depositional facies in the 385 Watts Bight sediments indicate tropical shallow shelf water conditions. The documented  $\delta^{18}$ O of the best preserved primary calcite precipitated from the Tremadoc seawater 386 387 during the deposition of the Watts Bight carbonates is estimated at  $\sim -9.8\%$  (Veizer et al. 1999; Shields et al. 2003), which translate to  $\delta^{18}O_{\text{seawater}}$  of a warm (20 to 25°C) tropical 388 zone of ~-6.3‰ VSMOW (Shields et al. 2003). Reinterpretation of  $\delta^{18}$ O isotopic data 389 390 from Silurian and Pennsylvanian carbonate fossils (Came et al. 2007) indicates that these values may underestimate both the  $\delta^{18}$ O values and seawater temperature. However the 391 392 very negative <sup>18</sup>O values recorded from Cambrian and Ordovician carbonates (Veizer at al. 1999) are consistent with low  $\delta^{18}$ O values of Tremadocian seawater and any 393 underestimation of  $\delta^{18}O_{\text{seawater}}$  is relatively minor (Came et al. 2007). 394

395	In similar present-day environments the $\delta^{18}$ O of seawater is ~ 0‰ and that of meteoric
396	water is depleted by $\sim 4\%$ (Clark and Fritz 1997). Assuming that the relationship
397	between the $\delta^{18}$ O of Tremadoc meteoric and seawaters were similar to those of our
398	present day environment, the $\delta^{18}$ O of meteoric fluids were likely about -10.3‰ VSMOW.
399	The calculated $\delta^{18}$ O values (-6.4 to -9.5‰ VSMOW) of the dolomitizing fluids of D1
400	(Fig. 9) seems to plot between the estimated $\delta^{18}$ O of the Tremadoc sea and meteoric
401	waters (-6.3 and -10.3 ‰ VSMOW, respectively) and likely indicate a mixing zone
402	model for early dolomitization as has been suggested for early dolomitization elsewhere
403	in the St. George Group (e.g., Azmy et al., 2008, in press).
404	The wide range of $\delta^{18}$ O recorded in D2 dolomite may reflect variations in the temperature
405	of dolomitisation, although this is considered unlikely due to the relatively narrow range
406	of homogenisation temperatures recorded from D2 dolomite. A number of alternative
407	possibilities for this wide range in $\delta^{18}$ O must be considered, including variations in the
408	$\delta^{18}$ O of the dolomitising fluids over time, and fractionation of oxygen isotopes during
409	diagenesis (Land, 1983). Fractionation of oxygen isotopes involves both the
410	recrystallization of D1 dolomite (and a corresponding depletion in $\delta^{18}$ O; Land 1983) and
411	the dolomitisation of C1/C2 (with corresponding enrichment in $\delta^{18}$ O; Land 1983).
412	However due to the complete replacement of precursor calcite and dolomite it is
413	impossible to determine which, if either, of these interpretations are valid.
414	Using the homogenization temperatures (estimates of minimum entrapment temperatures)
415	recorded from primary fluid inclusions in D2 and D3 dolomites as proxies for
416	temperature of dolomitization, the $\delta^{18}O_{fluid}$ has been calculated (Fig. 9). For D2 dolomite
417	$\delta^{18}O_{fluid}$ ranges from -4.5 to 4.7 ‰ VSMOW and for D3 dolomites $\delta^{18}O_{fluid}$ range from

418 1.4 to 8.4 ‰ VSMOW. These values are much higher than would be expected from 419 seawater or meteoric fluids and are consistent with dolomitization associated with an 420 influx of basinal brines, which are commonly enriched in  $\delta^{18}$ O (Goldstein and Reynolds 421 1994). The slightly lower  $\delta^{18}$ O calculated for D2 dolomites may represent mixing of these 422 brines with  $\delta^{18}$ O depleted residual meteoric fluids as was suggested by the wide scatter of 423 fluid inclusion salinity data.

424

#### 425 Strontium isotope data

426 The Sr isotopic signature of replacive dolomite usually reflects the Sr isotopic signature 427 of the precursor carbonate phase (Banner 1995) and therefore fabric retentive D1 428 dolomites formed during early diagenesis they are likely preserve the primary signature 429 of the Sr isotope composition of the original seawater. Due to difficulties with contamination of C1/C2 it was only possible to obtain one D1 sample for <sup>87</sup>Sr/<sup>86</sup>Sr 430 analysis. This sample plotted close to the range of <sup>87</sup>Sr/<sup>86</sup>Sr for early Ordovician 431 432 (Tremadocian) seawater (Fig. 8; Shields et al. 2003) as has been recorded in early 433 dolomite elsewhere in the St. George Group (Azmy et al. 2008; in press) and supports the 434 conclusions the earliest dolomitizing (D1) fluid is Tremadocian modified seawater. Our limited dataset of <sup>87</sup>Sr/<sup>86</sup>Sr signatures for D2 and D3 dolomite indicate significant 435 436 overlap with both D1 and the accepted value for Tremadocian seawater (Fig. 8). This 437 indicates the lack of significant interaction between the dolomitizing fluid with any 438 feldspar (Rubidium)-rich unit (crystalline basement or sandstone). This observation is 439 again supportive of our contention of a largely rock-dominated diagenetic system with 440 the ultimate source of the radiogenic strontium being Tremadocian limestones altered

through circulation of the hydrothermal fluid. This scenario is consistent with previous

442 studies of dolomitization in stratigraphically younger units of the St. George Group

443 (Azmy et al. 2008, in press) but strongly differs from the more open diagenetic conditions

444 recorded from the largely coeval Romaine Formation from nearby Anticosti Island

- 445 (Lavoie et al. 2005; Lavoie and Chi, in press).
- 446

#### 447 Major and trace element geochemistry

448 The major element geochemistry of Watts Bight Formation dolomites (Appendix 1, Table 449 2) show that all dolomites (D1 to D3) are non-stoichiometric and Ca-rich (Ca 450 concentrations range from 70.9 to 54.2%). D1 dolomites are the least stoichiometric (70.9 451 to 64.1% CaCO<sub>3</sub>), which may reflect some contamination from precursor calcite that was 452 very difficult to avoid during microsampling No trends of increasing stoichiometry in 453 hydrothermal dolomites from D2a (early) to D3 (late) has been observed, as would be 454 expected for dolomites formed in open diagenetic systems and continuing dissolution and 455 re-precipitation (Sperber 1984; Montanez and Read 1992). Non-stoichiometric 456 dolomitization indicate formation in a relatively close and rock-buffered diagenetic 457 systems characterized by low volumes of fluid throughout (Sperber 1984; Kirmaci and 458 Akdag 2005).

The trace element signature of D1 dolomites suggest that these dolomites precipitated from a meteoric/seawater mixture, consistent with stable isotope analyses. The Sr concentration of D1 dolomites  $(194 \pm 44 \text{ppm})$  are low compared with those of C1 and C2  $(302 \pm 54 \text{ppm})$ . These are lower than would be expected from dolomites associated with hypersaline waters (<550 ppm; Tucker and Wright 1990) as was believed by earlier

studies (e.g., Lane, 1990). The lower Sr concentration in D1 dolomites may be related to an influx of Sr depleted meteoric waters and are consistent with low Sr contents (70 to 250ppm) recorded from modern dolomites associated with mixed seawater/meteoric water zones (Vahrenkamp and Swart 1994). In addition D1 dolomites are characterised by relatively high Fe and Mn concentrations (1662  $\pm$  251ppm and 269  $\pm$  226ppm respectively), which may reflect an influx of terrestrial derived fluids which tend to be enriched in Fe and Mn (Rifia and Shaaban 2007).

471 Hydrothermal dolomites (D2 to D3) in the Watts Bight Formation have low Sr 472 concentrations (101  $\pm$  59 and 122  $\pm$  46ppm respectively), consistent with replacement of 473 Sr-poor C1 calcite and D1 dolomite. The iron and manganese content of hydrothermal 474 dolomite may reflect, at times, the redox state of the dolomitizing fluids since Fe and Mn 475 are commonly enriched in the later diagenetic phases (Tucker and Wright 1990). D2 and 476 D3 dolomites have low Fe contents (< 1250ppm) when compared to hydrothermal 477 dolomites from elsewhere in the St. George Group (Azmy et al. 2008). In addition the 478 general, but statistically not significant, trend of decreasing Fe from D2a (early 479 hydrothermal) dolomite to D3 (late-stage) dolomite in the Watts Bight Formation (Table 480 2) is contrary to what is normally seen during burial diagenesis (Tucker and Wright 481 1990). This may reflect lower Fe and Mn concentrations due to lack of sources of these 482 elements in the surrounding rocks, particularly under closed system conditions of 483 circulation, rather than variations in the redox state of the hydrothermal fluids.

484

485

#### 487 **Reservoir potential in the Watts Bight Formation**

488 Hydrothermal dolomites and associated limestones have been demonstrated to be one of 489 the most significant targets for hydrocarbon exploration in eastern North America (e.g. 490 Lavoie et al. 2005; Smith 2006; Azmy et al. 2008). Hydrothermal dolomitization during 491 early burial is an important process in the development of secondary porosity and 492 permeability in potential hydrocarbon reservoirs (e.g. Smith 2006; Wierzbicki et al. 2006; 493 Lavoie and Chi in press). Therefore, petrographic and geochemical investigations of 494 diagenesis in these dolomites provide a better understanding of the timing and origin of 495 diagenetic fluids, key controls on porosity evolution and distribution in potential reservoir 496 rocks (Smith 2006; Wierzbicki et al. 2006). 497 This study demonstrates that the pore network and diagenetic evolution of the Watts 498 Bight Formation recorded significant porosity development associated with the migration 499 of hydrothermal fluids and hydrothermal dolomitization. In many locations in Western 500 Newfoundland sucrosic dolomites (similar to D2) in the St. George Group are spatially 501 associated with faults e.g. Daniels Harbour, Port au Choix (Knight et al. 2008). At 502 Isthmus Bay the upper porous level is associated with a normal fault, now filled in with 503 C3 calcite. Although no major faults have been reported from the lower porous level, 504 hydrothermal fluids may have infiltrated along microfractures that have been recognised 505 during petrographic analysis of Watts Bight dolomites. The exact timing of the 506 hydrothermal dolomitization in western Newfoundland is unknown a Devonian age has 507 been suggested by Cooper et al. (2001). This age is supported by Devonian Rb-Sr ages 508 from sphalerite in the Daniel's Harbour MVT deposit in western Newfoundland (Lane 509 1990) which provides an upper age limit for D2 and D3 dolomitization. Potential

510 hydrocarbon charges postdate hydrothermal dolomitization as D2 rhombs are commonly 511 coated by a bituminous material; the original hydrocarbons might have utilized the 512 enhanced porosity and permeability of dolomite horizons in the Watts Bight Formation. 513 Cooper et al. (2001) demonstrated that potential source rock in western Newfoundland 514 entered the oil window in the late Devonian, by which time extensional faulting would 515 have juxtaposed the source rocks of the Green Point Formation and the dolomites of the 516 St. George Group. Lateral and vertical variations in the porosity distribution in the Watts 517 Bight Formation, due to the transition between porous hydrothermal dolomites and non-518 porous dolomites and limestones, would have facilitated the development of diagenetic 519 traps and potential diagenetic reservoirs for hydrocarbon accumulations, as has been 520 noted in other hydrothermal fields in eastern North America (e.g., Davies and Smith 521 2006; Wierzbicki et al. 2006).

522

#### 523 **Conclusions**

524 Petrographic, fluid inclusion, stable isotope and trace element data indicate that there are 525 at least three types of dolomite in the Watts Bight Formation, representing early 526 diagenetic to late-stage hydrothermal dolomites. Early replacement micritic dolomites 527 (D1) are fine grained and indicate that dolomitization began during the earliest stages of diagenesis. The  $\delta^{18}$ O of D1 dolomitizing fluids (-6.4 to -8.4% VSMOW) fall between the 528 529 estimated  $\delta^{18}$ O of the Tremadoc sea and meteoric waters and support a mixing-zone 530 dolomitization, as has been suggested elsewhere in the St. George Group. 531 Later-stage replacement dolomites (D2) are associated with enhancement in 532 intercrystalline porosity and permeability through the development of intercrystalline

533 pores, while latest stage saddle dolomite (D3) and late burial calcite cements,

534 significantly occluded the pores in some horizons. The D2 dolomite crystals are often

535 coated by a bituminous material which suggests possible hydrocarbon charge after D2

536 dolomitization. Fluid-inclusion microthermometric data imply that D2 and D3 dolomites

537 were formed from warm, saline fluids of hydrothermal origin. The  $\delta^{18}O_{\text{fluid}}$  of D2 ranges

538 from -4.5 to 3.6 % VSMOW and for D3 dolomites  $\delta^{18}O_{fluid}$  ranges from 1.4 to 8.4 %

539 VSMOW, suggesting an influx of  $\delta^{18}$ O heavier basinal brines.

540 The high porosity and permeability associated with D2 dolomites has been recorded in

541 other cored sections from the Watts Bight Formation and combined with tight limestone

542 beds, presence of favourable source rocks and thermal maturation, may suggest that the

543 Watts Bight Formation carbonates are possible potential hydrocarbon reservoirs and

suitable targets for future hydrocarbon exploration in Western Newfoundland.

545

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# 694 Figures



- 696 Figure 1: Geological map of the Port au Port peninsula showing locating of the Isthmus
- 697 Bay section (modified from Knight et al. 2008).





699 **Figure 2:** Simplified lithotratigraphy of the St. George Group, Port au Port peninsula

700 with detailed section of the Watts Bight Formation showing the distribution of dolostones

and limestones (adapted from Knight et al. 2008).



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703 **Figure 3:** Photomicrographs of the Watts Bight Formation. (a) Stained thin section

- showing D1 dolomite replacing C1 micrite peloidal and intraclasts; (b) Photomicrograph
- of D2a and D2 dolomite; (c) Stained thin section showing Fe-poor (pink) and Fe-rich
- 706 (purple) C3 infilling pores between D2b rhombs; (d) Stained thin section with D3
- dolomite showing undulose extinction; (e) D2b dolomite; (f) Same view as (a) under
- 708 cathodoluminescence showing zoned orange/red luminescence of D2 dolomite and
- 709 yellow fluorescence of C3 infilling pores.



- 711 **Figure 4:** Photomicrographs showing distribution of fluid inclusions. (a) Biphase
- 712 inclusions in D2 dolomite rhomb. Note inclusion is elongate in the direction of growth of
- the dolomite rhomb. (b) Large biphase inclusion in C3 calcite. Microfractures around
- 714 edge of inclusion may indicate leaking of trapped fluid.





**Figure 5:** Histogram of homogenization temperatures for fluid inclusions in the D2 and

717 D3 dolomites and C3 calcite of the Watts Bight Formation.





the D2 and D3 dolomites and C3 calcite of the Watts Bight Formation.



Figure 7: Oxygen vs. Carbon isotope values for (a) dolomite (D1, D2a, D2b, D3) and



Figure 8: Scatter diagram showing the <sup>87</sup>Sr/<sup>86</sup>Sr values in the Watts Bight dolomites (D1 to D3). The shaded area represents the range of <sup>87</sup>Sr/<sup>86</sup>Sr values for the Tremadoc (Veizer 

et al. 1999; Shields et al. 2003) 



**Figure 9:** Temperature vs.  $\delta^{18}O_{\text{fluid}}$  for various  $\delta^{18}O_{\text{dolomite}}$  derived from the fractionation equation:  $10^3 \text{ In}\alpha_{\text{dolomite-water}} = 3.2 \times 10^6 \text{T}^2 - 3.3$  (Lands, 1983). Hatched areas represent range of temperatures and  $\delta^{18}O$  for various dolomite generations (D1, D2, D3). Values for Tremadoc seawater from Shields et al. (2003). See text for details.

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Phase		Ti	T <sub>m</sub> (hydrohalite)	T <sub>m</sub> (ice)	Salinity (eq. wt% NaCl)	X <sub>NaCl</sub>	T <sub>h</sub> (L)
D2	п	11	7	31	31	7	58
	Mean	-53.3	-34.6	-14.3	17.5	0.22	79.7
	Max.	-51.3	-31.9	-6.2	25.6	0.28	108.6
	Min.	-55.2	-39.1	-25.0	9.5	0.14	44.6
	S.D.	1.4	2.9	5.2	4.5	0.06	13.4
D3	п	3	4	8	8	3	7
	Mean	-51.1	-31.1	-16.9	20.1	0.31	114.0
	Max.	-50.2	-30.2	-14.7	23.2	0.34	130.1
	Min.	-52.0	-31.5	-21.3	18.4	0.29	91.4
	S.D.	0.9	0.6	2.2	1.6	0.03	13.3
C3	п	8	11	15	15	11	28
	Mean	-53.3	-30.9	-16.4	19.7	0.32	94.1
	Max.	-51.0	-26.1	-14.7	22.4	0.54	126.7
	Min.	-55.2	-34.6	-20.1	18.4	0.22	68.0
	S.D.	1.7	2.5	1.7	1.3	0.10	17.3

**Table 1:** Fluid inclusion microthermometric data from the Watts Bight Formation

 $T_i = \text{temperature of first ice melting; } T_m(\text{hydrohalite}) = \text{temperature of hydrohalite melting; } T_m(\text{ice}) = \text{temperature of last ice melting; } X_{\text{NaCl}} = \text{NaCl/ (NaCl + CaCl_2); } T_h(L) = \text{homogenisation temperature (to the liquid phase)}$ 

Phase		CaCO <sub>3</sub> (%)	MgCO <sub>3</sub> (%)	Sr (ppm)	Fe (ppm)	Mn (ppm)	δ <sup>13</sup> C	δ <sup>18</sup> Ο
C1/C2	n	10	10	10	10	10	15	15
	Mean	98.4	1.6	302	451	65	-1.3	-7.9
	Max.	99.2	3.0	390	2251	99	-0.3	-7.4
	Min.	97.0	0.8	225	95	35	-2.2	-8.9
	S.D.	0.7	0.7	54	685	20	0.6	0.4
C3	n	3	3	3	3	3	4	4
	Mean	99.0	1.0	155	155	31	-2.5	-7.7
	Max.	99.4	1.2	192	403	37	-1.5	-7.0
	Min.	98.8	0.6	101	21	24	-5.1	-8.5
	S.D.	0.3	0.3	48	215	6	1.7	0.7
D1	n	4	4	4	4	4	5	5
	Mean	66.0	34.0	194	1662	269	-1.3	-6.7
	Max.	70.9	35.9	257	1920	600	-1.0	-5.5
	Min.	64.1	29.1	156	1323	96	-1.8	-7.4
	S.D.	3.3	3.3	44	251	226	0.3	0.8
D2a	n	8	8	8	8	8	14	14
	Mean	58.3	41.7	79	765	63	-1.3	-9.2
	Max.	62.2	45.6	225	1242	82	-0.6	-6.0
	Min.	54.4	37.8	27	411	45	-1.8	-11.4
	S.D.	3.2	3.2	70	249	14	0.3	1.8
D2b	n	10	10	10	10	10	12	12
	Mean	60.0	40.0	112	537	69	-1.4	-8.2
	Max.	63.7	45.8	183	1126	114	-1.0	-5.5
	Min.	54.2	36.3	32	201	43	-1.9	-11.4
	S.D.	3.0	3.0	58	258	21	0.3	2.0
D3	n	2	2	2	2	2	4	4
	Mean	62.1	37.9	122	617	139	-1.3	-7.5
	Max.	66.1	42.0	154	876	219	-1.0	-5.6
	Min.	58.0	33.9	89	357	59	-2.1	-9.6
	S.D.	5.8	5.8	46	367	113	0.5	1.9

**Table 2:** Ca, Mg, Sr, Fe, Mn,  $\delta^{13}$ C and  $\delta^{18}$ O data for the Watts Bight Formation 

Sample ID	Phase	Formation	Depth (m)	CaCO <sub>3</sub> (%)	MgCO <sub>3</sub> (%)	Sr (ppm)	Mn (ppm)	Fe (ppm)	δ <sup>18</sup> Ο (‰ VPDB)	δ <sup>13</sup> C (‰ VPDB)	<sup>87</sup> Sr/ <sup>86</sup> Sr	Visual estimate of porosity (%)	Permeability (mD)	
WB30	C1/C2	Watts Bight	68	98.50	1.50	239	70	320	-8.41	-1.54		<1		
WB27	C1/C2	Watts Bight	61.6	98.12	1.88	244	65	2251	-8.87	-1.42		<1		
WB26	C1/C2	Watts Bight	60	99.07	0.93	390	35	128	-8.06	-1.53		0		
WB25	D1	Watts Bight	58	64.25	35.75	156	158	1746	-7.36	-1.84				
WB24	D3	Watts Bight	55.9	57.97	42.03	89	59	357	-9.60	-2.07	0.709075	1		
WBDD	D2a	Watts Bight	58.3	54.37	45.63	32	75	1242	-11.43	-1.76		4		
WD22	D2b	Watts Bight	58.5	54.20	45.80	32	74	1126	-11.34	-1.86		4	1.30	
WP21	D2b	Watts Bight	56.8						-7.80	-1.55		<1		
WB21	D2a	Watts Bight	50.8						-9.60	-1.69		1		
WB20	C3	Watts Bight	53.9	98.78	1.22	173	24	21	-7.85	-5.06		0		
WB18	D2a	Watts Bight	50.4	54.82	45.18	41	67	744	-9.16	-1.48		4	1.42	
WB17	D2b	Watts Bight	48.1									4	1.28	
WB16	D2a	Watts Bight	45.8						-10.48	-1.61		<1		
WD15	C1/C2	Watts Bight	13.5	97.02	2.98	275	99	228	-7.63	-2.24		0		
WD15	D2a	Watts Bight	45.5	62.19	37.81	59	52	773	-10.29	-1.44		<1		
WB14	D2a	Watts Bight	42						-6.70	-0.98		<1		
WB13	C1/C2	Watts Bight	40.2	- 40.2	97.80	2.20	225	27	967	-8.18	-2.02		<i>~</i> 1	
WD15	D2a	Watts Bight	40.2						-8.91	-1.12		<b>N</b>		
WB12	C1/C2	Watts Bight	27.0						-7.78	-2.14		~1		
WD12	D1	Watts Bight	51.9	64.69	35.31	173	223	1920	-7.23	-1.21	_	<1		
WB11	D2b	Watts Bight	35.6	57.30	42.70	100	60	326	-8.20	-1.28	0.709015	1		
WDII	D3	Watts Bight	55.0						-8.62	-1.20	0.7091	1		
WB10	D1	Watts Bight	33.6	64.10	35.90	257	96	1323	-6.31	-1.01	0.709283	0		
WB9	D2b	Watts Bight	31.7	60.65	39.35	171	68	586	-6.85	-1.08	0.708953	1		

Appendix A. Samples, description, elemental and isotopic compositions, visual porosity estimates and permeability in the Watts Bight Formation.

Appendix A	(continued)

Sample ID	Phase	Formation	Depth (m)	CaCO <sub>3</sub> (%)	MgCO <sub>3</sub> (%)	Sr (ppm)	Mn (ppm)	Fe (ppm)	δ <sup>18</sup> Ο (‰ VPDB)	δ <sup>13</sup> C (‰ VPDB)	<sup>87</sup> Sr/ <sup>86</sup> Sr	Visual estimate of porosity (%)	Permeability (mD)
WDQ	C1	Watts Bight	20.8	98.41	1.59	335	81	182	-7.96	-1.29		<1	
W DO	D1	Watts Bight	29.0	70.89	29.11	190	600	1660	-6.96	-1.38		<1	
	C3	Watts Bight		98.94	1.06	101	33	41	-7.30	-1.50			
WB-7	D2a	Watts Bight	27.4						-6.54	-0.88		<1	
	D3	Watts Bight		66.14	33.86	154	219	876	-6.17	-1.04			
WDC	C1	Watts Bight	25.1	99.18	0.82	346	74	114	-7.81	-0.27		<1	
WDO	C1	Watts Bight	23.1						-7.81	-0.74			
WB5	D2a	Watts Bight	22.7	60.24	39.76	27	50	411	-11.28	-1.25		3	
WB4	C1	Watts Bight	21.2	99.15	0.85	341	42	108	-7.59	-1.00		1	
WB3	D2b	Watts Bight	18.3						-9.34	-1.18		7	16.50
WB2	D2b	Watts Bight	15.9	62.81	37.19	121	54	492	-7.29	-1.32		3	
WD 1	D2a	Watts Bight	14.6	55.25	44.75	38	54	554	-10.57	-1.39		4	
W D I	D2b	Watts Bight		60.81	39.19	32	54	460	-10.88	-1.73		4	
WBA1	D2b	Watts Bight	13.3						-8.87	-1.57		7	1.95
WBA2	D2b	Watts Bight	11.7									7	10.05
WBA3	D2b	Watts Bight	10						-6.84	-2.29		2	1.90
WDAA	D2a	a Watts Bight	0	62.07	37.93	225	82	940	-6.00	-0.64	0.70909	-1	
WDA4	D2b	Watts Bight	0	60.41	39.59	183	61	423	-5.54	-1.12	0.709106	cstimate of porosity (%)         <1	
WD A 5	D2a	Watts Bight	5.0	59.78	40.22	142	45	694	-7.80	-1.05		.1	
WDAJ	D2b	Watts Bight	5.9	63.72	36.28	134	43	386	-6.76	-1.14		<1	
	C1	Watts Bight		98.30	1.70	313	50	95	-7.42	-1.19		.1	
	C1	Watts Bight		98.51	1.49	315	51	121	-7.61	-0.66		<1	
WBA6	D2a	Watts Bight	4.5	57.36	42.64	64	77	759	-10.13	-1.34			
	D2b	Watts Bight		57.84	42.16	48	91	633	-11.36	-1.47		5	
	D3	Watts Bight							-5.57	-0.99			

Sample ID	Phase	Formation	Depth (m)	CaCO <sub>3</sub> (%)	MgCO <sub>3</sub> (%)	Sr (ppm)	Mn (ppm)	Fe (ppm)	δ <sup>18</sup> Ο (‰ VPDB)	δ <sup>13</sup> C (‰ VPDB)	<sup>87</sup> Sr/ <sup>86</sup> Sr	Visual estimate of porosity (%)	Permeability (mD)
	C1/C2	Watts Bight							-7.54	-1.71			
WBA7	C1/C2	Watts Bight	0.9						-8.12	-1.20		<1	
	C3	Watts Bight		99.39	0.61	192	37	403	-7.04	-1.84			
	C1/C2	Berry Head	0						-7.46	-1.05		-1	
WDAð	D2b	Berry Head	0	63.67	36.33	175	69	201	-6.19	-0.98	0.708984	<1	
WBA9	D2b	Berry Head	-1.6	52.45	47.55	43	49	424	-8.95	-1.59		4	2.00
WBA10	D1	Berry Head	-3.7						-5.54	-1.00		<1	
WBA11	D2b	Berry Head	-5.2	59.03	40.97	120	114	736	-6.76	-1.52		2	

Appendix A (continued)