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3	A re-appraisal of the petrogenesis and tectonic setting of the
4	Ordovician Fishguard Volcanic Group, SW Wales
5	
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19 Abstract

The Fishguard Volcanic Group represents an excellently preserved example of a volcanic 20 sequence linked to the closure of the Iapetus Ocean. This study re-examines the petrogenesis 21 and proposed tectonic setting for the Llanvirn (467-458 Ma) Fishguard Volcanic Group, 22 South Wales, UK. New major and trace element geochemical data and petrographic 23 observations are used to re-evaluate the magma chamber processes, mantle melting and 24 source region. The new data reveal that the Fishguard Volcanic Group represents a closely 25 related series of basalts, basaltic andesites, dacites and rhyolites originating from a spinel 26 lherzolite source which had been modified by subduction components. The rocks of the 27 Fishguard Volcanic Group are co-genetic and the felsic members are related to the more 28 primitive basalts mainly by low pressure fractional crystallisation. The geochemistry of the 29 lavas was significantly influenced by subduction processes associated with a coeval arc, 30 while significant amounts of assimilation of continental crust along with fractional 31 crystallisation appear to have contributed to the compositions of the most evolved lavas. The 32 Fishguard Volcanic Group was erupted into a back-arc basin where extensive rifting, but no 33 true seafloor spreading had occurred. 34

35

36 Keywords: Geochemistry, Magmatism, Back-arc basin, Subduction

37

39 1. Introduction

3

The closure of the Iapetus Ocean (510-410 Ma) was one of the most significant events 40 in the geological evolution of northwestern Europe (e.g., Trench & Torsvik, 1992; Cocks & 41 Torsvik, 2006; Murphy & Nance, 2008; van Staal et al., 2009; Cocks & Torsvik, 2011). 42 However, our understanding of the tectonomagmatic development, both of this ocean and of 43 the marginal subduction zones along which Iapetus oceanic crust was ultimately consumed, is 44 restricted to somewhat limited exposures. The igneous rocks of the Fishguard Volcanic 45 Group in southwest Wales represent an excellently preserved example of rocks associated 46 with the closure of the Iapetus Ocean. 47

During the emplacement of the Fishguard Volcanic Group in the early-middle 48 Ordovician (485-458 Ma), Wales formed part of the southern margin of the closing Iapetus 49 Ocean. Avalonia, (the microcontinent of which Wales was a part), and moved northward 50 from 55°S to 30°S in the period from the Arenig (478-467 Ma) to the Ashgill (451-444 Ma) 51 (Fitton et al., 1982). During this period a number of island arc chains and marginal basins 52 formed above the south-eastward dipping subduction zone on the north side of Avalonia 53 along a NE-SW-trending plate margin (Fitton & Hughes 1970; Phillips et al., 1976; Fitton et 54 al., 1982; Murphy & Nance, 2008; Murphy et al., 2011). Ordovician sequences of both basic 55 and/or silicic lavas are present not only in the Fishguard region but throughout Wales, in 56 areas such as Snowdonia and Ramsey Island (Fig. 1) (Kokelaar et al., 1984a). 57

A re-evaluation of the Fishguard Volcanic Group is required as earlier studies were based on a limited range of geochemical elements. Furthermore, previous geochemical investigations have shown that the rocks are similar in composition to present-day MORB, while also displaying a relative depletion in Nb (Bevins, 1982). However, the use of tungsten carbide crushing mills in these earlier studies may have affected the Nb concentration of the rocks (e.g., Hickson & Juras, 1986) and so to avoid this problem this study has used agate crushing mills. This study presents new major and trace element geochemical data and
petrographic observations to re-evaluate the petrogenesis and tectonic setting of the Llanvirnage (467-458 Ma) Fishguard Volcanic Group.

67

68 2. Local Geology and Previous Work

69 Most of the Ordovician volcanism in west Wales occurred in the period from Arenig to Llanvirn (478-458Ma) (Thomas & Thomas, 1956; Bevins & Roach, 1979). The overall 70 consensus is that the Ordovician rocks of Wales were emplaced in a supra-subduction back-71 arc basin, in a mostly submarine environment on immature continental crust on the south-east 72 side of the Iapetus Ocean (Bevins, 1982; Bevins et al., 1984; Kokelaar et al., 1984a; Leat et 73 al., 1986). All of the Lower Palaeozoic rocks in Wales and the Welsh Borderland were 74 affected by low-grade (zeolite to low grade greenschist facies) metamorphism, primarily due 75 to burial (Bevins & Rowbotham, 1983; Bevins & Robinson, 1988; Robinson et al., 1999). 76

Early work on the geology of the area was reported by Reed (1895), Cox (1930) and Thomas & Thomas (1956). The Fishguard Volcanic Group (Reed, 1985; Thomas & Thomas, 1956; Bevins, 1979; Bevins, 1982; Bevins et al., 1984; Kokelaar et al., 1984a; Leat et al., 1986; Bevins et al., 1991) is located in the Strumble Head to Fishguard area of north Pembrokeshire, Wales (Fig. 1), close to the older Trefgarn basaltic andesite to andesite lavas of Tremadoc age (485-478Ma) which are thought to represent the products of a volcanic arc (Bevins et al., 1984).

Fishguard Volcanic Group samples were collected from an area around the Strumble
Head Peninsula, (Fig. 1, Fig. 2). The volcanic group comprises three formations, from oldest
to youngest: 1) the predominantly rhyodacitic to rhyolitic Porth Maen Melyn Volcanic

Formation, 2) the pillowed basalts of the Strumble Head Volcanic Formation and 3) the
rhyolitic Goodwick Volcanic Formation (Bevins, 1982) (Fig. 2).

The Porth Maen Melyn Volcanic Formation (Bevins & Roach, 1979) is composed principally of rhyolitic tuffs, ash flow tuffs, rhyolite lavas, autobreccias, debris flow breccias and, to a lesser extent, massive and pillowed rhyodacite lavas. Small volumes of intermediate magmas were also emplaced as microtonalite intrusions (Bevins & Roach, 1979).

Conformably overlying this formation is the Strumble Head Volcanic Formation. 93 Basaltic pillow lavas with well-developed inter-pillow breccias, local elongate steeply 94 inclined lava tubes and necking structures are found at the contact with the Porth Maen 95 Melyn Volcanic Formation (Bevins, 1982). The formation also comprises massive lensoid 96 lava sheets, thin hyaloclastites, basaltic Tuffs, rare rhyolitic tuff horizons and high-level 97 intrusive basaltic sheets (Bevins & Roach, 1979). The boundary between this formation and 98 99 the overlying rhyolitic Goodwick Volcanic Formation is marked by a complex interdigitation of lavas and high-level intrusions (Kokelaar et al., 1984b). 100

101 The Goodwick Volcanic Formation progresses stratigraphically upwards through 102 thick rhyolitic domes and flows to autobrecciated rhyolites and fine tuffaceous silicic rocks 103 which are intruded by a thick basic sill. The sill is bulbous at the base and pillowed at the top 104 indicative of having been intruded into wet sediment (Kokelaar et al., 1984b).

The volcanic group is composed of up to 1.8 km of volcanic and intrusive rocks, deposited in locally subsiding basin (Kokelaar et al., 1984a; Kokelaar et al., 1984b). The rocks were extruded in a mostly submarine environment with restricted occurrences of subaerial volcanism (Bevins, 1982; Kokelaar et al., 1984a; Kokelaar et al., 1984b; Kokelaar 1988). Sediment gravity flow deposits are found interstratified with the pillow lavas, although these are not derived from the lavas and so no topographic highs are thought to have
developed (Kokelaar et al., 1984a; Kokelaar et al., 1984b; Kokelaar 1988).

The basaltic and intermediate lavas and intrusions were recognised as tholeiitic by Bevins (1982) and were proposed to be derived by low-pressure fractional crystallisation of a parental magma, which originated in the upper mantle. Bevins et al. (1991) noted that some of the silicic intrusive and extrusive rocks display a calc-alkaline trend as opposed to a tholeiitic trend. It is uncertain whether dacitic and rhyolite magmas were derived from the same source as the basaltic rocks, although limited trace element data for the rhyolites reported by Bevins et al. (1991) are consistent with such an origin.

119

120 **3. Petrography**

Eighteen basic, intermediate and silicic rocks from the samples collected by Bevins (1979) 121 were re-examined petrographically (Supplementary Material 1). Basalt sample SB34 is from 122 a pillow lava and contains tabular clinopyroxene, minor plagioclase, secondary chlorite and 123 pumpellyite. Other pillowed basalts (samples SB31 and SB58) contain zoned clinopyroxene 124 and plagioclase as microphenocrysts in a groundmass of spherulitic (quenched) 125 clinopyroxene and plagioclase. Sample SB33 consists of tabular plagioclase and interstitial 126 clinopyroxene with secondary chlorite and comes from a massive basaltic lava. The basaltic 127 intrusions (samples LG3, LG1 and LG5) contain euhedral, subophitic clinopyroxene, altered 128 tabular plagioclase, abundant chlorite and, in the case of LG5, epidote. Other intrusive 129 basaltic sheets (samples REB166, SB22, SB44, SB28 and SB59) contain albitised plagioclase 130 with subophitic clinopyroxene and chlorite. Samples SB54 and SB55 are also from intrusive 131 basaltic sheets; however they comprise clinopyroxene and plagioclase phenocrysts set in a 132 finer feldspathic groundmass along with Fe-Ti oxides. The rhyolitic flow samples (SA11 and 133

SA5) are fine grained with quartz-feldspar groundmass (most likely recrystallized from glass)
and also contain quartz and epidote veins. Both the rhyodacite lava (sample REB94) and the
microtonalite intrusion (sample REB342) contain plagioclase, quartz, chlorite and rare
biotite. The field relations of these samples are discussed in detail in Bevins (1982).

138

139 4. Analytical techniques

Eighteen samples (2 rhyolites, 2 dacites and 14 basalts) from the original sample set described and collected by Bevins (1979) from a transect through the Fishguard Volcanic Group (Fig 2 and Supplementary Material 1) were re-prepared and re-analysed for major and trace elements.

Following removal of weathered surfaces, the samples were crushed in a steel jaw crusher and powdered using an agate Tema mill at Cardiff University. Major and trace element abundances were analysed using a JY Horiba Ultima 2 inductively coupled plasma optical emission spectrometer (ICP-OES) and a Thermo X7 series inductively coupled plasma mass spectrometer (ICP-MS) at Cardiff University, Wales. Further information regarding methods and instruments are discussed in McDonald & Viljoen (2006).

Accuracy and precision of the data were assessed using the international reference materials NIM-G and JB-1A (Supplementary Material 2). Relative standard deviations show accuracy of 1-5% for most major and trace elements for the standard materials used. Duplicate standard deviations are also within error, ensuring precision. A representative data set can be found in Table 1 and the full data set in Supplementary Material 3.

155

156 5. Geochemical Results

157 **5.a. Element Mobility**

The rocks analysed in this study have all undergone low-grade sub-greenschist facies 158 metamorphism (Bevins & Rowbotham, 1983; Robinson & Bevins, 1986) and this is reflected 159 in the abundance of chlorite observed in the thin sections along with less common 160 pumpellyite, prehnite and epidote. Varying degrees of albitization have affected the rocks and 161 so the concentrations of Al, Ca and Na in particular have been modified (Bevins 1982). 162 Under such metamorphic conditions many elements, in particular the large-ion lithophile 163 elements, become mobile. Accordingly, these elements are not representative of the original 164 magmatic composition of the rocks and so cannot be used to assess the petrogenetic processes 165 that the magmas have undergone. In this study therefore we will use trace elements generally 166 regarded to be relatively immobile during low-grade metamorphism, i.e., HFSE (High Field 167 Strength Elements) and REE (Rare Earth Elements) (e.g., Pearce & Cann, 1973; Wood et al., 168 1979; Merriman et al., 1986). 169

170

171 **5. b. Classification**

A range of rock types is evident in the samples analysed, including basalts, basaltic andesites, dacites and rhyolites (Figs. 3a - c). In Figure 3a the basalts and dacites (with the exception of SB33) predominantly classify as tholeiitic, while the rhyolites are more calcalkaline. However, silica contents are susceptible to mobility during sub-solidus hydrothermal alteration, and so several classification diagrams which are based on relatively immobile elements have also been used, i.e., the Nb/Y vs. Zr/Ti and Co vs. Th diagrams (Figs. 3b, 3c).

In Figure 3b the samples all plot in the subalkaline field as a continuum from basalt to 179 rhyolite, and on this diagram two of the samples (SB59 and SB58) classify as basaltic 180 andesite. On the Co-Th diagram (Fig. 3c) the majority of the samples plot in the calc-alkaline 181 field, with four of the basalts falling in the island arc tholeite field. While the majority of 182 basalts plot close to the dividing line between basalt and basaltic andesite only sample SB55 183 plots in the basaltic andesite field. The rhyolites plot in the dacite/ rhyolite field while the 184 dacites, fall on the andesite side of the boundary between andesite and dacite. The most 185 evolved basalts appear to be samples SB59 and SB55, although the classification diagrams 186 are generally inconclusive in this regard. Overall, the basalts and dacites of the group show a 187 tholeiitic to tholeiitic/calc-alkaline transitional trend while the rhyolites have a more calc-188 alkaline type chemistry. 189

190

191 **5.c. Basalt Geochemistry**

The basalts show an overall trend of increasing SiO₂ with decreasing MgO and increasing TiO₂, K₂O, CaO and Fe₂O₃ with MgO. Al₂O₃ wt.% generally behaves erratically showing no correlation with MgO wt.%. Two trends are observed in the TiO₂, CaO and Fe₂O₃ vs. MgO plots (Fig. 4), one at lower MgO wt.% with a steep increase in the other major elements and the other which shows less variation in the other oxides with increasing MgO wt. %. Sample REB166 is the most primitive basalt in the suite, with a MgO of 10.8 wt. %, SiO₂ of 46.4 wt.% and Fe₂O₃ of 1.0 wt.%.

In Figure 5 selected trace elements are plotted against Zr. When immobile elements are plotted against Zr (itself immobile and incompatible in the basalts), a good correlation within the basalts indicates that the sequence of rocks are possibly co-genetic (Cann, 1970; Hastie et al., 2008). Many of the more immobile elements show a broad positive linear The La/Yb ratios of the basalts range from 1.9 to 3.3. The basaltic samples generally have flat to slightly LREE-enriched chondrite-normalised REE patterns with a slight depletion in the HREE (Fig. 6a). The basalts are somewhat enriched compared to N-MORB, particularly for the more incompatible elements on the left hand-side of the N-MORBnormalised plot and generally have variable negative Nb anomalies, except sample REB166 (Fig. 6b). As the basalts become more evolved so they become more enriched in incompatible trace elements and develop a progressively more marked negative Eu anomaly (Figs. 5e, 6b).

213

5.d. Dacite and Rhyolite Geochemistry

The evolved samples contain more Si and K along with less Ti, Ca and Fe than the 215 basaltic rocks. The major element compositions variations range from e.g. 64.7 - 76.1 wt.% 216 SiO_2 and from 1.3 - 6.3 wt. % K₂O in a linear array (Fig. 4). The trace element variations vs. 217 Zr (Fig. 5) for the more evolved samples are more scattered than for the basalts, although, a 218 broad linear trend is observed for La and Nb (Figs. 5a, 5d). The Zr content increases from the 219 basalts to the dacites before falling again in the rhyolites which may indicate Zr saturation 220 followed by fractionation. The lack of coherent trace element trends both between the basalts 221 and the evolved rocks, and within the evolved rocks themselves is most likely a reflection of 222 Zr's incompatibility in the basalts and dacites and compatibility in the rhyolites. This is 223 especially evident from the Th-Zr variation diagram, which shows a co-genetic trend for the 224 basalts, while the data for the rhyolites and dacites are more scattered (Fig. 5b). 225

226	The La/Yb ratio of the dacites varies from 3.8 - 7.3, while the rhyolites vary from 4.6
227	-6.1. The rhyolites and dacites are significantly more enriched in the LREE [(La/Sm) _{CN} >3]
228	(chondrite normalised; Sun and McDonough, 1989) than the basalts $[(La/Sm)_{CN} < 3]$ but have
229	similar, relatively flat, HREE patterns (Figs. 6a - b). All of the dacites and rhyolites possess
230	negative Nb anomalies (Fig. 6b). The more evolved rocks also show more marked negative
231	Eu anomalies than the basalts indicating separation of plagioclase during low pressure crysta
232	fractionation from a more basic magma (Figs. 5e, 6a).

234 6. Discussion

235

236 6.a. Mantle source composition

Modern day back-arc basins are underlain by both oceanic and continental crust 237 formed by either seafloor spreading, rifting of older arc or continental crust (Sinton et al., 238 2003; Martinez et al., 2006). However, the relative proportions of these types of crust are 239 often a matter of considerable debate, (Stern, 2002). The formation of oceanic crust in back-240 arc basins is thought to be mainly controlled by two different melt generation processes; 241 hydrous flux melting (e.g., Tatsumi & Eggins, 1995) and decompression melting during 242 seafloor spreading (e.g., Langmuir et al., 1992). With increasing distance of the back arc 243 spreading centre from the subduction zone decompression melting tends to dominate over 244 subduction flux melting (e.g., Gribble et al., 1998; Sinton et al., 2003). 245

Factors contributing to the geochemical heterogeneity of back-arc basin basalts include variations in the underlying mantle in terms of its fertility, composition and degree of partial melting, lithospheric thickness, and amount of water present in the system (Pearce & Stern, 2006). Variations in the subduction input, such as the nature and composition of the
subducted materials, will also influence the geochemistry of lava erupted in the back-arc
region. The melting, assimilation and crystallisation history of the lavas will also contribute
to the compositional diversity observed in back-arc basins. The influence of subduction zone
fluids should be greatest during the early stages of basin opening and then diminish as the
basin widens, although this is dependent on the geometry of the basin (Sinton et al., 2003;
Martinez et al., 2006).

The magmas represented by the basalts of the Fishguard Volcanic Group were most likely derived from a shallow, relatively garnet-free, source (spinel lherzolite) as indicated by the flat HREE patterns. This is most likely reflective of the E-W extension of immature continental crust that occurred from the Tremadoc – Caradoc (485-448Ma), resulting in locally subsiding grabens controlled by crustal discontinuities (Kokelaar, 1988).

261 More insights from the REE data for the rocks and so their source can be gained through the use of a Dy/Dy* - Dy/Yb plot (Fig. 7a). Dy/Dy* $(Dy/Dy* = DyN/(La_N^{4/13}))$ 262 $Yb_N^{9/13}$) is a measure of the concavity of a REE pattern (Davidson et al., 2013) and, in 263 addition to helping determine source components, this diagram can help elucidate if 264 amphibole has been significantly involved in the petrogenesis of a magma (either by 265 fractionation or by being residual in the source). Figure 7a shows that the Fishguard lavas and 266 intrusions plot mostly in the MORB field of the diagram and display a general trend of 267 increasing Dy/Dy* with Dy/Yb. Only samples SB59 and SB54 trend towards the amphibole 268 vector, indicating minimal involvement. 269

The Th/La-(Ce/Ce*)_{Nd} diagram (Fig. 7b) which is used to determine the affinity of sedimentary slab components that have contaminated the source region of subduction zone rocks. The majority of the basalts plot in an array between N-MORB and the volcanic detritus field, with some of the more evolved basalts plotting within the continental detritus field,
indicating a variable subduction component within the basalts. This suggests relative
proximity to the continental subduction zone.

All except one of the samples analysed in this study (REB166) are displaced above 276 the MORB-OIB array towards the continental arc region of the Th/Yb – Nb/Yb plot (Fig. 7c) 277 further implying proximity to a subduction zone. The subducting slab does not retain Th and 278 therefore an increase in its concentration in the mantle wedge, and so in back-arc basin 279 magmas, is indicative of input from a subduction zone (Pearce, 2008). An increase in the 280 Th/Yb ratio therefore implies an increasing subduction input, while an increase in the Nb/Yb 281 ratio is more indicative of increasing depth of melting. It is clear that the Fishguard rocks 282 show a variable subduction influence in the basalts with little change in melting depth, in 283 agreement with Figure 7b. Figures 7b and c suggest that sample REB166 contains a minimal 284 amount of subduction component. The sample, highlighted in Figure 6, also shows no Nb 285 depletion or significant Th enrichment seen in the other samples. Typically back-arc basin 286 basalts can contain variable amounts of subduction components (Pearce et al., 2005; Pearce 287 & Stern, 2006). In the case of the Fishguard Volcanic Group the majority of the basalts 288 contain a substantial subduction component. These components are derived during 289 subduction by the release of fluids and/or sediments from the slab or may have been via 290 inherited subduction components in the lithosphere, highlighting the complexity of the 291 292 source.

293

294 6. a. Crustal Processes

Bevins (1982) suggested that the various dacites, rhyodacites and rhyolites exposed in the Porth Maen Melyn and Fishguard area were derived by crystal fractionation from the basic magmas. The relative roles of fractional crystallisation from basalt and generation by
crustal melting for the silicic rocks of the Ordivician marginal basins of Wales have been
debated by many authors (Kokelaar et al., 1984a; Leat et al., 1986; Thorpe et al., 1993)

In order to test these different models both fractional crystallisation and assimilation 300 with fractional crystallisation (AFC) modelling were carried out. Rayleigh fractional 301 crystallisation modelling was undertaken using sample LG3 as a possible parental magma as 302 it has relatively high MgO (9.1 wt. %) and also has low LREE contents. Those models were 303 calculated using PELE, a PC platform version of the silicate liquid crystallization MELTS 304 software program (Boudreau, 1999). The major element geochemical trends were modelled at 305 different pressures and water content parameters, using a quartz – fayalite – magnetite (QFM) 306 oxygen buffer to define oxygen fugacity (fO_2). The modelling at 1kbar predicts initial 307 crystallization of olivine and spinel, until ~12% crystallisation at which point plagioclase 308 begins to crystallise followed by clinopyroxene at ~39% crystallisation. This sequence is 309 generally consistent with that observed in the thin sections. Most of the samples show ophitic 310 clinopyroxene plagioclase relationships with some of the clinopyroxene also containing 311 rounded pseudomorphs of what was previously olivine. The chemistry of this mineral 312 assemblage predicted by PELE was then modelled at 10% fractionation intervals using the 313 relevant distribution coefficients (Rollinson, 1993). 314

The results presented in Figure 8 are of the models which best predict the major geochemical trends observed in the data i.e., low pressure fractional crystallisation (1kbar) in a magma which was either anhydrous or contained 1% H₂O. The basalt trend is more accurately modelled than the dacite/ rhyolite data. The models can generally predict the SiO₂ and K₂O trends of the basalts and some of the more evolved rocks but cannot replicate the highest values. The models predict the low CaO trend of the basalts but have higher CaO than the dacites and rhyolites. The high CaO trend is most likely a result of alteration (Bevins, 1982). The FeO trend, however, is poorly replicated by the models. A low-pressure magma
chamber model is in accordance with previous studies (Bevins et al., 1991) but neither
fractional crystallisation model conclusively fits the observed data.

AFC modelling (DePaolo, 1981) was carried out for trace elements at 1kbar (Fig. 9). A significant limitation to the model is that Zr is incompatible in all instances, whereas in reality it is compatible in the more evolved rocks, meaning that their compositions are not reproduced in the models. Another limitation to the AFC models is that the exact nature of any potential contaminant is not known. An average composition for felsic continental crust has been used (from Rudnick & Gao, 2003) which represents a potential fusible crustal contaminant.

Marked enrichments in incompatible elements (particularly Yb, Tb, Y) observed in rhyolites and dacites (Figs. 9a, 9c, 9d) can be replicated by AFC modelling especially with higher crustal assimilation. The models cannot however replicate the elevated Th, Yb, Tb and Y concentrations of the basalts nor does any one single curve match the trend shown by the rest of the samples (Fig. 9b). These models do, however, suggest a shift from initial fractional crystallisation in the most primitive basalts to assimilation fractional crystallization in the more evolved basalts.

339

340 6.c. Tectonic Reconstruction

The evidence discussed in the previous section suggests that the rocks in this study: a) have been subjected to the influence of subduction zone components; b) have undergone some crustal contamination; and c) the basalts have a source that is compositionally similar to MORB, as evidenced by Figure 7a.

The mantle source composition for the primitive magmas of the Fishguard Volcanic 345 Group has been determined to be a mixture of MORB like-source mantle and subducted 346 sediment. Flat HREE patterns indicate melting took place in the spinel stability field at 347 shallow depths (<60km) and so along with the above indicate contamination of a spinel 348 lherzolite source with subduction components. Contamination of the source by continental 349 volcanic detritus (Fig. 7b) signifies an arc-proximal setting close to a continental margin 350 (e.g., Sinton et al., 2003; Martinez et al., 2006). In addition to this, the presence of spatially 351 and temporally close acidic and intermediate rocks and the lack of formation of true oceanic 352 crust (Kokelaar et al., 1984a) indicate proximity to a volcanic arc. The immature continental 353 crust on which the Fishguard basin formed (Kokelaar et al., 1984a) may have been part of a 354 rifted arc system, which originated from earlier arc volcanism in the Tremadoc represented 355 by the Trefgarn Volcanic Group (Kokelaar et al., 1984b). 356

The suite of investigated rocks shows a compositional range through basalts, basaltic 357 andesites, dacites and rhyolites. The rocks in this suite of lavas and intrusions are most likely 358 compositionally similar magmas, with any scatter likely due to variable subduction inputs, 359 fractional crystallisation and AFC. Fractional crystallisation at 1kbar can explain many of the 360 trends observed in the data. Low-pressure fractional crystallisation also generally agrees with 361 the estimated source depth and with previous studies such as Bevins et al. (1991). There may 362 have also been a hydrous component in the form of fluids or hydrous minerals (Fig. 8) 363 present during fractional crystallisation, which is consistent with a subduction zone setting 364 (Stern, 2002). AFC modelling indicates that crustal contamination also played a significant 365 role in the formation of the more evolved rocks. Furthermore, crustal contamination 366 reinforces the interpretation that the basin was formed on rifted volcanic arc crust (Kokelaar 367 et al., 1984a; Martinez et al., 2006). 368

The Fishguard Volcanic Group is therefore likely to have formed in a back-arc basin setting in which a compositionally continuous and genetically linked suite of lavas were erupted. It is suggested that the Fishguard Volcanic Group formed in a back-arc basin where extensive rifting but no true spreading (and oceanic crust generation) had yet occurred.

373

374 6.c.1 Regional Implications

Other Ordovician lavas erupted in the Welsh marginal basin show geochemical 375 similarities to the Fishguard Volcanic Group. The Ramsey Island rhyolites (Middle Arenig) 376 (Kokelaar et al., 1984a) in particular are markedly compositionally similar to the Fishguard 377 Volcanic Group (e.g., Figs. 7b - c), which may be a reflection of their spatial and temporal 378 proximity (Bevins et al., 1991). The rocks of the Snowdon Volcanic Group, North Wales 379 (Fig.1) (Arenig – early Caradog) (Kokelaar et al., 1984a) are enriched in incompatible 380 elements (Fig. 7c) with moderate to marked Fe enrichments and are transitional between 381 tholeiitic and calc-alkaline in composition (Kokelaar et al., 1984a). Figure 7c shows that the 382 rocks from the Snowdon Volcanic Group also contain generally higher Th/Yb ratios 383 highlighting the more pronounced influence of subduction due to their closer proximity to the 384 volcanic arc (e.g. Sinton et al., 2003). This may be a reflection of the control of crustal 385 discontinuities (steep fractures at shallow levels of considerable length that have undergone 386 repeated tectonic activity) on the volcanic location and development (Kokelaar, 1988). The 387 Fishguard Volcanic Group by comparison shows marked Fe enrichment (Kokelaar et al., 388 1984a) and a tholeiitic to tholeiitic – calc alkaline transitional trend. 389

Figure 10 illustrates the proposed tectonic model for the origin and evolution of the Fishguard Volcanic Group. The proposed model is that the Fishguard Volcanic Group formed in a back-arc basin in which rifting and extension of immature continental crust (which may represent rifted arc crust) had occurred. The locus of extension most likely occurred between
the active and rifted arc, where the system is rheologically weakest (Sdrolias and Müller,
2006). The magmas originated partially from flux melting in the mantle wedge, with this
melting facilitated by fluids expelled from subducting slab sediments. Decompression
melting at relatively shallow depths, associated with the rifting of the basin also took place.
The magmas underwent fractional crystallisation, during which crustal contamination from
the immature continental crust of the rifted arc took place.

There are several other known Ordovician Avalonian terranes that have comparable magmatism and originated in similar tectonic settings to the Welsh marginal basin rocks. Similar terranes from the southern flank of the Iapetus include the Antigonish Highlands Avalonian terrane in Nova Scotia, Canada and the Avoca volcanic rocks, Ireland; both formed due to rifting within an overall arc environment (McConnell et al., 1991; Murphy et al., 2011). These comparable origins therefore reinforce the suggested model of formation for the Fishguard Volcanic Group.

407

408 7. Conclusions

409 1. The Fishguard Volcanic Group formed in a back-arc basin in close proximity to a410 subduction zone.

Geochemical evidence and modelling indicates a shallow (garnet-free) lherzolite
source for the Fishguard Volcanic Group. Subduction inputs and low-pressure (1kbar)
fractional crystallisation are the more influential processes in the petrogenesis of the
basalts, while crustal contamination was a significant processes in the formation of
the more evolved rocks of the Fishguard Volcanic Group.

3. The immature continental crust nature of the basin may be derived from a rifted arc crust (Kokelaar et al., 1984a) which coupled with the suggested shallow source region, indicates that previous extension had occurred in the basin. This is consistent with the evidence for earlier volcanic arc activity in this region during the Tremadoc (485-478Ma) (Kokelaar et al., 1984a).
4. It is therefore suggested that the Fishguard Volcanic Group was generated in a back-

422 arc basin setting where extensive rifting (but no seafloor spreading) has occurred (Fig.423 10).

424

425 8. Acknowledgements

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431 9. References

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636 Figure 2

















661 Figure 7











Figure 1. Map adapted from Bevins et al. (1991) showing the location of the Ordovician

igneous rocks of North Pembrokeshire and the rest of Wales in relation to one another. The

rock types of the study area are also shown.

683

Figure 2. Geological map showing the locations of the samples analysed in this study. Also highlighted are the formations which have been sampled. The non patterned portion of the map represents unrelated rocks and quaternary cover.

687

Figure 3. (a) Miyashiro (1975) basalt classification diagram showing the compositions of the
studied samples. (b) Nb/Y – Zr/Ti classification diagram adapted from Pearce (2008). (c) ThCo discrimination diagram (Hastie et al., 2007). The compositional fields are IAT, island arc
tholeiite; CA, calc-alkaline; H-K, high-K calc-alkaline; SHO, shoshonite. (*indicates that
latites and trachytes also plot in the D/R fields).

Figure 4. Bivariate plots of major element data against MgO (wt.%) for the Fishguard
Volcanic rocks.

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Figure 5. Bivariate plots of trace elements variation with Zr (ppm) for the FishguardVolcanic rocks.

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Figure 6. (a) Chondrite normalised REE and (b) MORB normalised trace element diagrams
 (normalising values from Sun & McDonough 1989). Sample REB166, is highlighted.

Figure 7. (a) A Dy/Dy* - Dy/Y plot of Fishguard lavas adapted from Davidson et al. (2013). 703 DM -depleted mantle, PM - primitive mantle (Sun & McDonough, 1989), GLOSS - average 704 global subducting sediment (Plank & Langmuir, 1998). (b) Th/La-(Ce/Ce*)_{Nd} diagram 705 (Hastie et al. 2013) showing the potential sedimentary slab components that may have 706 contaminated the source region. SSC- HD - slow sediment clay-hydrogenous, SSC-FH - slow 707 sediment clay-fish debris/ hydrothermal. (c) The Th/Yb-Nb/Yb diagram from Pearce (2008). 708 An enrichment in Th relative to the equally-incompatible Nb acts as an effective proxy of 709 subduction input. Representative fields for other Welsh lower Ordovician volcanic complexes 710 are also plotted on (b) and (c) for comparison. The data for Ramsey Island and St Davids 711 Head were obtained from Bevins et al. (1991) and the Snowdon data from Thorpe et al. 712 (1993). 713

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Figure 8. Bivariate diagrams of selected major elements vs. MgO for the Fishguard
volcanics along with modelled fractional crystallisation trends (1kb anhydrous and 1kb with
1% H₂O added) for a parental magma with a composition of sample LG3. Markers on the
modelled trends are at intervals of 10% crystallisation.

720	Figure 9. Bivariate diagrams for representative trace elements and incompatible trace element
721	ratios showing the results for AFC and fractional crystallisation, where 0.3 AFC is equal to
722	30% additional contamination and represents the model which best replicates the
723	geochemical trend of the Fishguard Volcanic Group.
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725	Figure 10. Proposed model for the formation of the Fishguard Volcanic Group. The diagram
726	shows the tectonic configuration of the Iapetus slab subducting below the Avalonia continent,
727	with rollback of the subducting slab facilitating the rifting of the arc and opening of the back-
728	arc basin. This rifting leads decompression melting in the structurally weaker parts of the
729	basin; these magmas mix with melts associated with fluids from the subducting sediments. A
730	small amount of crustal contamination occurs as the melts rise through the rifted arc crust.
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Sample:	LG3	LG1	LG5	REB166	SB22	SB59	SB54	SB34	SB28
Major elem	nents								
(wt.%)									
SiO ₂	48.61	49.63	49.19	46.37	48.17	49.74	54.7	48.83	47.61
TiO ₂	1.40	1.38	1.75	1.44	2.14	1.16	2.47	1.26	1.67
AI_2O_3	14.57	13.5	12.94	15.45	13.95	14.92	11.79	13.96	15.15
MnO	0.20	0.18	0.23	0.17	0.19	0.17	0.24	0.15	0.16
MgO	9.09	7.97	8.12	10.75	5.58	4.98	4.15	6.65	8.13
CaO	9.03	10.60	10.28	8.70	13.29	12.04	5.47	16.42	9.73
Na₂O	3.55	3.52	2.81	2.35	2.12	3.69	4.41	0.66	3.46
K ₂ O	0.60	0.37	0.47	0.75	0.45	0.25	0.10	0.02	0.99
P_2O_5	0.13	0.14	0.19	0.13	0.29	0.17	0.31	0.13	0.21
FeO*	10.17	9.09	11.58	9.28	11.42	6.02	12.52	8.99	8.60
LOI	2.55	2.52	2.16	3.94	2.09	7.00	2.18	3.38	3.80
Total	99.90	98.90	99.72	99.33	99.69	100.14	98.34	100.45	99.51
Trace elem	ents								
(ppm)									
Ве	0.6	0.6	1.7	0.7	1	1	1.7	0.6	0.6
Sc	42	50	48	39	39	33	35	39	38
V	265	278	338	256	354	183	336	251	306
Cr	273	67	91	361	106	132	6	322	333
Со	46	41	98	47	42	43	35	39	64
Ni	24	18	13	52	23	49	16	29	87
Cu	126	105	106	79	84	87	82	133	70
Zn	55	19	49	47	46	32	81	23	64
Ga	18.8	19.2	44	21.4	25.4	25.4	24.5	26.6	23
Rb	13.3	6.8	22.2	10.2	5.8	4.5	1.8	0.2	20.7
Sr	244	293	200	259	490	85	106	363	239
Υ	33	34.8	89	31.3	51.7	64.6	63.3	31.3	43.2
Zr	88.6	95.6	120.8	78.6	136.5	184.8	196.4	69.2	89.5
Nb	5.25	3.76	9.70	9.14	5.82	6.04	8.17	2.77	4.04
Мо	0.32	0.14	0.59	0.14	1.72	0.10	0.13	0.41	0.25
Sn	1.88	1.62	2.64	1.83	0.52	0.67	0.82	0.44	0.54
Cs	1.56	0.85	0.88	1.39	0.46	0.34	2.22	0.30	1.07
Ва	114	81	157	134	160	71	226	7	52
La	6.72	6.12	14.60	5.71	9.44	14.76	18.47	5.59	6.89
Ce	17.38	16.60	40.66	15.07	25.56	37.72	45.39	14.77	19.35
Pr	2.69	2.64	6.59	2.42	3.94	5.35	6.35	2.23	3.06
Nd	13.15	13.10	33.25	12.23	19.95	25.30	29.58	11.12	15.88
Sm	4.14	4.15	10.62	3.83	6.02	7.13	8.11	3.55	5.12
Eu	1.34	1.40	3.37	1.47	2.05	1.77	2.38	1.21	1.81
Gd	4.22	4.54	11.32	4.06	7.14	8.20	9.06	4.12	5.86
Tb	0.81	0.86	2.18	0.79	1.27	1.49	1.57	0.76	1.06
Dy	5.33	5.62	13.93	5.11	8.08	9.49	9.84	4.88	6.79
Но	1.03	1.13	2.75	1.02	1.52	1.84	1.87	0.93	1.26

Table 1 Major and trace element data for the Fishguard Volcanic Group

Er	2.96	3.14	7.94	2.93	4.41	5.56	5.42	2.70	3.62		
Tm	0.46	0.49	1.21	0.45	0.73	0.93	0.88	0.45	0.59		
Yb	2.96	3.10	7.84	2.95	4.57	6.09	5.61	2.87	3.70		
Lu	0.46	0.47	1.19	0.46	0.66	0.88	0.82	0.39	0.53		
Hf	2.88	2.91	3.61	2.75	4.35	6.38	5.83	2.49	3.02		
Та	0.38	0.28	0.62	0.62	0.40	0.36	0.56	0.18	0.23		
Th	2.14	0.71	2.15	0.40	1.05	3.46	4.04	0.72	0.60		
U	0.31	0.28	0.62	0.17	0.34	0.94	1.07	0.27	1.10		
Sample:	SB31	SB58	SB33	SB55	SB44	SA5	SA11	REB342	REB94		
Major elements											
(wt.%)											
SiO ₂	50.16	48.2	61.9	56.37	43.95	76.13	75.4	64.65	67.29		
TiO ₂	2.78	1.27	1.86	1.84	3.54	0.23	0.15	0.95	1.08		
AI_2O_3	12.29	12.87	11.79	15.26	10.74	12	11.64	14.73	14.28		
MnO	0.22	0.15	0.16	0.21	0.28	0.04	0.02	0.16	0.15		
MgO	5.8	5.4	4.12	2.5	7	0.78	0.67	1.41	1.38		
CaO	8.2	11.95	4.87	5.19	9.82	0.35	0.003	0.55	2.8		
Na_2O	3.3	4.24	4.41	6.76	2.26	3.44	2.18	3.55	3.48		
K ₂ O	0.86	0.25	0.68	0.07	0.52	4.79	6.13	3.28	1.26		
P_2O_5	0.29	0.32	0.21	0.52	0.4	0.05	0.01	0.24	0.26		
FeO*	13.36	7.61	7.8	9.32	17.75	2.1	1.05	6.51	6.19		
LOI	2.02	7.96	1.69	1.41	2.35	0.74	0.66	2.39	1.93		
Total	99.28	100.22	99.49	99.45	98.61	100.65	97.91	98.42	100.1		
Trace elem	ients										
(ppm)	1 0	1 0	1 2	1 0	0.0	2.1	2.4	2.0	2.0		
ве	1.3	1.3	1.2	1.3	0.9	2.1	2.1	2.8	2.9		
SC	41	29	41	22	57	4	3	22	23		
V	440	179	325	98	854	10	1	96	129		
Cr	19	122	100	10	48	24	11	23	29		
Co	44	40	36	20	53	1	1	12	11		
NI	5	27	15	2	50	2	4	3	3		
Cu	40	64	20	40	100	31	12	34	39		
Zn	66	37	51	43	59	56	16	23	45		
Ga	23.1	23.6	17.7	26.9	24.5	19.5	19.3	26.8	27.3		
Rb	15.3	3.7	10.9	1.3	7.9	84.4	182.2	105.1	55.0		
Sr	200	133	163	242	193	56	38	101	303		
Y	56.4	81.5	35.7	76.3	47.9	62.3	56.6	59.2	62.0		
Zr	127	160	132	151	76	175	126	315	266		
Nb	6.90	6.12	4.94	5.96	3.25	12.59	8.55	8.17	16.05		
Mo	0.32	1.46	0.14	0.24	0.26	1.61	0.64	0.51	0.66		
Sn	0.60	0.72	0.63	0.57	0.50	3.40	0.73	0.54	0.98		
Cs	0.61	0.92	0.50	0.64	0.83	0.82	0.91	2.25	1.89		
Ва	278	171	191	45	147	752	493	931	594		
La	16.65	15.68	10.71	13.02	8.79	34.85	26.87	38.65	43.38		
Ce	40.01	38.52	26.49	36.32	23.74	75.13	59.46	85.33	93.95		
Pr	5.55	5.54	3.74	5.72	3.66	9.45	7.94	10.34	11.37		
Nd	25.96	27.07	17.76	29.20	19.22	36.66	33.95	41.80	47.02		

Sm	7.23	7.77	4.98	8.86	5.87	8.65	8.48	9.10	10.18
Eu	2.23	2.02	1.61	3.11	1.99	1.13	1.02	2.17	2.12
Gd	7.94	9.05	5.56	10.33	6.78	7.95	8.47	9.29	10.31
Tb	1.41	1.65	0.96	1.84	1.19	1.46	1.48	1.53	1.63
Dy	8.69	11.01	5.93	11.70	7.56	9.27	9.18	9.11	9.63
Но	1.65	2.19	1.10	2.26	1.43	1.85	1.73	1.71	1.76
Er	4.81	6.76	3.12	6.59	4.13	5.50	5.29	5.02	5.16
Tm	0.77	1.16	0.50	1.08	0.66	0.90	0.93	0.80	0.85
Yb	5.04	7.51	3.20	6.96	4.16	5.75	5.87	5.29	5.57
Lu	0.71	1.11	0.44	1.02	0.60	0.87	0.88	0.77	0.79
Hf	4.47	5.55	3.93	4.50	2.43	5.77	4.79	8.57	7.91
Та	0.46	0.30	0.26	0.40	0.21	1.09	0.49	0.44	0.88
Pb	6.76	7.22	5.34	2.45	7.34	18.68	21.89	19.23	24.86
Th	4.03	2.75	2.07	0.93	0.75	15.75	18.31	14.52	15.42
U	1.04	1.34	0.68	0.31	0.22	4.01	4.54	3.40	3.77

746 * denotes total iron