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- 1 Structural characterization by Raman hyperspectral mapping of organic carbon in the 3.46
- 2 billion-year-old Apex chert, Western Australia

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11 Abstract

The 3.46 billion years old Apex Chert, Pilbara Craton, Western Australia, is well-12 known for hosting the oldest, highly disputed microfossils on Earth. This rock has a complex 13 history of thermal alteration that includes circulation of hydrothermal fluids, lower 14 greenschist-facies regional metamorphism, and post-metamorphic weathering by meteoric 15 fluids. Carbonaceous material¹ occurs in the sedimentary stratiform part of the chert as well as 16 17 the underlying intruding hydrothermal black chert veins. In order to identify a least-altered remnant of early life it is necessary to develop a method that enables rapid evaluation of CM 18 structural order on a small spatial scale. Here we present the detailed characterization of CM 19 20 in the Apex chert by Raman hyperspectral mapping. It is shown that this approach gives better 21 estimates of average Raman band ratios than individual point analyses, and it is demonstrated

¹ Abbreviations: CM: Carbonaceous Material

22 that significant differences in structure exist between CM in the stratiform part of the Apex 23 chert and CM in an underlying black chert vein. The large Raman map-based datasets also 24 reveal that significant mixing took place between these two end members CM's, indicating that the Apex chert has been thoroughly altered by hydrothermal fluid circulation. At the 25 26 brecciated intersection between the stratiform chert and the intrusive hydrothermal chert vein 27 very poorly ordered CM was found that is not in line with lower greenschist-facies regional metamorphism. It is speculated here that this CM represents an organic fraction that was 28 29 introduced or thoroughly altered by late stage meteoric fluids. Alternatively, the ubiquitous 30 presence of hematite in this sample caused a perturbation in the Raman spectra of the CM, 31 leading to an artifact in the calculated Raman-based band ratios. Overall it can be concluded 32 that the best preserved CM occurs in the stratiform parts of the Apex chert, while earlier discussions on organic microfossils in this rock often focused on parts of the chert that either 33 represented the hydrothermal veins or the brecciated intersection between the veins and the 34 35 stratiform part.

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37 1. Introduction

Reconstructing the origin and early evolution of life on Earth is a difficult and 38 controversial task that for a large part is based on rare preserved organic microfossils in 39 ancient chert and carbonate deposits (Brasier et al., 2002, 2005, 2006; Lindsay et al., 2005; 40 41 Marshall et al., 2011; Olcott Marshall et al., 2012; Schopf et al., 2007 and references 42 therein). As many of these putative fossils occur in hydrothermally-influenced chert deposits, 43 the origin of the CM is questioned (Brasier et al., 2002, 2005; Lindsay et al., 2005). They 44 could represent sedimentary organic debris but could also have been introduced by circulating 45 hydrothermal fluids. Moreover, processes associated with sediment burial such as diagenesis,

secondary hydrothermal fluid circulation, metamorphism, post-depositional 46 and contamination make any unambiguous interpretation of the origin of the CM tenuous. The 47 prolonged effects of temperature, pressure, and strain have altered organic remains to highly 48 aromatic macromolecular structures (kerogen) and ultimately crystalline graphite (Buseck and 49 50 Huang, 1985; Bustin et al., 1995).

Raman spectroscopy is used since the early seventies for the study of CM as it can 51 rapidly and non-destructively distinguish kerogens from void fillings, modern endolithic 52 organisms, and opaque fluid inclusions. Moreover, it can also probe the degree of structural 53 54 order of the organic matter and, therefore, be used to estimate the peak metamorphic temperature experienced by the host rock (Aoya et al., 2010; Bernard et al., 2010; Beyssac et 55 al., 2002, 2003; Jehlicka and Beny, 1992; Jehlicka et al., 2003; Lahfid et al., 2010; Rahl et al., 56 57 2005; Wopenka and Pasteris, 1991, 1993). Finally, Raman analysis enables imaging of 58 structural heterogeneities on a microscopic scale either inherited from the organic precursor, 59 or formed by secondary processes such as deformation, fluid circulation or adsorption on 60 mineral surfaces (Bustin et al., 1995; Qu et al., 2012; Ross and Bustin, 1990; van Zuilen et al., 2012; Wopenka and Pasteris, 1993). Owing to these different sources of uncertainties, it is 61 62 therefore of crucial importance that Raman-based interpretations rely on a statistically meaningful dataset. For instance, Aoya et al. (2010) showed that the Raman spectral 63 indicators for structural order of CM can be confidently interpreted only by using the mean 64 65 value of at least 25 measurements within a single petrographical thin section. It has been shown in studies on CM in meteorites that hyperspectral Raman mapping can be used to 66 67 generate such large arrays of Raman spectral data (Busemann et al., 2007). This approach has 68 so far not been applied to the description of CMs in metamorphic rocks.

Here we present the application of Raman hyperspectral mapping for the 69 70 characterization of CM in diagenetically and metamorphically complex Archean terrains. The 71 information in map-based Raman datasets is used to generate a statistically meaningful histogram of band-intensity- and band-area-based indicators of structural order for CM that 72 73 occurs in a specific region within a rock thin section. Structural characterization is then based 74 on the information in several thousands of spectra instead of a few point-based analyses. We 75 apply this tool to study the structural variation of CM in the Apex chert. This rock has been 76 the focal point for discussions regarding the oldest microfossils on Earth (Brasier et al., 2002, 77 2005, 2006; De Gregorio et al., 2009; Marshall et al., 2011; Marshall and Olcott Marshall 78 2011; Olcott Marshall et al., 2012, 2013; Olcott Marshall and Marshall, 2013; Pinti et al., 79 2009, 2013; Schopf, 1993; Schopf et al., 2002; Schopf and Kudryatsev, 2012, 2013; Schopf and Packer, 1987). The controversy still exists regarding the distinction between primary 80 81 sedimentary CM and hydrothermally-derived fractions of CM. Our samples are derived from the bedded part of the formation as well as from the strongly brecciated and hydrothermally-82 influenced feeder vein (Fig. 1). An overview is given of the different ways in which Raman 83 spectral indicators for CM structure can be calculated and which one use when peak 84 85 metamorphism has reached only lower greenschist-facies. The application of Raman hyperspectral mapping to obtain large datasets is also discussed. Finally, an assessment of the 86 87 variations in CM structural order within some key regions of the Apex chert is provided.

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2. Theory: the Raman spectrum of carbonaceous material

A Raman spectrum of sedimentary CM is composed of a first-order region (1100-1800 cm⁻¹) and a second-order region (2200-3400 cm⁻¹). In an infinite structure such as pure crystalline graphite, only the G-band (~1580 cm⁻¹) is Raman active (Tuinstra and Koenig,

1970). It results from the vibration of the E_{2g} mode of sp^2 -bonded hexagonal ring structures. 93 The vibration of the A_{1g} breathing mode of sp^2 rings becomes possible if the in-plane crystal 94 domain size (L_a, in Å) of individual graphene layers within the crystal is limited by defects, 95 e.g. in micro-crystalline graphite or in disordered CM. This gives rise to the additional D-96 bands (D for disorder) at ~1350 cm⁻¹ (D1) and at ~1620 cm⁻¹ (D2) (Fig. 2a). Many out-of-97 98 plane defects, related to tetrahedrally coordinated carbons, dangling bonds and hetero-atoms, occur in natural CM. These heterogeneities give rise to a broad D3-band at ~1500 cm⁻¹ 99 (Beyssac et al., 2002; Wopenka and Pasteris, 1993). When the crystallinity of the organic 100 matter is very low, a shoulder can be observed on the D1-band at ~1190-1250 cm⁻¹ (Lahfid et 101 al., 2010; Sadezsky et al., 2005). This band (D4) has been tentatively attributed to sp^2-sp^3 102 bonds or C-C and C=C stretching vibration of polyene-like structure (Dippel and 103 Heintzenberg, 1999). The bands for the second order are located at ca. 2700, 2900 and 3200 104 cm⁻¹. They are attributed to overtone (2700 cm⁻¹: 2xD1, 3200 cm⁻¹: 2xD2) or combination 105 modes (2900 cm⁻¹: G+D) (Sadezsky et al., 2005). 106

Natural CMs are classically characterized by analysis of the first-order spectrum. 107 Since there is considerable overlap between the different disorder-related bands, different 108 109 protocols exist for band assignment and spectral profile fitting. A commonly used protocol assigns a single D-band at ~1350 cm⁻¹ and a G-band at ~1600 cm⁻¹ (Bonal et al., 2006; 110 111 Busemann et al., 2007; Ferrari and Robertson, 2000; Marshall et al., 2007; 2001; Olcott-112 Marshall et al., 2012; Quirico et al., 2003; Wopenka and Pasteris, 1993). From the intensity 113 ratio of these D- and G-bands the degree of structural order of CM can be determined. In 114 chemically pure graphitic materials a general inverse correlation is observed between in-plane crystal domain size L_a (Å) and the intensity-based I_D/I_G -ratio (Tuinstra and Koenig, 1970) or 115 116 area-based A_D/A_G -ratio (Larouche and Stansfield, 2010). However, when very poorly ordered CMs are considered, this relationship becomes inaccurate. Several other parameters have 117

118 therefore been used to describe the degree of maturation in such materials, including D-width (FWHM-D), G-width (FWHM-G), D-position, and G-position (Bonal et al., 2006; Busemann 119 120 et al., 2007; Ferrari and Robertson, 2000; Kelemen and Fang, 2001; Quirico et al., 2003). The most sensible way to study the maturity of poorly organized CM is to look at the width of the 121 D- and G-band in function of their intensity ratio. When poorly ordered CM is progressively 122 123 metamorphized, the width of the D-band and the one of the G-band decrease as I_D/I_G 124 increases. Here, we will use the widths ratio FWHM-D/FWHM-G in order to trace CM 125 maturity.

When CM is poorly ordered, another common protocol decomposes the D- and G-126 127 bands into the individual D1-, D2-, D3-, D4- and G-bands (Aoya et al., 2010; Bernard et al., 2010; Beyssac et al., 2002; Lahfid et al., 2010). From this band assignment several ratios can 128 be calculated that describe the structural order of natural CMs. As the overall process of 129 alteration and graphitization is non-reversible, studies of metamorphosed organic matter have 130 shown that such ratios faithfully record peak-metamorphic temperatures (Beyssac et al., 2002; 131 Lahfid et al., 2010; Rahl et al., 2005; Wopenka and Pasteris, 1993). Beyssac et al. (2002) 132 defined the ratio $R_2(1)$ and using progressively altered CMs in metapelites they were able to 133 134 determine a relationship (2) with peak metamorphic temperature in the range 330°-650°C.

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$$R_2 = A_{D1}/(A_{D1} + A_{D2} + A_G)$$
 (1) with T = - 445 * R_2 +641 (± 50°C) (2)

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Above 650°C, CM has attained the triperiodic structure of pure graphite and R_2 has virtually vanished. In very low-grade metamorphic terrains (T-range between 200° and 330°C), CMs display a broad D1-band, significant D4- and D3-bands, and a relative intense G-band. For this range Lahfid et al. (2010) defined the ratios R_{A1} (3) and R_{A2} (5) and determined relationships (4, 6) with peak metamorphic temperature:

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144	$R_{A1} = (A_{D1} + A_{D4}) / (A_{D1} + A_{D2} + A_{D3} + A_{D4} + A_G) (3) \text{ with } R_{A1} = 0.0008T (^{\circ}C \pm 50) + 0.3758 (4)$
145	
146	$R_{A2} = (A_{D1} + A_{D4}) / (A_{D2} + A_{D3} + A_G)$ (5) with $R_{A2} = 0.0045T (^{\circ}C \pm 50) + 0.27$ (6)
147	
148	The intensity of the D4- and D3-bands can also be very useful as the decrease of these
149	bands relative to D1 is linked to an increase in structural order. Accordingly, we define two
150	simple ratios that describe the degree of structure in poorly ordered CMs:
151	
152	$R_{D3}=I_{D3}/I_{D1}$ (7)
153	
154	$R_{D4} = I_{D4} / I_{D1} (8)$
155	
156	It is important to note that all these parameters and ratios can only properly describe
157	CMs that originate from a common precursor and have experienced the same conditions of
158	maturation. Certain spectral parameters are calibrated as a structural indicator using a specific
159	group of CMs (e.g. a coal series) or a specific type of rock matrix (e.g. metapelites). In
160	natural systems, however, CMs often have an unknown precursor or can represent a mixture
161	of different precursors (Behar et al., 2008; Barker, 1996; Waples, 1994). Moreover, many
162	CMs occur in rocks that have highly complex metamorphic histories. In the present paper a
163	type of CM is described that has been the focus of controversy, since the precursor has been
164	interpreted as remnant cyanobacteria (Schopf and Packer, 1987), methanogens (Ueno et al.,
165	2004), or the products of Fischer-Tropsch synthesis (Brasier et al., 2002, 2005, 2011; Lindsay
166	et al., 2005). It is thus arbitrary what exact spectral parameters and ratios are used to describe
167	the variation in structure in this CM. We therefore decided to apply two different but widely

used band-fitting protocols and to calculate the full range of spectral parameters and ratios.Since the CMs in this study occur in chert matrix, a direct comparison with metapelite-based

170 geothermometers R_{A1} , R_{A2} , R_2 is not possible. We use these ratios as indicators of structural

succession of the second secon

171 order, but only calculate 'tentative' peak-temperatures.

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- 173 **3.** Geological setting and sample description
- 174

175 **3.1 Regional geology**

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The East Pilbara Granite-Greenstone Terrane (Fig. 1), located in Western Australia, 177 represents a continental volcanic plateau consisting of granitoid complexes embedded in the 178 greenstone belts of the Pilbara supergroup that formed between 3.72 and 2.85 Ga (van 179 Kranendonk et al., 2002). It represents one of the three ancient crustal nuclei of the Pilbara 180 Craton within the West Pilbara Granite-Greenstone Terrane and the Kurrana Granite-181 Greenstone Terrane (for further details see van Kranendonk et al., 2002; van Kranendonk, 182 2006). Some of the oldest and best preserved Archean rocks on Earth can be found in the 183 3.515 and 3.427 Ga Warrawoona group, within the lower part of the East Pilbara stratigraphy 184 185 (van Kranendonk, 2006). It consists of a thick section of dominantly basaltic rocks 186 intercalated with subordinate felsic volcanic rocks (tuffs and flows), hydrothermal barite and 187 sulfide deposits, and cherty metasedimentary rocks including carbonate and jasper deposits 188 (van Kranendonk and Pirajno, 2004). Metamorphic conditions range from prehnite-189 pumpellyite to greenschist facies (Brasier et al., 2005; van Kranendonk, 2006).

The 3.46 Ga old Apex Basalt Formation, which forms the middle section of the 190 191 Warrawoona Group, is composed mainly of komatiitic basalts with minor amounts of 192 tholeiitic basalts, dolerite flows and dikes (van Kranendonk, 2000). The formation shows evidence of submarine volcanism, including chilled margins, hyaloclastic breccias, and pillow 193 basalt textures. The Apex Basalt Formation is divided into three structural blocks (North, 194 195 Central and South), which are limited by syn-sedimentary fault systems (Brasier et al., 2005). The Apex chert (Fig. 1b), located near Marble Bar in the Chinaman Creek Area, is one of the 196 197 chert horizons that occur in these blocks. It consists of white and black stratiform chert up to 25 m thickness. The bedding is defined by moderate to good local sorting and orientation 198 which testify to a primary sedimentary origin (Brasier et al., 2005). Results from mapping 199 200 showed that these stratiform cherts were fed by individual swarms of black chert veins that were attributed to a stage of pervasive hydrothermal fluid circulation attending the deposition 201 of the stratiform sedimentary chert (Brasier et al., 2002, 2005; Nijman et al. 1998). 202

203 The contact between the black chert veins and the stratiform chert is marked by 204 strongly brecciated material composed of angular fragments of both stratiform chert and black 205 chert veins within a white to translucid secondary chert matrix of likely secondary 206 hydrothermal origin (Brasier 2002, 2005, 2011; van Kranendonk, 2006). Both the bedded chert and black chert veins are cut across by numerous secondary quartz veins that formed 207 during a later regional metamorphic overprint. Similar stratigraphic relationship between 208 209 stratiform sedimentary chert and intrusive black chert veins were recognized on a regional scale in the different formations forming the Warrawoona Group (van Kranendonk, 2006). 210

Some of the oldest carbonaceous microstructures resembling fossilized bacteria were reported in the black chert (Schopf and Packer, 1987; Schopf, 1993). At the time of discovery, the origin of the chert was thought to be sedimentary and represent a shallow marine

214 depositional setting in which photosynthetic bacteria could thrive (Schopf and Packer, 1987). 215 Brasier et al. (2002, 2005) have subsequently shown that the samples studied by Schopf et al. 216 (2002) were derived from a feeder black chert vein representing a conduit for silica-rich hydrothermal fluids. A study based on Si isotopes confirmed that the Apex stratiform chert 217 218 was derived from seawater silica rather than hydrothermal silica (van den Boorn et al., 2007). 219 In contrast, the silica in the black chert veins precipitated from hydrothermal fluids upon 220 cooling and/or mixing with seawater at depth (van den Boorn et al., 2007). It is highly 221 unlikely that photosynthetic organisms occurred in such a sub-seafloor hydrothermal setting. Rather, it has been suggested that CM was produced abiogenically by Fischer-Tropsch 222 223 synthesis (Brasier et al., 2002; De Gregorio and Sharp, 2003, 2006a, 2006b) or by partial decomposition of iron carbonate (Garcia-Ruiz et al., 2003). As Lindsay et al. (2005) 224 225 suggested, the abiotic organic output of such systems may overwhelm the signatures of primitive life and therefore makes it the most difficult environment in which to recognize a 226 record of the early biosphere. Alternatively, it has been proposed that the 3.5 Ga subseafloor 227 hydrothermal system was inhabited by chemoautotrophic organisms (Ueno et al., 2004), 228 sulphur bacteria (Wacey et al., 2011), and/or hyperthermophile bacteria (Brasier et al., 2002; 229 230 De Gregorio et al., 2009).

It should be stressed that the different rock types forming the Apex Formation have been influenced by multiple episodes of hydrothermal circulation, regional metamorphic overprint and by various extent of weathering alteration, complicating the alteration history of the organic material. Low degree of alteration primarily induced by peak regional metamorphism is expected in the stratiform chert while a more intense thermal alteration induced by hydrothermal activity is expected in and near the feeder black chert veins. The late

stage meteoric alteration may have also affected the overall structure of the originalcarbonaceous material.

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240 **3.2.** Sample description

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242 A total of six samples (sampled in 2002 and 2004) from different parts of the Apex chert (Fig. 3) were selected for analysis. These include: three samples of black chert veins 243 244 (#1, #2 and #6), one from the brecciated zone at the intersection between the stratiform chert 245 and black chert vein (#3), and two from the stratiform chert (#4, #5). All these samples are cut 246 across with white quartz veinlets of secondary origin. Mineralogy was determined by optical microscopy and Raman spectroscopy on polished thin section. Sample #1 consists mainly of 247 248 quartz with minor amounts of randomly distributed CM. The different colors of the matrix are 249 due to the relative amount of CM, the black chert being richer in organic matter than the gray chert. Sample #2 represents an altered green-glassy tuff that has been pervasively silicified by 250 hydrothermal fluids. Sample #3 consists of small blocks of fractured stratiform chert 251 252 composed of quartz, CM, hematite, anatase and minor goethite, embedded in a matrix of 253 black chert composed of quartz, CM and minor hematite, goethite, anatase and pyrite. Samples #4 and #5 are stratiform cherts consisting of alternating bands of CM-rich (black) 254 and CM-poor (white, gray) cherty material with minor hematite, goethite, anatase, and pyrite. 255 256 Sample #6 consists mainly of dark chert with a small amount of CM dispersed in a matrix mainly composed of quartz and minor anatase. This phase is cut across with a white chert. 257 258 Based on this petrographic description a distinction is made here between CM associated with

259	stratiform chert (Fig. 3, white dots) and CM associated with hydrothermal chert (Fig. 3, black
260	dots).

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4. Methods 262

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264 4.1 **Spectral acquisition**

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SCRIP All Raman analyses were performed with a Renishaw In Via Raman spectrometer 266 coupled to an Olympus BX61 confocal microscope and with an Ar-ion-20 mW 267 monochromatic 514 nm as laser source. Laser excitation was adjusted to an on-sample 268 intensity of *ca*. 2 mW (measured with a Coherent Lasercheck Analyser) and focused through 269 a 50x objective to obtain a 1-2 µm spot size. Acquisitions were obtained with a 1800 l/mm 270 grating and 100 cm⁻¹ cut-off edge filter, in 'Streamspot'-mode with an air cooled (-70°C) 271 1024x256 pixel CCD array detector. This method enabled a 2000 cm⁻¹ spectral detection 272 range. Beam centering and Raman spectra calibration were performed daily on a Si-Al 273 microprocessor chip with a characteristic Si Raman band at 520.4 cm⁻¹. A 1 s static mode 274 analysis (fixed at 520 cm⁻¹) was used to tune the focal plane position and optimize the count 275 rate. Point analysis measurements were made in static mode (fixed at 1150 cm⁻¹) for 2x10 s 276 running time and in extended mode for 1×10 s over a 100-4000 cm⁻¹ range. Raman 277 hyperspectral maps were acquired in 'Streamspot' static mode fixed at 1150 cm⁻¹, with a 278 motorized Prior Stage allowing XYZ displacement with step precision better than 1 µm. Each 279 spectrum was acquired at 1x9 s running time. 280

A total of 87 spectra have been acquired by punctual analysis for the 6 samples (19 for #1, 11 for #2, 16 for #3, 13 for #4, 14 for #5 and 15 for #6). Eight maps have been acquired (squares on Fig. 3), one for each sample with the exception of samples #1 and #6, for which two maps have been obtained (see ST1). A total of 33,813 individual spectra have been collected. Spectra obtained by hyperspectral mapping have lower resolution than punctual analysis. In order to obtain good estimation for the calculated ratios, we excluded the spectra in which the intensity of the D1-band was less than 10,000 counts. Moreover, spectra for which one or more of the assigned bands could not be fitted were excluded. Overall, 7,296 spectra have been considered in this study (n#1= 3994, n#2= 101, n#3= 1758, n#4=1225, MAN

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4.2 Spectral decomposition 292

n#5=101, n#6-1=23, n#6-2=82).

293 The recorded spectral data were treated with the software 'Renishaw Wire 3.3'. The 294 baseline due to the background fluorescence must be removed and the 5 bands of the 295 spectrum must be fitted in order to determine the different spectral parameters (height, area, 296 full-width at half maximum (FWHM)) for quantitative description of the CM structure, (Lahfid et al., 2010; Sadezsky et al., 2005). The baseline was subtracted by third order 297 polynomial fit. We performed the subsequent peak fitting and characterization by band 298 299 decomposition based on a Gaussian-Lorentzian function with 10,000 iterations per fit. For 300 comparison with literature data and discussion of the various aspects of CM structural order, 301 we followed two different band-fitting protocols; 1) D1-, D2-, D3-, D4- and G-bands are 302 decomposed (Sadezky et al. (2005) and Lahfid et al. (2010)) (Fig. 2a), and 2) only D- and G-303 bands are taken into account (Bonal et al. 2006; Ferrari and Robertson, 2000; Olcott Marshall 304 et al., 2012; Quirico et al., 2003; Wopenka and Pasteris, 1993) (Fig. 2b). In the first protocol,

the band positions were constrained to lie between 1340-1360 cm⁻¹ for D1, 1615-1625 cm⁻¹ 305 for D2, 1490-1510 cm⁻¹ for D3, 1150-1250 cm⁻¹ for D4 and 1580-1610 cm⁻¹ for G (Fig. 2a). 306 These intervals are based on the mean values given by Sadezky et al. (2005). In addition, the 307 intensity of D2 was constrained to be less than half of that of the G-band. For the D4 band, the 308 FWHM was limited at a maximum of 200 cm⁻¹ in agreement with the interval of FWHM 309 310 determined for D4 by Sadezsky et al. (2005). These constraints were necessary to assure 311 proper band assignment by the fitting software. The same fitting procedure was applied to the 312 Raman hyperspectral map arrays. In the second protocol the band positions were constrained between 1340-1360 cm⁻¹ for D and 1580-1610 cm⁻¹ for G (Fig. 2b). 313

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315 **4.3. Artifacts due to sample polishing**

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Polishing of rock thin sections can induce damage in the CM structure and therefore 317 have a strong influence on the resulting Raman spectra (Beyssac et al., 2003; Mostefaoui et 318 al., 2000; Nasdala et al., 2003; Nemanich and Solin, 1979; Pasteris, 1989; Wang et al., 1989; 319 Wopenka and Pasteris, 1993). To avoid this problem it has been proposed that measurements 320 321 should be performed on chemically isolated CM (Wopenka and Pasteris, 1993), freshly cracked surfaces (Tice et al., 2004), or by focusing the laser beam on CM grains embedded in 322 323 a transparent mineral matrix below the surface of a petrographic thin section (Beyssac et al., 324 2003; Lepland et al., 2011; Nasdala et al., 2003). However, several studies have shown that 325 polishing has little damaging effect on organic matter displaying R_2 ratio \geq to 0.5 (Beyssac et 326 al., 2003; Qu et al., 2011, 2013). To determine the magnitude of polishing artifacts on the 327 Apex chert samples, we performed a specific test on samples #1, #4 and #5. Based on the

standard deviation (2σ), the Raman spectrum parameters of sub-surface CM structures that are fully embedded in the quartz matrix are exactly similar to that of the polished CM structures (Table 1). Based on this observation, we conclude that CM in the Apex chert can be directly studied by surface Raman analysis, and thus that Raman hyperspectral maps reflect real variations in CM structures.

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334 4.4. Artifacts from laser-induced heating

335 CM is very sensitive to laser-induced heating that can provoke severe damages 336 (Everall et al., 1991; Kagi et al., 1994). It leads to a downward shifting for the G- and Dbands. It is possible to prevent these effects with the use of low level laser (<5mW, Beyssac et 337 al., 2003; Everall et al., 1991). We used a 2mW laser intensity, which is comparable with or 338 lower than that used in other Raman spectroscopy-based studies of CM in greenschist-facies 339 metamorphic grade rocks (Lepot et al., 2013; Olcott-Marshall et al., 2012; Tice and Lowe, 340 2004; Wopenka and Pasteris, 1993). In addition, Qu et al., (2013) specifically tested the effect 341 342 of laser heating for intensities up to 5 mW on the surface of CM-rich rocks of the greenschist-343 facies Zaonega Formation, Russia, and found no evidence of thermal damage.

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345 5. Results and discussion

The average Raman band characteristics and calculated band ratio are listed for all samples in Table 2. Individual spectral data are listed in the Supplementary Data (ST3). Protocol 1 was used to enable the calculation of the ratios R_{D3} , R_{D4} , R_{A1} , R_{A2} , and R_2 , while protocol 2 was used to obtain the ratios I_D/I_G , A_D/A_G , and the positions and widths of the Dand G-bands. In order to provide a complete set of Raman characteristics the positions and

widths of the decomposed D1- and G-bands from protocol 1 are also reported. The mean values of the map-based ratios slightly differ from those calculated from punctual analyses (Table 2). Hyperspectral maps derived-ratios have smaller standard deviation leading to a better estimation of the calculated ratio and thus of the ordered/disordered state of the CM in the Apex chert.

All Raman spectra (from point analysis and hyperspectral maps, Fig. 4 and S1, 356 respectively) obtained in this study display well-developed D3 and D4 bands, and have a D1 357 band (FWHM $< 70 \text{ cm}^{-1}$) that is more intense than the G band. These spectral characteristics 358 are indicative for rather poorly ordered CM that is typically encountered in lower greenschist-359 facies Archean terrains (Lepot et al., 2013; Marshall et al., 2007). The spectra of the CM from 360 the brecciated chert (Fig. 4 f, S1f) are also typical of low-ordered CM. However, the intensity 361 362 of D1 is nearly the same as that of the G-band, D1 is wider (FWHM= $96 \pm 22 \text{ cm}^{-1}$), and D3 is 363 higher than for the other samples. All spectra of punctual analyses display well-developed second order bands (Fig. 4). This observation differs from that of Olcott Marshall et al. 364 (2012), in which a secondary phase is described that does not contain bands in the second 365 366 order spectrum.

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5.1. CMs in stratiform cherts (#4, and #5) vs CMs in black chert veins (#1)

Some important spectral differences are observed between CMs from the stratiform chert and CMs from the black chert vein. CM spectra for black chert vein (Fig. 4 a-c, Fig. S1 a-c) have always a narrower D1-band (55-60 cm⁻¹) than CM spectra of the stratiform chert (Fig.4 d-e, S1de, 64-67 cm⁻¹). Moreover, black chert vein spectra display less intense D3- and D4-bands (Fig. 6).

374 A distinction between the stratiform chert CM (#4, #5) and the black chert vein CM 375 (#1) is also observed in several of the calculated Raman spectral ratios (Table 2). This is most 376 clearly seen when R_2 is plotted against I_D/I_G (Fig. 5a). Samples #4 and #5 display lower I_D/I_G and R₂ ratios than sample #1. Since R₂ decreases with increasing degree of order in the CM, 377 378 lower R₂ ratios for the stratiform chert CM imply that the CM is more organized than that of the black chert veins. However the spectra of the stratiform cherts have a lower RA1 and ID/IG 379 than those of the black chert veins (Fig. 5b). Since RA1 increases with increasing degree of 380 381 order (Lahfid et al., 2010), lower RA1 values are in agreement with a more disordered CM in 382 the stratiform cherts.

This apparent contradiction can be explained if the structural order of these CMs 383 covers the upper limit of the RA1 ratio (Lahfid et al., 2010) and the lower limit of the R2 ratio 384 385 (Beyssac et al., 2003). In metapelitic rocks this would correspond to peak metamorphic 386 temperatures that range from slightly below to slightly above 330°C. The RA1 and R2 ratios 387 are therefore not well suited to estimate the degree of organization of CM in these rocks. The 388 ratio I_D/I_G is also problematic, since it increases with increasing structural order until a certain 389 maximum is reached, and then decreases with increasing structural order until the D-band is absent (Busemann et al., 2007; Ferrari and Robertson, 2000). Stratiform cherts CMs have 390 lower I_D/I_G ratios that black chert CMs (Fig. 5a-b), and it is not clear whether this reflects a 391 higher or lower degree of structural order. The only effective indicators are then the positions 392 393 and widths of the D- and G-bands, or - in case of total decomposition of the first-order spectrum – the presence of the specific disorder-related bands D3 and D4. The FWHM of the 394 395 D1 band and the intensities of the D3 and the D4 bands are significantly higher in the CM 396 spectra of the stratiform cherts (Fig. 6a-b) than in those of the black chert veins. As shown in 397 Figure 6c, the stratiform chert CM displays high R_{D3} and R_{D4} ratios of 0.10 and 0.09

respectively, which correspond to a low structural order. In contrast, the black chert vein CM have lower R_{D3} and R_{D4} ratios of 0.06 and 0.06, respectively, which are indicative of an higher structural order. FWHM-D, R_{D3} and R_{D4} are then in good agreement, and indicate that the CM in the stratiform cherts is more disordered than that in the black chert veins.

402 While the punctual analyses reveal two distinct CMs within the samples #1, #4 and #5, the large datasets of the hyperspectral maps show a considerably more complex situation. The 403 modes of the normalized histograms for R_{D3} ratio are different for sample #1 and #4/5 (Fig. 7) 404 405 in line with two populations of CM, but a considerable overlap between the population is 406 observed as well. The non-gaussian distributions of the different calculated ratios (normality tested with the Kolmogorov-Smirnov test) show an asymmetry in the low values of RD3 for 407 samples #4/#5 and in the high values of RD3 for sample #1(Fig. 7a). These asymmetric 408 409 distributions could reflect the preservation of a primary phase in the black chert despite the 410 influence of hydrothermal fluids and the influence of the hydrothermal fluids that crossed the 411 stratiform chert. There is thus no distinct separation between CM types, but rather a 412 continuum of mixed CM components. On average, the most altered CM particles occur within 413 the deep regions of the hydrothermal black chert veins whereas the least altered are preserved 414 in the stratiform cherts. This suggests either that there was a significant difference between 415 the organic precursors in the deep hydrothermal veins and the organic precursors in the stratiform chert, or that CM in the hydrothermal fluids experienced a higher peak temperature 416 then the CM in the stratiform chert. 417

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5.2. Sample-scale heterogeneities in sample #6

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420	The Raman hyperspectral maps enable the characterization of sample-scale structural
421	heterogeneities. This is most clearly seen in the spectral data of sample #6. Two maps have
422	been obtained for sample #6; one in the dark grey chert (#6-1, Fig. 3f) and one in the
423	secondary cross-cutting white chert (#6-2, Fig. 3f). The map-derived R_{D3} and R_{D4} values for
424	the darker chert are in the range of the stratiform cherts (0.08 and 0.09, respectively; Fig. 7b).
425	The map-derived ratios in the white chert portion of the thin section are lower (0.06 and 0.07,
426	respectively) and fall in the range of black chert veins (Fig. 7b). They could therefore reflect
427	an episode of channeled hydrothermal fluid infiltration within the stratiform cherts. This
428	interpretation is further supported by the intrusive character of the white chert within the dark-
429	colored chert seen at the thin-section scale (Fig. 3) and by the intrusive character of the black
430	chert veins within the stratiform chert seen at the outcrop scale (Fig. 1).
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5.3 High-temperature hydrothermal alteration in sample #2. 432

The ratios R_{D3} and R_{D4} of sample #2, calculated from the hyperspectral map data, are 433 434 much lower (0.03 and 0.03, respectively) than those calculated for sample #1 (Fig. 7a). These low values are consistent with a higher peak temperature, likely reflecting a different 435 generation of hydrothermal alteration overprint. However, the spectral data from punctual 436 analyses strongly differs from this trend. All but two points in this dataset have R_{D3} and R_{D4} 437 values that fall in the range of the stratiform chert CM. Apparently, by chance some low-438 439 ordered CM was analyzed by punctual analysis, which does not reflect the bulk CM. This 440 discrepancy demonstrates the limited use of a few punctual analyses, as they reflect only a 441 limited range in the CM structural variability.

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443 **5.4 Tentative geothermometry**

444 As discussed in theory part, the ratios RA1, RA2, and R2 can only be used tentatively as 445 indicators of peak-temperature. These geothermometers were calibrated with CMs that 446 represent a common precursor and occur in metapelitic metamorphic rocks (Beyssac et al., 447 2002; Lahfid et al., 2010). The CMs that are studied here do not necessarily represent a 448 common precursor, and occur in cherts. Regional metamorphism and local hydrothermal fluid circulations have resulted in a wide range of peak temperatures of the Apex chert CM through 449 450 time. As follows from the discussion in paragraph 5.1 this temperature range apparently spans 451 the higher end of the R_{A1} geothermometer and the lower end of the R_2 geothermometer. This implies that the two thermometers based on R2 and RA1 should be used for temperature 452 453 estimates of samples #1, #2, and samples #4 and #5, respectively. As shown in paragraph 5.2., 454 sample #6 consists of both a stratiform phase and an infiltrating hydrothermal phase. For this 455 sample, the R2-thermometer should be used for the CM in white chert (#6-2) and the RA1thermometer for the CM in dark gray chert (#6-1). Using equation 2, a tentative temperature 456 457 of 350°C is obtained for sample #1 and of 360°C for #2 and #6-2. Using equation 4 a tentative 458 temperature of 300°C is obtained for samples #4 and #5 and of ~265°C for #6-1. It is 459 speculated here that these temperatures indeed could represent the peak temperature of hydrothermal fluids and of regional metamorphism, respectively. 460 These ranges of temperatures are in line with temperature estimated from the mineralogical assemblages (van 461 Kranendonk, 2006). 462

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5.5 A disordered pool of Organic Matter in Apex chert

The punctual and the hyperspectral map analyses have shown that there is significant variability in CM structure throughout the Apex chert and the associated hydrothermal chert veins. The lowest degree of structural order is preserved in the stratiform chert (samples #4, #5 and #6-1), and appears to have experienced only greenschist-facies regional metamorphism. The highest degree of structural order is preserved in the cross-cutting chert veins (samples #1 and #2), and represents temperatures that are slightly higher than that of regional metamorphism.

In a R_{D4} vs R_{D3} diagram (Fig. 8a), these different CMs define a linear correlation of 472 473 slope ~ 1 . In contrast, analyses performed on sample #3 show a random distribution that strongly deviates from this linear trend (Fig. 8a). The spectral data of sample #3 show high 474 475 FWHM-D/FWHM-G (1.1-2.1) and low I_D/I_G (0.8-1.2) relative to the stratiform chert CM 476 (FWHM-D/FWHM-G =1.2-1.45; I_D/I_G =1.2-1.5) and black chert vein CM (FWHM-D/FWHM-G =1.1-1.3; I_D/I_G =1.35-1.65) (Fig. 8b). This indicates that the CM of sample #3 is 477 478 less organized than that of the stratiform cherts. The random pattern of sample #3 in the diagram RD3 vs RD4 indicates a lack of correlation between the bands D3 and D4. 479 480 Importantly, the CM in sample #3 is in close association with randomly distributed hematite of secondary origin. It was shown by Marshall et al. (2011) that one of the Raman bands of 481 hematite (1320 cm⁻¹) is partially overlapping with the D-band of CM. The cloud pattern 482 defined by sample #3 (Fig. 8a) may therefore be an artifact of the Raman-band-fitting 483 484 protocol. Alternatively, this degree of disorder in the CM of sample #3 is real, and reflects features inherited either from different metamorphic/hydrothermal overprints or from a late 485 stage meteoric alteration. 486

487 A similar 'loss of order' due to post-emplacement phenomena could explain the 488 Raman spectra reported by Olcott Marshall et al. (2012). In their study, the primary CM

phase (E1-E2) is concordant with the signature of 'feeder vein chert CM' and the secondary CM phase (E3) is concordant with that of 'stratiform chert CM' when plotted in a I_D/I_G vs FWHM-D/FWHM-G diagram (dotted lines, Fig. 8b). The occurrence of hematite in the sample analyzed by these authors suggests that this CM could have been influenced by different metamorphic/hydrothermal overprints or a late stage contamination during meteoric alteration.

Likewise, the spectrum of the CM in the microfossils described by Schopf et al. (2007) is, to the naked eye, very similar to the spectra of the sample #3 (Fig. 4f). Indeed, the D-band displays the same pointy shape, ID~IG and FWHM-D>1.5FWHM-G. The field where this spectrum is the more likely to fall is the upper range of the diagram (crosshatched zone, Fig. 8b). This suggests that the CM in putative microfossils can represent altered organic matter that bears little information on the original precursor.

501

502 **7. Conclusions**

Many Archean chert deposits have experienced sub-seafloor hydrothermal fluid 503 504 circulation followed by low-grade regional metamorphism. The CM in such rocks can thus 505 display various degrees of structural ordering. In rocks with a complex thermal history such as the Apex Chert and particularly when peak metamorphism has reached only lower 506 507 greenschist-facies, it is essential that proper and representative Raman spectral data of each 508 stage of thermal alteration are obtained. We demonstrated here that this depends critically on a precise and complete band decomposition protocol, use of the appropriate ratios, and 509 510 acquisition of a statistically meaningful set of spectra.

511

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518 9. References

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518	9. References
519	Aoya M., Kouketsu Y., Endo S., Shimizu H., Mizukami T., Nakamura D., and Wallis S.
520	(2010) Extending the applicability of the Raman carbonaceous-material
521	geothermometer using data from contact metamorphic rocks. Journal of Metamorphic
522	Geology 28 , 895-914.

- Barker C. (1996) Thermal modeling of petrol generation: theory and application. 523 524 Developments in petroleum science 45, Elsevier.
- Behar F., Lorant F. and Lewan M. (2008) Role of NSO compounds during primary cracking 525 of a type II kerogen and a Type III lignite. Organic Geochemistry **39**, 1-22. 526
- Bernard S., Beyssac O., Benzerara K., Findling N., Tzvetkov G., and Brown Jr G. E. (2010) 527 XANES, Raman and XRD study of anthracene-based cokes and saccharose-based 528 chars submitted to high-temperature pyrolysis. Carbon 48, 2506-2516. 529
- 530 Beyssac O., Goffé B., Chopin C., and Rouzaud J.-N. (2002) Raman spectra of carbonaceous material in metasediments: a new geothermometer. Journal of Metamorphic Geology 531 20, 859-871. 532
- 533 Beyssac O., Goffé B., Petitet J.-P., Froigneux E., Moreau M., and Rouzaud J.-N. (2003) On the characterization of disorded and heterogeneous carbonaceous materials by Raman 534 535 spectroscopy. Spectrochimica Acta Part A 59, 2267-2276.
- 536 Bonal L., Quirico E., Bourot-Denise M., and Montagnac G. (2006) Determination of the petrologic type of CV3 chondrites by Raman spectroscopy of included organic matter. 537 Geochimica et Cosmochimica Acta 70, 1849-1863. 538
- Brasier M. D., Green O. R., Jephcoat A. P., Kleppe A. K., van Kranendonk M. J., Lindsay J. 539 F., Steele A., and Grassineau N. V. (2002) Questioning the evidence for Earth's oldest 540 fossils. Nature 416, 76-81. 541

Brasier M. D., Green O. R., Lindsay J. F., McLoughlin N., Steele A., and Stoakes C. (2005) 542 543 Critical testing of Earth's oldest putative fossil assemblage from the 3.5 Ga Apex chert, Chinaman Creek, Western Australia. Precambrian Research 140, 55-102. 544

545	Brasier M. D., Green O. R., Lindsay J. F., McLoughlin N., Stoakes C. A., Brasier A. T., and
546	Wacey D. (2011). Geology and putative microfossil assemblage of the c. 3460 Ma
547	'Apex chert', Chinaman creek, Western Australia - A field and petrographic guide.
548	Geological Survey of Western Australia, Record 2011/7, 60p.
549 550 551	Brasier M. D., McLoughlin N., Green O. R. and Wacey D. (2006) A fresh look at the fossil evidence for early Archean cellular life. <i>Philisophical Transactions of the Royal Society B</i> 361 , 887-902.
552 553	Buseck P. R. and Huang BJ. (1985) Conversion of carbonaceous material to graphite during metamorphism. <i>Geochimica et Cosmochimica Acta</i> 49 , 2003-2016.
554 555 556	Busemann H., Alexander C. M. O. D., and Nittler L. R. (2007) Characterization of insoluble organic matter in primitive meteorites by microRaman spectroscopy. <i>Meteoritics & Planetary Science</i> 42, 1347-1416.
557 558	Bustin R. M., Ross J. V., and Rouzaud JN. (1995) Mechanisms of graphite formation from kerogen: experimental evidence. <i>International Journal of Coal Geology</i> 28 , 1-36.
559 560 561	De Gregorio B. T. and Sharp T. G. (2003) Determining the biogenecity of microfossils in the Apex chert, Western Australia. Thirty-fourth Lunar and Planetary Science conference, Houston, Texas, Abstracts, 1267.
562	De Gregorio B. T. and Sharp T. G. (2006a) The structure and distribution of carbon in 3.5 Ga
563	Apex chert: implications for the biogenecity of Earth's oldest putative microfossils.
564	<i>American Mineralogist</i> 91 , 784-789.
565	De Gregorio B. T. and Sharp T. G. (2006b) Possible abiotic formation of kerogen-like carbon
566	in the Strelley Pool chert. Thirty-seventh Lunar and Planetary Science conference,
567	Houston, Texas, Abstracts, 2318.
568 569	De Gregorio B. T., Sharp T. G., Flynn G. J., Wirick S., and Hervig R. L. (2009) Biogenic origin for Earth's oldest putative microfossils. <i>Geology</i> 37 , 631-634.
570	Dippel B. and Heintzenberg J. (1999) Soot characterization in atmospheric particles from
571	different sources by NIR FT Raman spectroscopy. <i>Journal of Aerosol Science</i> 30,
572	Supplement 1, S907-S908.
573	Everall N. J., Lumsdon J., and Christopher D. J. (1991) The effect of laser-induced heating
574	upon the vibrational raman spectra of graphites and carbon fibres. <i>Carbon</i> 29, 133-
575	137.
576 577	Ferrari A. C. and Robertson J. (2000) Interpretation of Raman spectra spectra of disordered and amorphous carbon. <i>Physical reviews</i> 61 , 14095-14107.
578	Garcia-Ruiz J. M., Hyde S. T., Carnecup A. M., Christy A. G., van Kranendonk M. J., and
579	Welham N. J. (2003) Self-assembled silica-carbonate structures and detection of
580	ancient microfossils. <i>Science</i> 302 , 1194-1197.
581	Jehlicka J. and Bény C. (1992) Application of Raman microspectrometry in the study of
582	structural changes in Precambrian kerogens during regional metamorphism. Organic
583	Geochemistry 18, 211-213.

584 585 586	Jehlicka J., Urban O., and Pokorny J. (2003). Raman spectroscopy of carbon and solid bitumens in sedimentary and metamorphic rocks. Spectrochimica Acta Part A: Molecular and Biomolecular Spectroscopy 59, 2341-2352.
587 588 589 590	 Kagi H., Tsuchida I., Wakatsuki M., Takahashi K., Kamimura N., Iuchi K., and Wada H. (1994) Proper understanding of down-shifted Raman spectra of natural graphite: Direct estimation of laser-induced rise in sample temperature. <i>Geochimica et Cosmochimica Acta</i> 58, 3527-3530.
591 592	Kelemen S.R. and Fang H.L. (2001) Maturity trends in Raman spectra from kerogen and coal. Energy &Fuels 15, 653-658.
593 594 595	Lahfid A., Beyssac O., Deville E., Negro F., Chopin C., and Goffé, B. (2010) Evolution of the Raman spectrum of carbonaceous material in low-grade metasediments of the Glarus Alps (Switzerland). <i>Terra Nova</i> 22 , 354-360.
596 597	Larouche N. and Stansfield B. L. (2010) Classifying nanostructured carbons using graphitic indices derived from Raman spectra. <i>Carbon</i> 48 , 620-629.
598 599 600	Lepland A., van Zuilen M. A., and Philippot, P. (2011) Fluid-deposited graphite and its geobiological implications in early Archean gneiss from Akilia, Greenland. <i>Geobiology</i> 9 , 2-9.
601 602 603 604	Lepot K., Williford K.H., Ushikubo T., Sugitani K., Mimura K., Spicuzza M.J., Valley J.W. (2013) Texture-specific isotopic compositions in 3.4 Gyr old organic matter support selective preservation in cell-like structures. <i>Geochimica et Cosmochimica Acta</i> 112, 66-86.
605 606 607	Lindsay J. F., Brasier M. D., McLoughlin N., Green O. R., Fogel M., Steele A., and Mertzman S. A. (2005) The problem of deep carbone-An Archean paradox. <i>Precambrian Research</i> 143, 1-22.
608 609	Marshall C. P., Emry J. R., and Olcott Marshall A. (2011) Haematite pseudomicrofossils present in the 3.5-billion-year-old Apex chert. <i>Nature Geoscience</i> 4 , 240-243.
610 611 612 613	Marshall C. P., Love G. D., Snape C. E., Hill A. C., Allwood A. C., Walter M. R., Van Kranendonk M. J., Bowden S. A., Sylva S. P., and Summons R. E. (2007) Structural characterization of kerogen in 3.4 Ga Archaean cherts from the Pilbara Craton, Western Australia. <i>Precambrian Research</i> 155, 1-23.
614 615	Marshall C. P., Mar G. L., Nicoll R. S., and Wilson M. A., (2001) Organic geochemistry of artificially matured conodonts. <i>Organic Geochemistry</i> 32 , 1055-1071.
616 617 618	Marshall C. P. and Olcott Marshall A. (2011) Hematite and carbonaceous materials in geological samples: A cautionary tale. <i>Spectrochimica Acta Part A: Molecular and Biomolecular Spectroscopy</i> 80 , 133-137.
619 620 621	Mostefaoui S. L., Perron C., Zinner E., and Sagon G. R. (2000) Metal-associated carbon in primitive chondrites: structure, isotopic composition, and origin. <i>Geochimica et Cosmochimica Acta</i> 64, 1945-1964.

622	Nasdala L., Brenker F. E., Glinnemann J., Hofmeister W., Gasparik T., Harris J. W., Stachel
623	T., and Reese I. (2003) Spectroscopic 2D-tomography: Residual pressure and strain
624	around mineral inclusions in diamonds. <i>European Journal of Mineralogy</i> 15, 931-935.
625	Nemanich R. J. and Solin S. A. (1979) First- and second-order Raman scattering from finite-
626	size crystals of graphite. <i>Physical Review B</i> 20 , 392-401.
627	Nijman W., de Bruijne K. (C.) H., Valkering M.E. (1998) Growth fault control of Early
628	Archaean cherts, barite mounds and chert-barite veins, North Pole Dome, Eastern
629	Pilbara, Western Australia. Precambrian Research 88 , 25-52.
630	Olcott Marshall A., Emry J. R., and Marshall C. P. (2012) Multiple generations of carbon in
631	the Apex chert and Implications for preservation of microfossils. <i>Astrobiology</i> 12,
632	160-166.
633	Olcott Marshall A., Jehlicka J., Rouzaud JN., and Marshall C. P. (2013) Multiple
634	generations of carbonaceous material deposited in Apex chert by basin-scale pervasive
635	fluid flow. <i>Gondwana Research</i> In press.
636	Olcott Marshall A. and Marshall C. P. (2013) Comment on "Biogenicity of Earth's earliest
637	fossils: A resolution of the controversy" by J. W. Schopf and A. B. Kudryavtsev,
638	Gondwana Research, Volume 22, Issue 3–4, Pages 761–771. Gondwana Research 23,
639	1654-1655.
640 641	Pasteris J. D. (1989) In situ analysis in geological thinsections by laser Raman microprobe spectroscopy: a cautionary note. <i>Applied Spectroscopy</i> 43 , 567-570.
642	Pinti D. L., Hashizume K., Sugihara A., Massault M., and Philippot P. (2009) Isotopic
643	fractionation of nitrogen and carbon in Paleoarchean cherts from Pilbara Craton,
644	Western Australia: Origin of 15N-depleted nitrogen. <i>Geochimica et Cosmochimica</i>
645	<i>Acta</i> 73, 3819-3848.
646	Pinti D. L., Mineau R., and Clement V. (2013) Comment on "Biogenicity of Earth's earliest
647	fossils: a resolution of the controversy" by J. William Schopf and Anatoliy B.
648	Kudryavtsev, Gondwana Research 22 (2012), 761–771. <i>Gondwana Research</i> 23,
649	1652-1653.
650	Qu Y., van Zuilen M. A., and Lepland A. (2011) Raman spectroscopic analysis of
651	heterogeneous carbonaceous matter in the 2.0 Ga Zaonega Fm, Karelia, Russia.
652	Goldschmidt Geochemistry Conference, Prague, Czech Republic.
653	Qu Y., Črne A. E., Lepland A., and van Zuilen M. A. (2012) Methanotrophy in a
654	Paleoproterozoic oil field ecosystem, Zaonega Formation, Karelia, Russia. <i>Geobiology</i>
655	10 , 467-478.
656	Qu Y., van Zuilen M. A., Lepland A, Črne A. E., Fliegel D., Somelar P. (2013) Structural
657	variation of metamorphosed carbonaceous material in the 2.0 Ga Zaonega Formation,
658	Karelia, Russia. <i>Precambrian Research</i> , In Review.
659	Quirico E., Raynal PI., and Bourot-Denise M. (2003) Metamorphic grade of organic matter
660	in six unequilibrated ordinary chondrites. <i>Meteoritics & Planetary Science</i> 38, 795-
661	811.

662	Rahl J. M., Anderson K. M., Brandon M. T. and Fassoulas C. (2005) Raman spectroscopic
663	carbonaceous material thermometry of low-grade metamorphic rocks: Calibration and
664	application to tectonic exhumation in Crette, Greece. <i>Earth and Planetary Science</i>
665	Letters 240, 339-354.
666 667	Ross J. V. and Bustin R. M. (1990) The role of strain energy in creep graphitization of anthracite. <i>Nature</i> 343 , 58-60.
668 669 670	Sadezsky A., Muckenhuber H., Grothe H., Niessner R., and Pöschl U. (2005) Raman microspectroscopy of soot and related carbonaceous materials: Spectral analysis and structural information. <i>Carbon</i> 43 , 1731-1742.
671 672	Schopf J. W. (1993) Microfossils of the Early Archean Apex chert: New evidence of the antiquity of life <i>Science</i> 260 , 640-646.
673	Schopf J. W. and Kudryavtsev A. B. (2012) Biogenicity of Earth's earliest fossils: A
674	resolution of the controversy. <i>Gondwana Research</i> 22, 761-771.
675	Schopf J. W. and Kudryavtsev A. B. (2013) Reply to the comments of D.L. Pinti, R. Mineau
676	and V. Clement, and of A.O. Marshall and C.P. Marshall on "Biogenicity of Earth's
677	earliest fossils: A resolution of the controversy" by J. William Schopf and Anatoliy B.
678	Kudryavtsev, Gondwana Research 22 (2012), 761-771. Gondwana Research 23, 1656-
679	1658.
680	Schopf J. W., Kudryavtsev A. B., Agresti D. G., Wdowiak T. J., and Czaja A. D. (2002).
681	Laser-Raman imagery of Earth's earliest fossils. <i>Nature</i> 416, 73-76.
682	Schopf J. W., Kudryavtsev A. K., Czaja A. C., and Tripathi A. B. (2007) Evidence of Archean
683	Life: Stromatolites and microfossils. <i>Precambrian Research</i> 158, 141-155.
684	Schopf J. W. and Packer B. M. (1987) Early Archean (3.3-Billion to 3.5-Billion-Year-Old)
685	Microfossils from Warrawoona Group, Australia. Science 237, 70-73.
686 687 688	Thébaud N. and Rey P. F. (2013) Archean gravity-driven tectonics on hot and flooded continents: Controls on long-lived mineralised hydrothermal systems away from continental margins. <i>Precambrian Research</i> 229 , 93-104.
689	Tice M., Bostick B. C., and Lowe D. R. (2004) Thermal history of the 3.5-3.2 Ga
690	Onverwacht and Fig Tree Groups, Barberton greenstone belt, South Africa, inferred
691	by Raman microspectroscopy of carbonaceous material. <i>Geology</i> 32 , 37-40.
692 693	Tuinstra F. and Koenig J. L. (1970) Raman spectrum of graphite. <i>Journal of Chemical Physics</i> 53 , 1126-1130.
694 695 696	Ueno Y., Yoshioka H., Maruyama S., and Isozaki Y. (2004) Carbon isotopes and petrography of kerogens in 3.5-Ga hydrothermal silica dikes in the North Pole area, Western Australia. <i>Geochimica et Cosmochimica Acta</i> 68 , 573-589.
697 698 699	van den Boorn S. H. J. M., van Bergen M. J., Nijman W., and Vroon P. Z. (2007) Dual role of seawater and hydrothermal fluids in Early Archean chert formation: evidence from silicon isotopes. <i>Geology</i> 35 , 939-942.

silicon isotopes. Geology 35, 939-942.

700 701	van Kranendonk M. J. (2000) Geology of the North Shaw 1:100 000 sheet: Western Australia Geological Survey, 1:100 000. Geological Series Explanatory Notes, 86 p.
702 703 704 705	van Kranendonk M. J. (2006) Volcanic degassing, hydrothermal circulation and the flourishing of early life on Earth: a review of the evidence from c. 3490-3240 Ma rocks of the Pilbara supergroup, Pilbara craton, western Australia. <i>Earth-Science Reviews</i> 74 , 197-240.
706 707 708	van Kranendonk M. J., Hickman A. H., Smithies R. H., and Nelson D. R. (2002) Geology and tectonic evolution of the Archean North Pilbara Terrain, Pilbara craton, western Australia. <i>Economic Geology</i> 97 , 695-732.
709 710 711 712	van Kranendonk M. J. and Pirajno F. (2004) Geochemistry of metabasalts and hydrothermal alteration zones associated with c. 3.45 Ga chert and barite deposits: implications for the geological setting of the Warrawoona Group, Pilbara craton, Australia. <i>Geochemistry: Exploration, Environment, Analysis</i> 4 , 253-278.
713 714 715	van Zuilen M.A., Fliegel D., Wirth R., Lepland A., Qu Y., Schreiber A., Romashkin A. E., and Philippot P. (2012) Mineral-templated growth of natural graphite films. <i>Geochimica et Cosmochimica Acta</i> 83, 252-262.
716 717 718	Wacey D., Saunders M., Brasier M.D., Kilburn, M.R. (2011) Earliest microbially mediated pyrite oxidation in ~3.4 billion-year-old sediments. <i>Earth and Planetary Science</i> <i>Letter</i> 301, 393-402.
719 720 721 722	Wang A., Dhamenincourt P., Dubessy J., Guerard D., Landais P., and Lelaurain M. (1989) Characterization of graphite alteration in an uranium deposit by micro-Raman spectroscopy, X-ray diffraction, transmission electron microscopy and scanning electron microscopy. <i>Carbon</i> 27, 209-218.
723 724 725	Waples D.W. (1994) Maturity modeling: thermal indicators, hydrocarbon generation, and oil cracking. In <i>The petroleum system – from source to trap (eds. L.B. Magoon and W.G. Dow)</i> , AAPG Memoir 60 , 285-306
726 727	Wopenka B. and Pasteris J. D. (1991) Raman spectra of graphite as indicators of degree of metamorphism. <i>Canadian Mineralogist</i> 29 , 1-9.
728 729 730	Wopenka B. and Pasteris J. D. (1993) Structural characterization of kerogens to granulite- facies graphite: Applicability of Raman microprobe spectroscopy. American Mineralogist 78, 533-557.
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732 Figure captions

733

Figure 1: Simplified geological maps of a) the North Pilbara Craton (modified from Thébaud

and Rey, 2013) and of b) the Apex chert (modified from Brasier et al., 2005). The numbers

correspond to the location of the samples analyzed in this study (see Fig. 3).

737

Figure 2: Decomposition of the Raman spectrum of disordered CM. a) Disorder-associated 738 bands D2, D3 and D4 are distinguished as well as the main D1 and G bands (see Sadezky et 739 740 al. (2005) for band labeling and fitting intervals). Each of the five fitted bands has a Gaussian-Lorentzian shape where the percentage of Gaussian can vary freely. The band position are 741 fixed between 1340-1360 cm⁻¹ for D1, 1615-1625cm⁻¹ for D2, 1490-1510 cm⁻¹ for D3, 1150-742 1250cm⁻¹ for D4 and 1580-1610cm⁻¹ for G. The intensity of D2 was limited to less than half 743 744 of that of the G-band. For the D4-band, the maximum band width (FWHM) was limited at a maximum of 200 cm⁻¹. This constraint was necessary to prevent the generation of 745 unreasonably large D4-bands in some of the spectra. b) Following protocol 2, only the main 746 bands of D and G were distinguished. The band positions are fixed between 1340-1360 cm⁻¹ 747 for D and 1580-1610 cm^{-1} for G. 748

749

Figure 3: Thin sections of all samples analyzed in this study. White dots represent Raman
spectral analyses of carbonaceous matter in stratiform chert. Black dots represent
carbonaceous matter in intrusive black cherts zones. Squares correspond to locality where
Raman maps were acquired. Sample #1 maps are located in the same area, while the maps #61 and #6-2 are located in the dark gray chert and in the white chert, respectively.

755

Figure 4: Representative spectra for the 6 samples obtained by punctual analyses showingfirst and second order bands.

758

Figure 5: Example of apparent ambiguity in the use of the ratios R2 and RA1 for punctual 759 760 analysis in samples #1, #4 and #5. a) I_D/I_G vs R_2 ratios. R_2 and I_D/I_G ratios are lower for the stratiform chert CMs (white dots and diamonds) than for the black chert veins CMs (black 761 762 dots) indicating that the CM order is higher for the stratiform cherts than for the feeder black chert veins. b) I_D/I_G vs R_{A1} ratios. R_{A1} and I_D/I_G are lower for the stratiform cherts than for the 763 feeder black chert veins indicating that the CM order is higher for the hydrothermal cherts 764 than for the stratiform cherts. Filled and unfilled grey points correspond to the mean of the 765 analyses for sample #1 and samples #4-5, respectively. Error bars correspond to 1 standard 766 deviation. 767

768

Figure 6: Two pools of CM in the Apex chert. a) Mean spectrum for stratiform chert CM 769 770 normalized to D1. I_{D3} and I_{D4} are relatively intense corresponding to a poorly organized CM, 771 which argues for a low peak. b) Mean spectrum for black chert veins normalized to D1. I_{D3} and I_{D4} are relatively low, which correspond to a better organized CM and thus to a higher 772 peak temperature. c) R_{D4} vs R_{D3} for samples #1, #4 and #5. Two well separated groups of CM 773 774 can be recognized; black chert vein CM with low R_{D3} and R_{D4} and stratiform chert CM with 775 high R_{D3} and R_{D4}. Filled and unfilled grey points correspond to the mean of the analyses for sample #1 and samples #4-5, respectively. Error bars correspond to 1 standard deviation. d) 776 777 R_{D4} vs R_{D3} for all samples. Samples #2 and #6 have a 'stratiform chert signature' which is contrary to what is expected. Sample #3 shows a low-ordered CM. 778

779

Figure 7: Distribution of the CM in the stratiform chert and the black chert veins showed by a) histograms of R_{D3} ratios for samples #1 (white), #2 (light gray), #4 and #5 (black). b) Histogram of R_{D3} ratios for sample #6; light gray histogram is for the intrusive white chert and the dark gray histogram is for the darker chert. The white chert CMs fall within the range of black chert veins CMs whereas the darker chert CMs fall within the range of the stratiform chert CMs.

786

Figure 8 Poorly ordered CM in the Apex chert, showed in a) R_{D3} vs R_{D4} diagram. All samples 787 align themselves on a slope with a coefficient of ~ 1 except for measurements of sample #3 788 suggesting post-emplacement phenomena b) FWHM-D/FWHM-G vs I_D/I_G. The data for 789 790 sample #3 have high FWHM-D/FWHM-G and low I_D/I_G relative to the feeder veins and 791 stratiform cherts, which indicate poorly organized CM with a metamorphic temperature lower 792 than regional metamorphism and hydrothermal alteration. The range where the spectrum of 793 Apex chert showed in Schopf et al. (2007) is represented with crosshatches. It coincides with 794 the post-emplacement altered CM range. The mean ID/IG ratio calculated by Olcott Marshall 795 et al. (2012) for the two identified groups is shown with dashed lines. Primary phase (E1) has feeder vein chert signature (hydrothermal origin) and their secondary phase has a stratiform 796 797 chert signature (regional metamorphism) linked with high hematite content. Their secondary 798 phase could arise from the oxidation of the hydrothermally-derived chert during weathering.

799

Figure S1: Representative spectra for the 6 samples obtained with hyperspectral mappinganalysis.

802

Table 1: Comparison between ratios calculated for embedded CM and CM at the samplesurface.

805

Table 2: Mean values calculated for the different ratio from the punctual analysis and the map analysis. I_D/I_G , A_D/A_G are calculated with the protocol 2. R_2 , R_{A1} , R_{A2} , R_{D3} and R_{D4} , T_{R2} , T_{RA1} and T_{RA2} are calculated with protocol 1. The tentative temperatures that must be considered are in bold case. Positions of D (ω -D1) and G (ω -G), FWHM-D and FWHM-G and the ratio FWHM-D/FWHM-G are calculated from protocol 1 (*) and from protocol 2 (**). Errors correspond to 2 standard deviations.

812

813 **Table ST1:** Characteristics of the hyperspectral maps for each sample

814

Table ST2: Values calculated for the different ratio from the punctual analysis for each punctual analysis. I_D/I_G , A_D/A_G are calculated with the protocol 2. R_2 , R_{A1} , R_{A2} , R_{D3} and R_{D4} , T_{R2} , T_{RA1} and T_{RA2} are calculated with protocol 1. The tentative temperatures that must be considered are in bold case. Positions of D (ω -D1) and G (ω -G), FWHM-D and FWHM-G and the ratio FWHM-D/FWHM-G are calculated from protocol 1 (*) and from protocol 2 (**). Errors correspond to 2 standard deviations.

821

822 Table ST3: Values calculated for the different ratio from the map analysis for each individual

823 spectrum. I_D/I_G, A_D/A_G are calculated with the protocol 2. R₂, R_{A1}, R_{A2}, R_{D3} and R_{D4}, T_{R2},

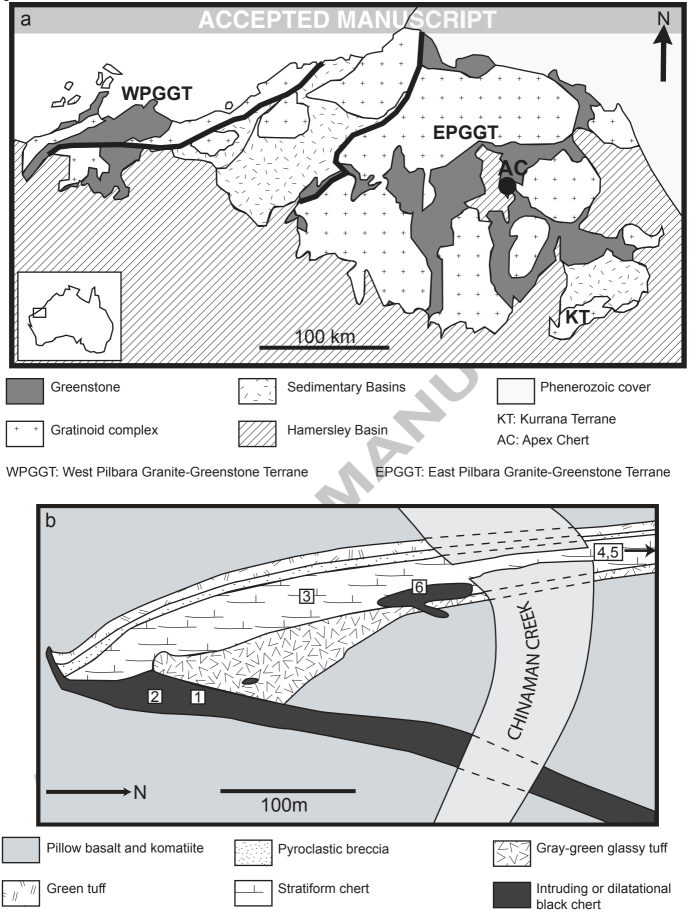
 T_{RA1} and T_{RA2} are calculated with protocol 1. The tentative temperatures that must be 824

considered are in bold case. Positions of D (ω -D1) and G (ω -G), FWHM-D and FWHM-G 825

Jm J and the ratio FWHM-D/FWHM-G are calculated from protocol 1 (*) and from protocol 2 826

827 (**). Errors correspond to 2 standard deviations.

Figure 1



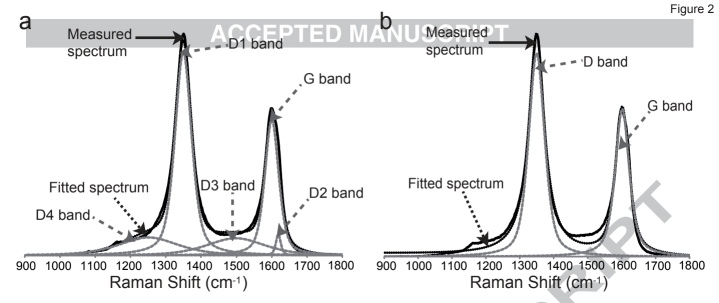
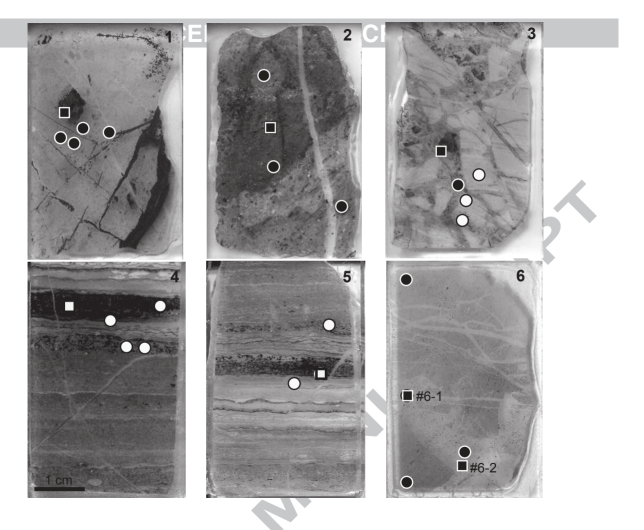
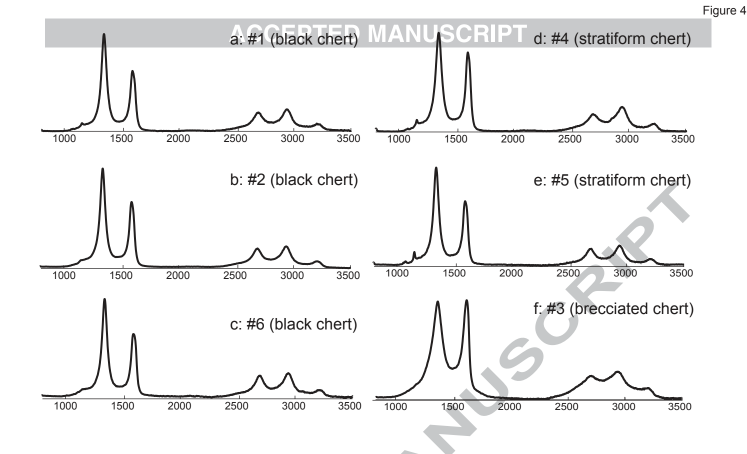
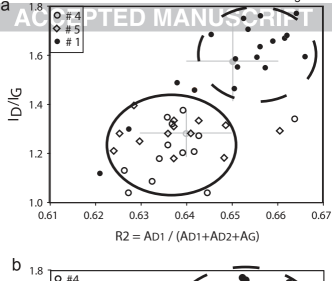


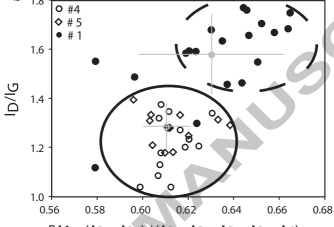
Figure 3



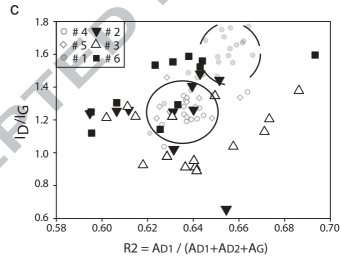








RA1 = (AD1+AD4) / (AD1+AD2+AD3+AD4+AG)



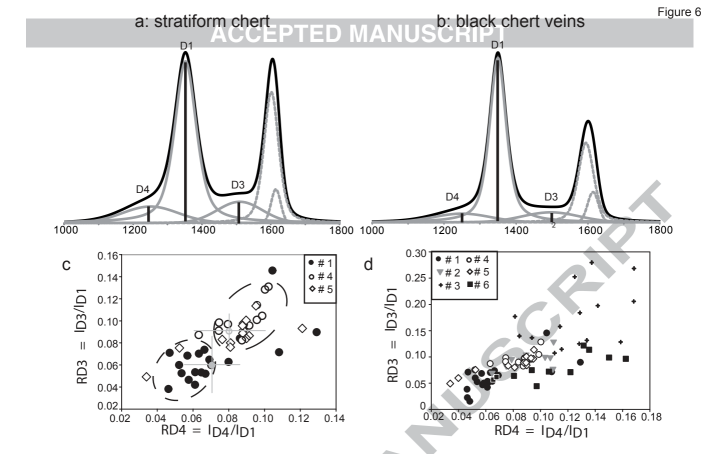


Figure 7

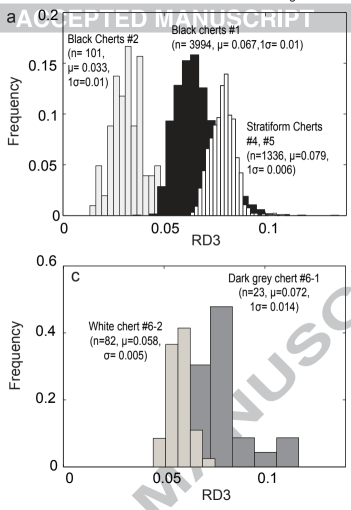
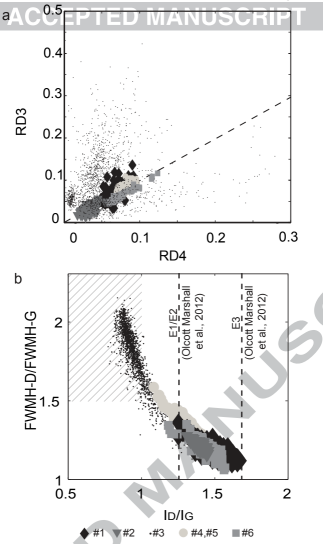


Figure 8



1 1	Туре	D1-w	R2	RA1	RA2	ID/IG	AD/AG	RD3	RD4	Туре	D1-w	R2	RA1	RA2	ID/IG	AD/AG	RD3	RD4
1	Embedded	53.54	0.66	0.65	1.82	1.76	1.92	0.05	0.06	Surface	53.67	0.66	0.64	1.80	1.75	1.92	0.05	0.07
	Embedded	54.25	0.66	0.66	1.92	1.67	1.90	0.05	0.07	Surface	60.60	0.63	0.61	1.57	1.45	1.78	0.08	0.09
1	Embedded	56.94	0.66	0.65	1.86	1.55	1.91	0.04	0.05	Surface	56.96	0.65	0.64	1.79	1.54	1.88	0.06	0.07
1	Embedded	53.14	0.65	0.65	1.88	1.71	1.90	0.08	0.09	Surface	54.80	0.64	0.63	1.68	1.62	1.84	0.06	0.08
1	Embedded	53.02	0.65	0.66	1.98	1.68	1.85	0.05	0.06	Surface	53.10	0.65	0.64	1.80	1.72	1.88	0.04	0.06
1	Embedded	52.32	0.65	0.66	1.98	1.75	1.90	0.06	0.07	Surface	56.83	0.65	0.64	1.80	1.55	1.84	0.06	0.07
1	Embedded	54.13	0.66	0.65	1.83	1.66	1.90	0.04	0.06	Surface	53.58	0.66	0.65	1.84	1.71	1.87	0.04	0.05
1	Embedded	57.68	0.64	0.64	1.76	1.46	1.78	0.06	0.08	Surface	60.51	0.63	0.63	1.68	1.39	1.73	0.06	0.07
1	Mean	54.38	0.65	0.65	1.88	1.65	1.88	0.05	0.07	Mean	56.26	0.65	0.64	1.75	1.59	1.84	0.06	0.07
1	2σ	3.84	0.01	0.02	0.16	0.21	0.09	0.03	0.03	2σ	6.04	0.02	0.03	0.18	0.27	0.12	0.03	0.02
4	Embedded	61.35	0.64	0.61	1.57	1.35	1.84	0.08	0.09	Surface	63.73	0.61	0.56	1.29	1.37	1.62	0.13	0.10
4	Embedded	68.01	0.64	0.62	1.62	1.20	1.80	0.13	0.10	Surface	65.20	0.64	0.62	1.60	1.32	1.83	0.10	0.09
4	Embedded	67.28	0.64	0.62	1.66	1.21	1.79	0.10	0.10	Surface	68.16	0.64	0.62	1.64	1.19	1.80	0.10	0.10
4	Embedded	63.88	0.64	0.61	1.55	1.32	1.82	0.10	0.09	Surface	61.51	0.64	0.62	1.63	1.34	1.82	0.09	0.09
4	Embedded	66.33	0.64	0.62	1.63	1.23	1.77	0.10	0.09	Surface	68.19	0.64	0.62	1.60	1.20	1.79	0.10	0.10
4	Embedded	66.13	0.64	0.62	1.61	1.27	1.81	0.10	0.10	Surface	68.67	0.64	0.62	1.62	1.19	1.82	0.11	0.10
4	Mean	65.50	0.64	0.62	1.61	1.26	1.81	0.09	0.08	Mean	65.91	0.64	0.61	1.56	1.27	1.78	0.09	0.08
4	2σ	4.93	0.01	0.01	0.09	0.12	0.05	0.06	0.05	2σ	5.83	0.03	0.04	0.27	0.16	0.16	0.07	0.06
5	Embedded	64.35	0.63	0.62	1.63	1.25	1.73	0.08	0.09	Surface	66.85	0.61	0.60	1.49	1.20	1.77	0.09	0.12
5	Embedded	66.13	0.64	0.62	1.61	1.33	1.68	0.09	0.12	Surface	58.67	0.64	0.62	1.60	1.37	1.81	0.08	0.08
5	Embedded	63.61	0.62	0.60	1.53	1.21	1.72	0.10	0.09	Surface	61.78	0.64	0.62	1.63	1.30	1.80	0.09	0.09
5	Embedded	66.66	0.64	0.61	1.57	1.28	1.86	0.08	0.08	Surface	66.57	0.64	0.62	1.63	1.27	1.80	0.10	0.09
5	Embedded	60.69	0.64	0.61	1.57	1.28	1.69	0.09	0.09	Surface	64.67	0.63	0.62	1.63	1.19	1.78	0.09	0.10
5	Embedded	63.33	0.63	0.61	1.58	1.28	1.69	0.09	0.09	Surface	64.71	0.64	0.62	1.61	1.28	1.80	0.10	0.09
5	Embedded	63.09	0.64	0.60	1.52	1.33	1.81	0.10	0.09	Surface	62.35	0.64	0.61	1.57	1.34	1.80	0.09	0.09
5	Embedded	64.69	0.64	0.60	1.52	1.31	1.81	0.10	0.09	Surface	62.59	0.64	0.62	1.62	1.24	1.80	0.09	0.09
5	Mean	64.07	0.63	0.61	1.57	1.29	1.75	0.09	0.09	Mean	63.52	0.64	0.61	1.60	1.27	1.79	0.09	0.09
5	2σ	3.74	0.01	0.01	0.08	0.08	0.13	0.02	0.02	2σ	5.45	0.02	0.01	0.10	0.13	0.03	0.01	0.02

Table. 1 Acceptin

Sample	Туре	n	ω-D1*	ω-G*	ω-D1**	ω-G**	FWHM-D1*	FWHM-G*	FWHM-D1/ FWHM-G**	FWHM-D1**	FWHM-G**	FWHM-D1/ FWHM-G**
1	Punctual	19	1350.0 ± 1.3	1597.2 ± 2.0	1349.6 ± 1.5	1600.0 ± 0.9	56.80 ± 10.56	48.58 ± 4.17	1.05 ± 0.36	57.88 ± 12.70	50.88 ± 5.12	1.14 ± 0.20
	Мар	3994	1349.9±0.7	1596.5±1.2	1349.6 ± 0.7	1600.2 ± 0.5	57.3 ± 3.61	48.12 ± 2.04	1.19 ± 0.05	<i>59.75</i> ± 5.5	50.44 ± 2.29	1.18 ± 0.06
2	Punctual	11	1349.4 ± 2.4	1599.1 ± 2.9	1349.1 ± 3.4	1602.4 ± 1.8	62.07 ±13.0	48.38 ± 4.1	1.28 ± 0.35	61.54 ± 15.90	50.02 ± 5.13	1.23 ± 0.4
	Мар	101	1348.7±0.5	1600.4±1.3	1348.5 ± 0.5	1602.3 ± 0.4	56.84 ± 2.03	46.24 ± 1.72	1.19 ± 0.05	<i>55.41</i> ± 1.95	<i>45.72</i> ± 1.38	1.21 ± 0.05
	Punctual	15	1348.3 ± 3.0	1597.4 ± 4.5	1347.2 ± 3.4	1600.9 ± 2.4	61.56 ± 18.18	47.48 ± 2.48	1.09 ± 0.69	67.33 ± 24.78	52.65 ± 7.38	1.27 ± 0.31
6	Map 6-1	23	1347.9 ± 2.2	1595.5 ± 2.1	1347.5± 2.5	1599.3 ± 2.4	57.50±7.17	47.70 ± 2.19	1.21 ± 0.15	60.00 ± 9.64	50.77 ± 2.67	1.18±0.17
	Map 6-2	82	1349.5 ± 0.4	1598.1 ± 0.9	1349.1 ± 0.5	1602.0 ± 0.5	56.81±3.13	47.73 ± 1.11	1.19 ± 0.06	58.67± 4.22	48.26 ± 1.10	1.22 ± 0.08
4	Punctual	13	1349.7 ± 2.5	1599.4 ± 2.1	1349.1 ± 2.8	1602.6 ± 2.1	66.17 ±6.87	48.72 ± 3.54	1.03± 0.65	69.26 ± 8.83	49.64 ± 1.96	1.40 ± 0.18
	Мар	1225	1350.7±0.7	1598.9 ± 1.2	1350.3 ± 0.8	1603.2± 0.7	<i>61.18</i> ± 4.3	49.15 ± 1.97	1.24 ± 0.09	<i>64.08</i> ± 4.78	<i>49.29</i> ± 1.58	1.30 ± 0.09
5	Punctual	14	1348.9 ± 1.3	1599.1 ± 2.4	1348.5 ± 1.7	1602.1 ±1.4	63.78 ±5 .78	47.74 ± 2.27	0.98 ± 0.60	64.53 ± 8.73	49.14 ± 2.08	1.31 ± 0.17
	Мар	111	1349.6 ± 0.4	1599.0±0.7	1349.0±0.5	1602.8 ± 0.5	<i>63.26</i> ± 1.90	47.05 ± 1.54	1.34 ± 0.06	66.67±2.39	47.94 ± 1.31	1.39 ± 0.06
3	Punctual	15	1350.6 ± 3.9	1599.8± 3.5	1349.8 ±5.5	1601.9 ±2.4	91.82 ±40.68	55.68±8.89	1.40 ± 1.0	106.56± 53.02	60.56 ± 5.63	1.75 ± 0.78
Ũ	Мар	1758	1350.2± 3.7	1598.1 ± 4.5	1349.0± 2.4	1603.0 ± 2.9	94.95 ± 22.88	57.40 ± 12.34	1.72± 5.03	96.02±22.19	53.38± 7.38	1.80 ± 0.37
							~					

Sample	Туре	n	ID/IG	AD/AG	R2	RA1	RA2	RD3	RD4	TR2	TRA1	TRA2
1	Punctual	19	1.58 ± 0.33	1.89 ± 0.13	0.65 ± 0.02	0.63 ± 0.05	1.73 ± 0.36	0.06 ± 0.05	0.07 ± 0.04	351.3 ± 10.6	320.6 ± 63.2	324.7±79.5
	Мар	3994	1.51 ± 0.13	1.81 ± 0.08	0.64 ± 0.01	0.62 ± 0.03	1.61 ± 0.18	0.07 ± 0.02	0.07 ± 0.01	349.3 ± 5.8	300.5 ± 33.2	297.5 ± 39.6
2	Punctual	11	1.23 ± 0.45	1.66 ± 0.83	0.63 ± 0.04	0.60 ± 0.06	1.51 ± 0.35	0.09 ± 0.03	0.09 ± 0.03	360.5 ± 16.8	279.7 ± 75.0	275.4 ± 78.5
	Мар	101	1.4 ± 0.06	1.71 ± 0.05	0.63 ± 0.02	0.63± 0.02	1.67 ± 0.14	0.03 ± 0.02	0.03 ± 0.01	359.0 ± 3.5	312.2 ± 24.8	311.5 ± 30.1
	Punctual	15	1.41 ± 0.37	1.91 ± 0.62	0.63 ± 0.05	0.63 ± 0.07	1.70 ± 0.52	0.08 ± 0.05	0.10 ± 0.07	360.7 ± 24.3	313.4 ± 84.2	317.5 ± 115.0
6	Map 6-1	23	1.40 ± 0.25	1.66± 0.08	0.62 ± 0.02	0.59 ± 0.02	1.43 ± 0.11	0.08 ± 0.03	0.09 ± 0.03	365.3 ± 7.3	266.0 ± 24.27	258.3 ± 25.0
	Map 6-2	82	1.45 ± 0.11	1.78± 0.06	0.63 ± 0.01	0.62 ± 0.01	1.64 ± 0.07	0.06 ± 0.01	0.07 ± 0.01	358.9± 3.02	306.7 ± 12.3	304.4 ± 15.2
4	Punctual	13	1.25 ± 0.20	1.82 ± 0.12	0.64 ± 0.02	0.61 ± 0.02	1.59 ± 0.12	0.10 ± 0.03	0.08 ± 0.03	356.6 ± 8.4	297.6 ± 21.7	293.6 ± 26.1
	Мар	1225	1.35 ± 0.12	1.77± 0.05	0.63 ± 0.01	0.61 ± 0.01	1.56 ± 0.07	0.08 ± 0.01	0.08 ± 0.01	352.7 ± 2.8	292.3 ± 13.6	287.1 ±15.9
5	Punctual	14	1.31 ± 0.17	1.83 ± 0.23	0.64 ± 0.02	0.61 ± 0.02	1.56 ± 0.09	0.09 ± 0.03	0.08 ± 0.04	356.8 ± 9.0	296.4 ± 28.6	292.3 ± 35.0

	Мар	111	1.29 ± 0.06	1.81 ± 0.03	0.64 ± 0.004	0.62 ± 0.01	1.62 ± 0.04	0.08 ± 0.01	0.08 ± 0.01	350.3 ± 2.0	302.4 ± 8.1	299.2 ± 9.8
3	Punctual	15	1.11±0.34	2.10 ± 0.79	0.64 ± 0.05	0.58 ± 0.11	1.46 ± 0.61	0.17± 0.11	0.12 ± 0.05	356.1 ± 21.6	263.8 ± 133.3	263.7 ± 135.6
-	Мар	1758	0.95 ± 0.16	1.77± 0.38	0.62± 0.03	0.60 ± 0.05	1.51 ± 0.28	0.10 ± 0.14	0.05 ± 0.09	357.3 ± 12.8	280.1 ± 57.2	275.5 ± 61.2
												Table 2
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