1 A novel, dichloromethane-fermenting bacterium in the *Peptococcaceae* family,

- 2 'Candidatus Formamonas warabiya', gen. nov. sp. nov.
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9 **Running Title**

10 A novel, dichloromethane fermenting bacterium

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A novel, dichloromethane fermenting bacterium

Holland *et al* (2020)

20 Abstract

21 Dichloromethane (DCM; CH₂Cl₂) is a toxic groundwater pollutant that also has a detrimental effect 22 on atmospheric ozone levels. As a dense non-aqueous phase liquid, DCM migrates vertically 23 through groundwater to low redox zones, yet information on anaerobic microbial DCM 24 transformation remains scarce due to a lack of cultured organisms. We report here the 25 characterisation of strain DCMF, the dominant organism in an anaerobic enrichment culture 26 (DFE) that is capable of fermenting DCM to the environmentally benign product acetate. Stable 27 carbon isotope experiments demonstrated that the organism assimilated carbon from DCM and bicarbonate via the Wood-Ljungdahl pathway. Strain DCMF is the first anaerobic DCM-degrading 28 29 bacterium also shown to metabolise non-chlorinated substrates. It appears to be a methylotroph 30 utilising the Wood-Ljungdahl pathway for metabolism of methyl groups from methanol, choline, and glycine betaine, which has implications for the flux of climate-active compounds from 31 32 subsurface environments. Community profiling and enrichment of the cohabiting taxa in culture 33 DFE to the exclusion of strain DCMF suggest that it is the sole organism in this culture responsible 34 for substrate metabolism, while the cohabitants persist via necromass recycling. Genomic and 35 physiological evidence support placement of strain DCMF in a novel genus, 'Candidatus Formamonas warabiya'. 36

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Abbreviations: DCM, dichloromethane; FISH, fluorescence *in situ* hybridisation; GC-FID, gas
chromatography with flame ionisation detector; GC-PDD, gas chromatography with pulse
discharge detector; GC-TQMS, gas chromatography with triple quadrupole mass spectrometry;
LC-MS/MS, liquid chromatography with tandem mass spectrometry; MRM, multiple reaction
monitoring; WLP, Wood-Ljungdahl pathway.

A novel, dichloromethane fermenting bacterium

Holland *et al* (2020)

43 Introduction

44 Dichloromethane (DCM, CH₂Cl₂) is one of the most commonly encountered subsurface pollutants in industrial areas (1). Current global production of DCM exceeds 900 Gg y⁻¹, of which 70% is 45 46 manufactured by humans (2). The remaining 30% comes from natural sources including biomass 47 burning, oceanic sources, and geothermal activity (2). Due to widespread production and use of 48 DCM, both surface and tropospheric levels of this toxic chemical continue to rise (3-6). 49 Atmospherically, DCM has recently been recognised as a potent greenhouse gas with detrimental 50 effects on ozone (7). The compound also poses a threat to human health (8,9) and microbial function (10–12). 51

52 Microbial transformation of DCM is an option for remediation of oxic and anoxic environments. 53 Aerobic DCM transformation is catalysed by a DCM dehalogenase found in facultative methylotrophs (13,14), but the enzyme responsible for anaerobic DCM dechlorination has not yet 54 been identified. DCM is a dense non-aqueous phase liquid that descends through groundwater to 55 56 low redox zones, and so anaerobic degradation plays a vital role in its removal from contaminated 57 sites. Yet study of anaerobic DCM degradation has been hindered by the highly limited number of 58 organisms capable of this metabolism. *Dehalobacterium formicoaceticum* strain DMC is the only 59 isolate (15) and from the handful of enrichment cultures (16-18), only 'Candidatus Dichloromethanomonas elyunquensis' strain RM has been characterised (19,20). Both D. 60 61 formicoaceticum and 'Ca. Dichloromethanomonas elyunguensis' are obligate anaerobic DCMdegrading bacteria and have genome sequences available (15,19,21,22). A combination of 62 genomic, physiological and proteomic work has demonstrated the central role of the Wood-63 Ljungdahl pathway (WLP) in DCM metabolism in both organisms, however variations on the 64 pathway result in different end products (15,20,23). D. formicoaceticum ferments DCM to formate 65 66 and acetate in a 2:1 molar ratio (15), whilst 'Ca. Dichloromethanomonas elyunquensis' completely mineralises DCM to H_2 and CO_2 (23). 67

A novel, dichloromethane fermenting bacterium

Holland *et al* (2020)

68 A new anaerobic DCM degrading bacterium, strain DCMF, was recently subjected to whole genome sequencing and also encoded a complete WLP (24). Strain DCMF is the dominant 69 70 organism in a non-methanogenic bacterial consortium, designated culture DFE (24). The 71 community was enriched from an organochlorine-contaminated aquifer near Botany Bay, 72 Sydney, Australia and culture DFE has been maintained on DCM as the sole energy source (24). 73 However, genome-based metabolic modelling suggested that strain DCMF may have a wider 74 substrate repertoire due to the presence of 81 full-length MttB superfamily methyltransferases and glycine/betaine/sarcosine reductase genes (24). 75

76 Here, we report that strain DCMF is the first non-obligate anaerobic DCM degrading bacterium 77 and characterise its metabolism of DCM, guaternary amines, and methanol, whilst also considering the role of the cohabiting bacteria in culture DFE. Stable carbon isotope labelling was 78 79 used to determine the fate of DCM carbon and function of the WLP. Based on its genomic and 80 physiological novelty, strain DCMF is proposed to form a novel genus within the Peptococcaceae family. Using contemporary molecular and traditional cultivation techniques (including exclusion 81 82 cultivation – removal the dominant taxon), this study represents a thorough and robust 83 characterisation of a novel bacterium despite its presence in a multi-lineage enrichment culture. 84 This supports recent initiatives to redefine how bacterial lineages are formally recognised (25).

85 Materials and Methods

86 *Culture medium*

Culture DFE was grown in anaerobic, defined bicarbonate-buffered mineral salts medium as
previously described (24). To investigate the requirement for exogenous bicarbonate during DCM
degradation, cultures were instead buffered with 3-morpholinopropane-1-sulfonic acid (MOPS,
4.2 g l⁻¹), either with or without 4 mM NaHCO₃. To study the metabolic fate of DCM, ¹³C-labelled
DCM ([¹³C]DCM, 1 mM) was used. To study the assimilation of inorganic carbon, ¹³C-labelled
bicarbonate (NaH¹³CO₃, 5 mM) was added to MOPS-buffered culture medium.

A novel, dichloromethane fermenting bacterium

Holland *et al* (2020)

93 To test alternative growth substrates, DCM was replaced with the following (5 mM unless stated 94 otherwise): carbon monoxide (2 mM), choline chloride, dibromomethane, dimethylglycine, 95 formic acid, H₂, glycine betaine, methanol, sarcosine, syringic acid, trimethylamine. Cultures 96 amended with choline chloride, glycine betaine, and trimethylamine were also amended with the 97 following compounds as electron acceptors (15 mM unless otherwise stated): fumarate (80 mM, 98 tested with trimethylamine only), NaNO₂, NaNO₃, Na₂SO₃ and Na₂SO₄. Acetate, H₂, and lactate 99 were tested as electron donors with Na_2SO_3 and Na_2SO_4 as electron acceptors. Glycine betaine and 100 sarcosine (5 mM) were tested as electron donors with H_2 (10 mM) as electron acceptor.

101 Analytical methods

DCM, dibromomethane, acetate, formate, methanol, and trimethylamine were quantified using a
Shimadzu Plus GC-2010 gas chromatograph with flame ionisation detector (GC-FID) equipped
with a headspace autosampler (PAL LHS2-xt-Shim; Shimadzu, Rydalmere, Australia; Table S1).
HCO₃- (as gaseous CO₂) and H₂ were quantified using a Shimadzu GC-2010 gas chromatograph
with pulsed discharge detector (GC-PDD; Table S1). In all analyses, the inlet temperature was
250°C, split ratio 1:10, FID temperature 250°C or PDD temperature 150°C.

108 Choline and glycine betaine were quantified using liquid chromatography with tandem mass 109 spectrometry (LC-MS/MS). The Agilent 1200 Series LC (Agilent Technologies, Mulgrave, Australia) was fitted with a Luna C18(2) column (150 \times 4.6 mm, 5 μ m; Phenomenex, Lane Cove 110 West, Australia). The mobile phases were 0.5 mM ammonium acetate in water (A) and 100% 111 methanol (B). Samples (5 µl) were eluted with a linear gradient from 95:5 (A:B) to 0:100 (A:B) 112 113 over 10 min, then held at 0:100 (A:B) for 1 min. The LC was coupled to an Applied Biosystems 114 QTRAP 4000 quadrupole mass spectrometer (SCIEX, Mulgrave, Australia) and electrospray 115 ionization performed in the positive mode. The machine was operated in multiple reaction monitoring (MRM) mode and the following precursor-product ion transitions were used for 116 quantification: $m/z \ 104.0 \rightarrow 59.0$ (choline) and $m/z \ 118.0 \rightarrow 57.7$ (glycine betaine). 117

A novel, dichloromethane fermenting bacterium

Holland *et al* (2020)

Labelled and unlabelled acetate, CO_2 , and HCO_3^- were quantified via GC with triple quadrupole mass spectrometry (GC-TQMS) performed with an Agilent 7890A GC system (Table S1). The TQMS was operated in MRM mode identifying the following precursor-product ion transitions: $m/z 43 \rightarrow 15.2$ (unlabelled acetate), $m/z 44 \rightarrow 15.1$ ([1-¹³C]acetate), $m/z 44 \rightarrow 16$ ([2-¹³C]acetate), $m/z 45 \rightarrow 16.1$ ([1,2-¹³C]acetate), $m/z 45 \rightarrow 29$ (¹³CO₂), $m/z 44 \rightarrow 28$ (¹²CO₂).

123 GC-TQMS in MRM mode was also used to quantify dimethylamine, methylamine, sarcosine, and 124 glycine, using alanine as an internal standard. Following derivatisation (26) (Table S1), the 125 following precursor-product ion transitions were used: m/z 117.2 \rightarrow 89.1 (dimethylamine), m/z126 $103.2 \rightarrow 74.9$ (methylamine), m/z 116.2 \rightarrow 44.1 (sarcosine and alanine), and $102 \rightarrow 30.1$ 127 (glycine).

128 Bacterial quantification

129 Genomic DNA was extracted from 2 ml liquid culture as previously described (24). Strain DCMF 130 and total bacterial 16S rRNA genes were quantified via quantitative real-time PCR (qPCR) with 131 primers Dcm775/Dcm930 and Eub1048/Eub1194 (27), respectively (Table S2). Standard curves were prepared by making serial 10-fold dilutions of plasmid DNA carrying cloned strain DCMF 132 16S rDNA or *Dehalococcoides* sp. 16S rDNA (for total bacterial quantification). Reactions were 133 134 carried out on a CFX96 thermal cycler (Bio-Rad) and the data was analysed with CFX Maestro v1.0 software (Bio-Rad). Strain DCMF 16S rRNA gene copy numbers were converted to cell 135 136 numbers by dividing by four (the number of 16S rRNA genes in the genome).

137 16S rRNA gene amplicon sequencing

Community profiling was carried out on the above DNA samples. The 16S rRNA gene was amplified with the 515F/806R primer pair with adapters (Table S2). Samples were sequenced with Illumina MiSeq technology by The Hawkesbury Institute for the Environment Next Generation Sequencing Facility. Amplicon reads were processed in QIIME2 (28) using the dada2 pipeline (29): forward and reverse reads were trimmed and joined, chimeras were removed, and

A novel, dichloromethane fermenting bacterium

Holland *et al* (2020)

143 samples were rarefied to the lowest sequencing depth. Taxonomy was assigned to genus level 144 using a Naïve Bayes classifier trained on a full-length 16S rRNA gene SILVA database (release 145 133) and the lowest 1% abundant reads were filtered out. Alpha diversity was assessed with 146 Shannon's diversity index and pairwise comparisons made with a Kruskal-Wallis test. A two-147 dimensional PCA plot was created from the weighted Unifrac distance matrix. Samples were 148 compared by the proportion of substrate consumed, as well as timepoint, to account for differing 149 metabolic rates between substrates (Table S3).

150 Exclusion cultivation of DFE cohabitant bacteria

151 To eliminate strain DCMF and enrich the cohabiting bacteria in culture DFE, two rounds of 152 dilution to extinction cultures (20 ml) were set up in 30 ml glass serum bottles (Fig S1). These 153 were prepared with the standard medium amended with one of: casamino acids (5 g l-1), ethanol 154 (10 mM), glucose (10 mM), peptone (5 g l^{-1}), 1-propanol (10 mM), yeast extract (5 g l^{-1}). Following 155 qPCR confirmation that strain DCMF was below the limit of detection in the lowest active dilution 156 culture, these cultures were subject to Illumina 16 rRNA gene amplicon sequencing and used to 157 inoculate triplicate microcosms amended with one of: 1 mM DCM, 5 mM choline chloride, or 5 158 mM glycine betaine (Fig S1), which were monitored for eight weeks.

159 Fluorescence in situ hybridisation microscopy

160 Fluorescence in situ hybridisation (FISH) was carried out with a strain DCMF-specific 161 oligonucleotide probe (Dcm623, 5'-/Cy3/CTCAAGTGCCATCTCCGA-3'), designed using ARB (30), 162 and probe Eub338i (5'-/6-FAM/GCTGCCTCCCGTAGGAGT-3') (31) to target all bacteria. FISH was 163 carried out as per an established protocol for fixation on a polycarbonate membrane, using 164 minimal volumes of reagents (32). Cells were fixed with protocols for both Gram negative (31) and Gram positive cell walls (33). Hybridisation was carried out with a formamide-free buffer. 165 166 Cells were counterstained with VECTASHIELD® Antifade Mounting Medium containing 1.5 µg ml ¹ 4,6-diamidino-2-phenylindole (Vector Laboratories, Burlingame, CA, USA). Images were 167 168 captured on a BX61 microscope equipped with a DP80 camera (Olympus Australia, Notting Hill,

A novel, dichloromethane fermenting bacterium

- 169 Australia) using Olympus cellSens Dimension software v2.1. Strain DCMF cell length and width
- 170 was determined from a sample of 20 cells using the linear measurement tool within the program.

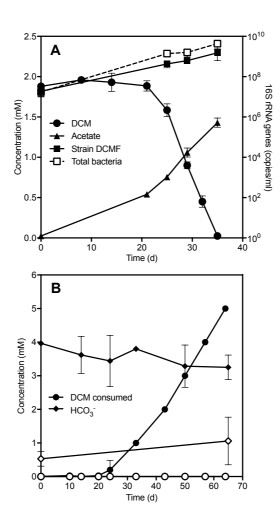
171 Results

172 Dichloromethane fermentation

- 173 Culture DFE consumed 1.9 ± 0.0 mM DCM within 35 days, yielding 3.7 ± 2.2 × 10⁸ strain DCMF
- 174 cells ml⁻¹ (Fig 1A), or $2.0 \pm 1.2 \times 10^{14}$ strain DCMF cells per mole of substrate consumed. The
- 175 product of DCM fermentation was acetate (1.4 ± 0.1 mM; Fig 1A), which was not observed in
- abiotic controls. DCM attenuation did not proceed in MOPS-buffered cultures free of bicarbonate
- 177 Fig 1B). However, in analogous cultures amended with 4 mM NaHCO₃, DCM attenuation was
- 178 observed, yet HCO_{3} concentrations did not significantly change (p = 0.11, two-tailed t-test
- 179 between days 0 and 65; Fig 1B).

A novel, dichloromethane fermenting bacterium

Holland et al (2020)



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181Figure 1 Strain DCMF ferments DCM to acetate, reliant on exogenous bicarbonate. (A) Strain DCMF182growth was concomitant with the depletion of DCM and formation of acetate. Error bars represent standard183deviation, n = 2. Substrate and product concentrations are quantified on the left y-axis; strain DCMF and184total bacterial 16S rRNA gene copies are quantified on the right y-axis. (B) In MOPS-buffered medium, DCM185consumption was only observed in the presence of bicarbonate (filled circles). Cumulative DCM186consumption is from repeat amendment of 1 mM DCM. Empty circles represent cultures with no exogenous187bicarbonate. Error bars represent standard deviation, n = 3.

188 Metabolism of quaternary amines and methanol

Of the additional substrates tested as sole energy source or with an electron acceptor, strain DCMF growth was observed when methanol, choline or glycine betaine (5 mM each) were supplied (Fig 2). Culture DFE consumed methanol (4.3 ± 0.2 mM) over 30 days, yielding 3.1 ± 0.1 mM acetate and $2.4 \pm 0.6 \times 10^9$ strain DCMF cells ml⁻¹ (Fig 2A). This corresponded to $5.7 \pm 1.4 \times$

A novel, dichloromethane fermenting bacterium

Holland *et al* (2020)

10¹⁴ strain DCMF cells per mole substrate utilised. No methanol depletion was observed in the
abiotic (cell-free) control.

195 Culture DFE consumed choline $(4.8 \pm 0.2 \text{ mM})$ within 25 days, producing $15 \pm 0.6 \text{ mM}$ acetate and 196 6.0 ± 1.1 mM methylamine (Fig 2B). Glycine betaine (4.7 ± 0.3 mM) was consumed within 21 days, 197 with production of 11 ± 0.4 mM of acetate and 4.5 ± 0.6 mM methylamine (Fig 2C). Trimethylamine, dimethylamine, sarcosine (methylglycine), and glycine were not detectable 198 199 throughout. Neither acetate nor methylamine were detected in abiotic controls, and the latter was 200 also absent from cultures amended with DCM. Strain DCMF cell proliferation aligned with the 201 consumption of these two substrates, yielding an increase of $1.4 \pm 0.4 \times 10^9$ and $5.3 \pm 0.4 \times 10^8$ 202 cells ml⁻¹ in choline- and glycine betaine-amended cultures, respectively (Fig 2B, C). This 203 corresponded to $3.0 \pm 0.9 \times 10^{14}$ cells per mole of choline, and $1.1 \pm 0.1 \times 10^{14}$ cells per mole of 204 glycine betaine utilised.

A novel, dichloromethane fermenting bacterium

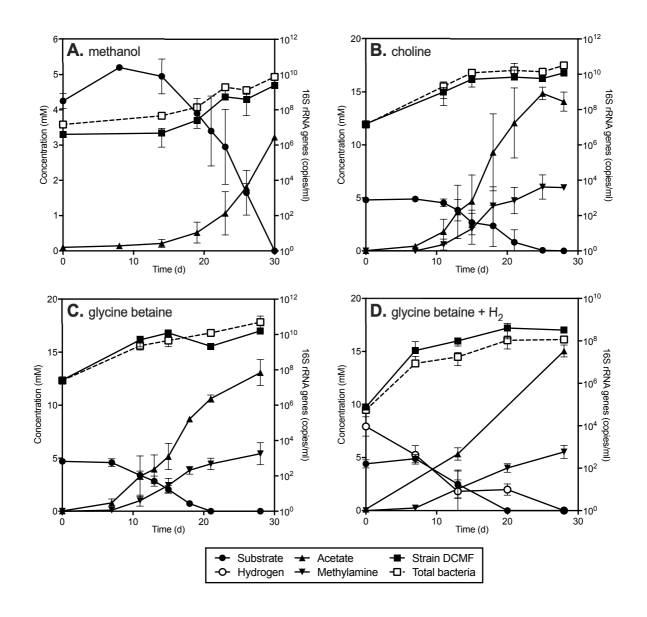




Figure 2 Strain DCMF growth was also correlated with the metabolism of methanol and quaternary amines. Strain DCMF growth correlated with the depletion of methanol and formation of acetate (A) and the depletion of choline (B) and glycine betaine (C) with formation of acetate and methylamine. Cultures amended with glycine betaine and hydrogen (D) did not produce trimethylamine, rather acetate and methylamine were once again the products. Substrate and product concentrations are quantified on the left y-axis; strain DCMF and total bacterial 16S rRNA gene copies are quantified on the right y-axis. Error bars represent standard deviation, *n* = 3.

DFE cultures amended with quaternary amine metabolic pathway intermediates dimethylglycine
and sarcosine + H₂ also demonstrated production of acetate and methylamine, which again
aligned with strain DCMF cell proliferation (Fig S2). Sarcosine was not degraded in the absence

A novel, dichloromethane fermenting bacterium

Holland *et al* (2020)

of H₂ (data not shown). Following the observation of strain DCMF growth and methylamine production in cultures amended with sarcosine + H₂, DFE cultures were also set up with glycine betaine + H₂ to determine whether glycine betaine could be reductively cleaved to trimethylamine and acetate. These cultures consumed all glycine betaine (4.4 ± 0.4 mM) and hydrogen (7.9 ± 0.9 mM) within 28 days, producing 15 ± 0.6 mM acetate and 5.5 ± 0.6 mM methylamine, but no trimethylamine (Fig 2D). Strain DCMF cell yields ($4.0 \pm 2.8 \times 10^8$ cells ml⁻¹) were similar to that when glycine betaine was the sole energy source.

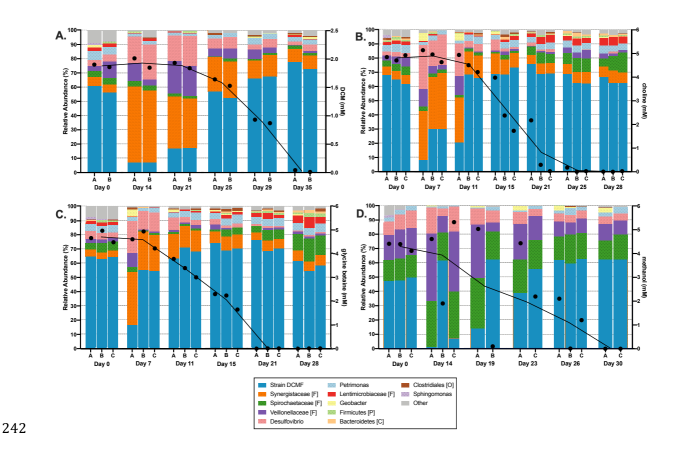
Culture DFE was unable to utilise CO, dibromomethane, ethanol, formate, syringic acid or
trimethylamine as sole energy sources (no growth and/or acetogenesis observed). Strain DCMF
was further unable to use any of the tested pairs of electron donors (acetate, choline, glycine
betaine, H₂, lactate, trimethylamine) and acceptors (CO₂, fumarate, Na₂SO₄, Na₂SO₃, NaNO₂, and
NaNO₃).

228 Shifts in DFE community composition in response to substrate consumption

229 Community profiling with Illumina 16S rRNA gene amplicon sequencing showed that culture DFE is composed of a limited number of taxa - only 12 amplicon sequencing variants (ASVs) were 230 present at $\geq 2\%$ relative abundance in > 1 sample (Fig 3). Community composition was similar in 231 232 cultures amended with DCM, choline, and glycine betaine, which had a common, DCM-amended 233 inoculum (Fig 3A,B,C), but was simplified in cultures that had been maintained on methanol for 234 two sub-cultivations and had a methanol-amended inoculum (Fig 3D; Fig S3A). While strain 235 DCMF was the dominant organism at the time of inoculation and during substrate consumption, its relative abundance decreased markedly in the lag phase prior to substrate consumption, 236 237 falling to as little as 0.96% in a methanol-amended replicate at day 14 (Fig 3). Taxa such as Synergistaceae (except in methanol-amended cultures, where this taxon was absent), 238 239 Desulfovibrio and Veillonellaceae increased in relative abundance during this lag phase, while Spirochaetaceae and Lentimicrobiaceae increased towards the end of and following substrate 240 241 depletion, particularly in quaternary amine-amended cultures (Fig 3).

A novel, dichloromethane fermenting bacterium

Holland et al (2020)



243 Figure 3 Culture DFE is subject to temporal shifts in community composition, with strain DCMF 244 dominant during substrate degradation. Illumina 16S rRNA amplicon sequencing was used to determine 245 DFE community composition (left y axis) at timepoints across the growth experiments amended with (A) 246 DCM, (B) choline, (C) glycine betaine, and (D) methanol reported in Figs 1 and 2. Taxa are reported down 247 to genus level where possible, otherwise taxonomic level is indicated in the legend ([F] = family, [P] =248 phylum, [C] = class, [O] = order). Reads with <1% abundance were filtered out in OIIME2. Unassigned reads 249 and taxa consistently <2% relative abundance were classed together as 'Other'. Substrate concentration (black circles, right y-axis) and a line connecting the mean substrate concentration at each time point is 250 251 overlaid on the community composition graphs. These are aligned with the time points written on the x-252 axis, not drawn to scale.

Differences in the DFE community were driven by the degree of substrate consumption (defined
in Table S3), more than the various substrates (Fig S3B). While there was no significant difference
in the Shannon diversity index between the samples when grouped by substrate (Kruskal-Wallis

A novel, dichloromethane fermenting bacterium

Holland et al (2020)

- p-value 0.0976; Fig S3C), there was a highly significant difference between all groups when
 clustered by substrate consumption (Kruskal-Wallis p-value <0.00001; Fig S3D).
- 258 *Exclusion of cohabitants as DCM and quaternary amine consumers*
- Attempts to isolate strain DCMF proved unsuccessful (24). Therefore, to test the hypothesis that strain DCMF was the sole consumer of DCM and quaternary amines, the cohabiting bacteria in culture DFE were enriched to the exclusion of strain DCMF (exclusion cultivation). This was achieved by dilution to extinction cultures on rich media amended with casamino acids, glucose, peptone or yeast extract. These growth conditions variously enriched *Bacillus, Desulfovibrio, Geobacter, Petrimonas*, and *Veillonellaceae*, but not strain DCMF (Fig S4E-H). *Spirochaetaceae* and *Synergistaceae* phylotypes did not grow on the tested rich media.
- 266 The strain DCMF-free cohabitant cultures were then tested for their ability to utilise DCM, choline,
- and glycine betaine. There was no significant substrate depletion in these cultures (Fig S4A-D),
- and therefore no evidence of DCM, choline, or glycine betaine degradation by the *Bacillus*,
- 269 *Desulfovibrio, Geobacter, Petrimonas,* or *Veillonellaceae* phylotypes in culture DFE.
- 270 Strain DCMF morphology
- 271 FISH microscopy enabled selective visualisation of strain DCMF cells, which appeared rod-shaped
- and occurred singly or in chains (Fig 4A). On average, strain DCMF cells were $1.69 \pm 0.27 \mu m \log 1000$
- and $0.64 \pm 0.12 \,\mu$ m wide. FISH images confirmed that strain DCMF numerically dominated culture
- 274 DFE during DCM dechlorination, congruent with community profiling results (Fig 4C).

A novel, dichloromethane fermenting bacterium

Holland et al (2020)

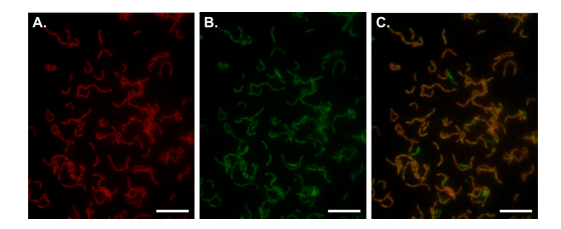




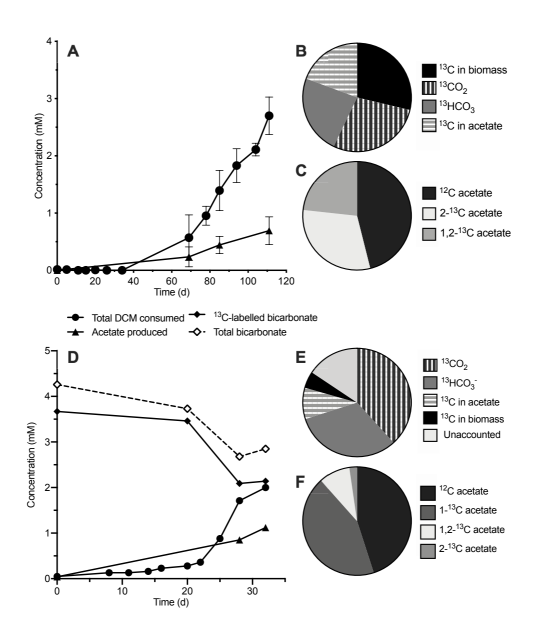
Figure 4 Morphology of strain DCMF. Fluorescence in situ hybridisation (FISH) microscopy images with
strain DCMF cells stained red with the Cy3-labelled Dcm623 probe (A), all bacterial cells stained green with
the 6-FAM-labelled Eub338i probe (B), and the overlay of Cy3- and 6-FAM-labelling in these images (C).
The scale bars represent 10 μM.

280 Strain DCMF is mixotrophic

281 To ascertain the fate of DCM carbon, triplicate DFE cultures were amended with [13C]DCM. When 282 2 700 \pm 328 μ M DCM had been consumed, 666 \pm 160 μ M of acetate was produced (Fig 5A), of which $47.1 \pm 5.5\%$ was unlabelled, $30.4 \pm 2.8\%$ was methyl group labelled ([2-13C]acetate), and 283 284 22.5 \pm 4.3% was both methyl and carboxyl group labelled ([1,2-13C]acetate; Fig 5C). A ¹³C mass balance was achieved by summing the measured concentrations of ¹³C-labelled carbon in acetate 285 (670 ± 289 μ M) and H¹³CO₃⁻ (815 ± 120 μ M) with the calculated concentrations of ¹³CO₂ in the 286 287 flask headspace (982 ± 144 μ M) and [¹³C]acetate equivalents in biomass (994 ± 121 μ M; Fig 5B, Table S4). This amounted to 128 ± 8.2% recovery of the labelled carbon, indicating no unknown 288 fate of DCM in culture DFE. 289

A novel, dichloromethane fermenting bacterium

Holland et al (2020)



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291 Figure 5 Strain DCMF assimilates carbon from DCM and bicarbonate to form acetate. 292 (A) Cumulative [¹³C]DCM consumption with concomitant with acetate production, including the ¹³C mass 293 balance from [¹³C]DCM (B) and proportion of labelled and unlabelled acetate (C). Error bars represent 294 standard deviation, n = 3. (D) Cumulative DCM consumption and acetate production in cultures amended 295 with H13CO3⁻. Total (labelled and unlabelled) aqueous HCO3⁻ is also shown (i.e. gaseous CO2 is not accounted 296 for here). Values in (D) are from a single representative culture as all triplicates had similar dechlorination 297 rates and product concentrations but began dechlorinating at different times. The ¹³C mass balance from 298 $H^{13}CO_3^{-}$ (E) and proportion of labelled and unlabelled acetate (F) is again shown. All pie charts represent 299 the average of triplicate cultures at the final time point.

A novel, dichloromethane fermenting bacterium

Holland *et al* (2020)

300	Analogous work was then carried out with unlabelled DCM in MOPS-buffered medium amended
301	with $^{\rm 13}\text{C}\xspace$ labelled bicarbonate, showing that strain DCMF incorporated carbon from CO $_{\rm 2}$ into the
302	carboxyl group of acetate. The culture consumed 2000 μM DCM and 2150 ± 492 μM ^{13}C from
303	bicarbonate. It produced 973 \pm 140 μM acetate (Fig 5D), of which 45.0 \pm 2.3% was unlabelled,
304	43.5 \pm 1.8% was labelled on the carboxyl group ([1-13C]acetate), 2.2 \pm 1.3% was labelled on the
305	methyl group, and 9.3 \pm 0.1% was labelled on both carbons (Fig 5F). A mass balance indicated
306	84.5 \pm 7.0% recovery of the labelled carbon in acetate (600 \pm 84.9 μM), the remaining $H^{13}CO_{3^{-}}$
307	(2280 ± 170 μM) and $^{13}CO_2$ (2740 ± 204 μM), and strain DCMF biomass (710 ± 9.74 μM ; Fig 5E,
308	Table S4).

309 **Discussion**

310 The DFE community

311 Strain DCMF is a novel bacterium present in enrichment culture DFE, which has been maintained with DCM as sole external source of energy for five years and at least 20 consecutive transfers 312 313 (24). Of the five other phylotypes previously reported in culture DFE, based on 16S rRNA genes 314 identified from genome sequencing data (24), four remained amongst the most abundant in the 315 present work (Desulfovibrio, Lentimicrobiaceae, Spirochaetaceae and Synergistaceae), while one 316 was no longer detected (Ignavibacteria). In combination with the similar community profiles observed across four different substrates, this suggests that culture DFE is a long-term stable-317 318 state community.

Illumina amplicon sequencing, FISH microscopy and qPCR all supported the previous observation
(24) of strain DCMF as the dominant organism in culture DFE during substrate consumption, and
linked growth of strain DCMF to depletion of DCM, methanol, choline, and glycine betaine.
Attempts to generate an axenic culture of strain DCMF have been unsuccessful, similar to the
DCM-mineralising bacterium '*Ca.* Dichloromethanomonas elyunquensis' in culture RM (19,23).
How the cohabiting organisms in both cultures persist despite numerous transfers and addition
of only a simple chlorinated compound (DCM) to minimal, anaerobic medium is a question of

A novel, dichloromethane fermenting bacterium

interest. While hydrogenotrophic acetogens and methanogens form major sub-populations in
 culture RM (18,19,23), culture DFE is non-methanogenic (24) and was unable to grow on H₂+CO₂
 alone.

329 Instead, the timing of the changes in relative abundance and known substrate range of major 330 phylotypes in culture DFE suggest that the cohabiting bacteria use cellular detritus resulting from 331 expired strain DCMF cells as an energy source (i.e. necromass fermentation). Some of the most 332 abundant cohabiting phylotypes in culture DFE - Desulfovibrio, Bacteroidetes (containing the 333 families Lentimicrobiaceae and Petrimonas), Spirochaetes/Treponematales, Synergistetes – have 334 previously been associated with hydrocarbon and organohalide-degrading mixed cultures (34-335 38), although their abundance was not linked to degradation of the primary substrate (with the 336 exception of some *Desulfovibrio* species) and some reports also suggested that they persist via necromass recycling (16,37–39). 337

Five of the 12 taxa in culture DFE were categorically excluded from being primary metabolisers of DCM, choline, and glycine betaine when tested in the absence of strain DCMF (Fig S4). *Spirochaetaceae* and *Synergistaceae* phylotypes could not be enriched to the exclusion of strain DCMF. However, their relative abundance during growth on DCM, choline, and glycine betaine diminished relative to strain DCMF, suggesting that it is unlikely that they are primary consumers of these substrates. This needs to be confirmed by proteomic assessment of the DFE community.

344 The role of the WLP in DCM metabolism

Amongst anaerobic DCM-dechlorinating bacteria, strain DCMF is unique in producing solely acetate as an end product (Fig 1A). *D. formicoaceticum* produced formate and acetate in a 2:1 molar ratio (15), while '*Ca.* Dichloromethanomonas elyunquensis' completely mineralised DCM to H₂, CO₂ and Cl⁻ (23). The latter organism is unique in also encoding and expressing reductive dehalogenases during growth with DCM (19,20). Despite these differences, both organisms utilise the WLP for DCM metabolism (15,20,23) as is likely the case with strain DCMF. Removal of

A novel, dichloromethane fermenting bacterium

bicarbonate from the culture medium precluded DCM dechlorination and ensuing work with ¹³Clabelled DCM and bicarbonate demonstrated that strain DCMF is mixotrophic, i.e. assimilates
carbon from both DCM and CO₂, similar to *D. formicoaceticum* (23).

354 These experiments also provided compelling evidence for the transformation of DCM to a WLP 355 intermediate, mostly likely methylene-tetrahydrofolate ($CH_2=FH_2$; Eq. 1). The production of 356 $H^{13}CO_3^{-1}$ from [13C]DCM suggested that $CH_2=FH_2$ is disproportionated into the WLP where it is 357 oxidised to HCO_3^{-1} (Eq. 2, Fig 6). The electrons released then reduce the remaining $CH_2=FH_2$ into 358 the methyl group of acetate (Eq. 3). However, the production of unlabelled acetate (47%) 359 indicates that the excess unlabelled HCO_{3} - (30 mM) in the medium is an alternative electron 360 acceptor to $CH_2=FH_2$ for acetogenesis (Eq. 4; Fig 6). The reduction of HCO_3^- to acetate requires twice as many electrons for acetate synthesis than CH₂=FH₂ (i.e. eight vs. four). Taking this ratio 361 into account, along with ~1:1 ratio of unlabelled to labelled acetate suggests that approximately 362 67% of electrons derived from DCM oxidation were directed toward HCO₃- reduction and 33% to 363 364 $CH_2 = FH_2$.

365	$4 \operatorname{CH}_2\operatorname{Cl}_2 + 4 \operatorname{FH}_4 \xrightarrow{} 4 \operatorname{CH}_2 = \operatorname{FH}_2 + 8 \operatorname{H}^+ + 8 \operatorname{Cl}^-$	Eq. 1
366	$3 \text{ CH}_2 = \text{FH}_2 + 9 \text{ H}_2\text{O} \rightarrow 3 \text{ HCO}_3^- + 12 \text{ e}^- + 3 \text{ FH}_4 + 15 \text{ H}^+$	Eq. 2
367	$CH_2=FH_2 + 4 e^- + HCO_3^- + 4 H^+ \rightarrow CH_3COO^- + H_2O + FH_4$	Eq. 3
368	$2 \text{ HCO}_{3^-} + 9 \text{ H}^+ + 8 \text{ e}^- \rightarrow \text{CH}_3\text{COO}^- + 4 \text{ H}_2\text{O}$	Eq. 4

The production of $[1,2^{-13}C]$ acetate from $[^{13}C]$ DCM is consistent with the reduction of $H^{13}CO_{3^{-1}}$ outlined above. However, the proportion (22.5%) was surprisingly high, given the relatively small contribution that labelled $H^{13}CO_{3^{-1}}$ from 2.7 mM $[^{13}C]$ DCM would make to the 30 mM unlabelled HCO₃⁻⁻ present in the culture medium. It is possible that co-localisation of WLP proteins in the cytoplasm may cause the reduction of $H^{13}CO_{3^{-1}}$ at a higher ratio than expected (i.e. 9%). Studies with $[^{13}C]$ DCM in *D. formicoaceticum* detected the ^{13}C label solely in the methyl group of acetate ([2-13C] acetate), congruent with DCM oxidation stopping at formate (23,40), while studies with

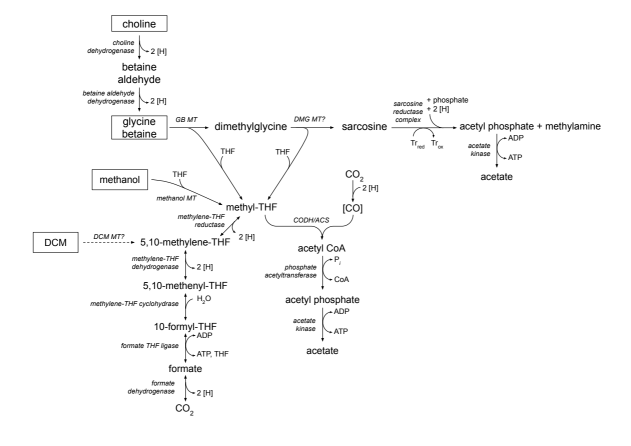
A novel, dichloromethane fermenting bacterium

Holland et al (2020)

another *Dehalobacterium* species in mixed culture that was capable of formate oxidation similarly

377 detected [1,2-¹³C]acetate (41).

378



379 Figure 6 Proposed model for metabolism of DCM, methanol and quaternary amines by strain DCMF.

The Wood-Ljungdahl pathway is central to transformation of all substrates into acetate. Proteins (with the exception of that catalysing the putative transformation of DCM to CH₂=FH₂, indicated by dotted arrow) are all identified in the strain DCMF genome and listed in Table S5. Abbreviations: CODH/ACS, carbon monoxide dehydrogenase/acetyl-CoA synthase; DCM, dichloromethane; DH, dehydrogenase; DMG, dimethylglycine; GB, glycine betaine; MT, methyltransferase; ox, oxidised; red, reduced; THF, tetrahydrofolate; Tr, thioredoxin.

386 DFE cultures amended with unlabelled DCM and ¹³C-labelled HCO_{3} - in MOPS-buffered medium 387 produced an analogous proportion of $[1-1^{3}C]$ acetate. A similar proportion of acetate (45.0%) to 388 that observed in the $[1^{3}C]$ DCM work was unlabelled, in this case evidently formed using unlabelled 389 HCO_{3} - produced from DCM. Thus, the ¹³C-labelling experiments support the hypothesis that DCM 390 metabolism involves the WLP and are consistent with the oxidation of formate to HCO_{3} -. As an

A novel, dichloromethane fermenting bacterium

Holland *et al* (2020)

exogenous supply of formate was unable to stimulate growth of culture DFE, strain DCMF alone is likely responsible for formate metabolism, which contrasts with the inability of *D. formicoaceticum* to further transform this metabolite (15). The production of $HCO_{3^{\circ}}$ from formate balances with its uptake during acetogenesis, congruent with a net flux of approximately zero, leading to the proposal that DCM is transformed as per Equation 5.

396
$$2 CH_2Cl_2 + 2 H_2O \rightarrow CH_3COO^2 + 5 H^2 + 4 Cl^2$$
 Eq. 5

397 Metabolism of non-chlorinated substrates

398 Strain DCMF is the first non-obligate anaerobic DCM-degrading bacterium. A genome-based 399 metabolic model previously suggested that the abundance of MttB superfamily 400 methyltransferases (named for their founding member, a trimethylamine:corrinoid 401 methyltransferase) encoded by strain DCMF may permit growth on methylamines amines and/or 402 glycines (24). While strain DCMF was unable to metabolise trimethylamine, growth was observed 403 with glycine betaine and the closely related compound choline. Both compounds are quaternary 404 amines with significant environmental roles. Glycine betaine is an osmoprotectant widely used by bacteria (42-44), marine algae (45), marine invertebrates (46), plants (47), and some 405 406 vertebrates (48). It is also an important source of nitrogen, comprising up to 20% of the total 407 nitrogen in hypersaline environments (49). Choline is typically more abundant, albeit as a part of 408 larger molecules including eukaryotic phospholipids, and can be converted to glycine betaine by 409 a near ubiquitous pathway in soil and water environments (50).

Accordingly, strain DCMF encodes both the choline dehydrogenase (Ga0180325_11215) and betaine aldehyde dehydrogenase (Ga0180325_114191) required for this transformation to glycine betaine. Based on the stoichiometry of observed end products, growth on putative pathway intermediates, and genomic information, we propose that strain DCMF likely stepwise demethylates glycine betaine to dimethylglycine and then sarcosine (methylglycine), which is then reductively cleaved to form acetate (via acetyl-phosphate) and methylamine

A novel, dichloromethane fermenting bacterium

Holland *et al* (2020)

(Supplementary Discussion and Fig 6). The electron equivalents produced from demethylation can be used for additional reduction of CO₂ to acetate via the WLP, as well as the reductive cleavage of sarcosine. This combination of demethylation and reductive cleavage has previously only been observed in *Sporomusa* spp. (51,52) and is a novel metabolic pathway within the *Peptococcaceae* family. A theoretical energy balance of the product formation and strain DCMF cell yield suggested that no other organisms in culture DFE were involved in quaternary amine metabolism (Supplementary Discussion).

423 The strain DCMF genome also encodes a number of methanol methyltransferases, which are likely 424 utilised for transformation of methanol into $CH_2=FH_4$ prior to its entry into the WLP 425 (Supplementary Discussion and Fig 6).

426 Environmental significance

The ability of strain DCMF to utilise choline, glycine betaine and methanol suggests that its 427 428 environmental relevance extends beyond DCM contaminated sites. Coastal salt marshes and 429 intertidal mudflats represent significant sources of methane from the demethylation of trimethylamine, which is in turn derived from quaternary amines (53–55). Both trimethylamine 430 and methanol are non-competitive methane precursors, which may allow large methanogen 431 432 populations to develop in environments where sulphate reduction would typically dominate (56,57). Indeed, trimethylamine is responsible for 60 - 90% of methane production in coastal salt 433 marshes and intertidal sediments (54,56). The transformation of quaternary amines to 434 methylamine by strain DCMF provides a pathway of lower methanogenic potential that could 435 operate in coastal subsurface environments. Strain DMCF does create acetate as a major end 436 product, which can be utilised by acetoclastic methanogens. However, unlike methylated amines, 437 438 methanogens have to compete with more thermodynamically favourable processes such as sulphate reduction for this substrate. 439

A novel, dichloromethane fermenting bacterium

Holland *et al* (2020)

Furthermore, DCM has recently also been recognised as a potent greenhouse gas with ozonedepleting potential (7), and oxygenated hydrocarbons such as methanol can influence atmospheric ozone formation through reactions with nitrous oxides (58). Therefore, although DCM, methanol, and quaternary amines are seemingly disparate substrates, they are closely linked to the atmospheric flux of climate-active gasses from anoxic, subsurface environments. This is both via the direct influence that DCM and methanol can have on ozone, and the indirect influence of quaternary amines on the flux of methylated amines and methane.

447 Provisional classification of strain DCMF as a novel genus and species

In addition to previously reported 16S rRNA gene phylogeny (24), whole genome analysis of 448 449 universally conserved marker genes and amino acid identity methods showed the closest relative 450 of strain DCMF to be *D. formicoaceticum* strain DMC (Supplementary Methods and Results). While 451 the former two methods support the placement of strain DCMF in a novel genus, the latter was on the borderline of suggested thresholds (Supplementary Results). However, the physiological 452 453 information presented here distinguishes strain DCMF from the sole representative of the genus 454 Dehalobacterium, which has thus far only proved capable of growth on DCM (15). Strain DCMF 455 also harbours a significantly larger genome than *D. formicoaceticum* (6.44 Mb for the former, 3.77 456 Mb for the latter) (22), which may account for its wider substrate range. Strain DCMF appears to be an anaerobic methylotroph, capable of metabolising a unique range of one-carbon compounds 457 458 (DCM, methanol) or substrates from which it can utilise methyl groups (choline, glycine betaine, 459 dimethylglycine, sarcosine). Thus, multiple lines of evidence support the placement of strain 460 DCMF within a novel genus in the family *Peptococcaceae*. As strain DCMF is not yet represented 461 in pure culture despite intensive efforts to isolate the organism, we propose it be classified in the *Candidatus* category (59) as '*Candidatus* Formamonas warabiya' strain DCMF gen. nov. sp. nov. 462

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A novel, dichloromethane fermenting bacterium

Holland *et al* (2020)

464 Description of 'Candidatus Formamonas' gen. nov.

465 *'Candidatus* Formamonas [Form.a.mon'as. L. adj. *formicum* relating to formic acid or, more
466 generally, one-carbon compounds; Gr. n. *monas* unit; ML. n. *Formamonas* the one-carbon utilising
467 unit.

'Candidatus Formamonas' is strictly anaerobic and metabolises one-carbon and methylated
compounds including DCM, methanol and quaternary amines glycine. Methylene/methyl groups
are metabolised via incorporation into the WLP. The type species is *'Candidatus* Formamonas
warabiya'.

472 Description of 'Candidatus Formamonas warabiya' sp. nov.

473 *'Candidatus* Formamonas warabiya [war.a.bi'ya N.L. n. *warabiya* the Dharawal name for the area
474 between Botany Bay and Bunnerong, honouring the Traditional Custodians of the land where this
475 bacterium was sampled from]. Permission was granted from the Dharawal Language Program
476 research group for use of this word as the species name.

477 Utilises DCM, methanol, choline, glycine betaine, dimethylglycine as sole sources of electrons 478 under anoxic conditions. Can also utilise the electron donor and acceptor pair H_2 and sarcosine. 479 The aforementioned substrates plus CO_2 are carbon sources. The primary fermentation product 480 is acetate. Methylamine is also produced from choline, glycine betaine, dimethylglycine, and 481 sarcosine + H_2 . The WLP is likely used for carbon fixation and metabolism of the methyl groups 482 removed from substrates. Cells are rod shaped (1.69 × 0.27 μm).

Type strain DCMF^T is not available in pure culture. The source of inoculum was contaminated sediment from the Botany Sands aquifer, adjacent to Botany Bay, Sydney, Australia. The type material is the finished genome of '*Candidatus* Formamonas warabiya' strain DCMF, which is 6.44 Mb and has a G+C content of 46.4% (GenBank accession number CP017634.1; IMG genome ID 2718217647).

A novel, dichloromethane fermenting bacterium

Holland *et al* (2020)

488 Acknowledgements

- 489 We are grateful to Kate Montgomery in the School of Biotechnology and Biomolecular Science,
- 490 UNSW, for her assistance with FISH microscopy. Thanks to Dr Valentina Wong for her initial
- 491 culturing assistance. SH was supported by an Australian Government Research Training Program
- 492 Scholarship.

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Holland et al (2020)

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