

Assessment of composition and origin of airborne bacteria in the free troposphere over Japan

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2 Japan

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5 Bacteria of the free troposphere over Japan

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24 **Abstract**

25

26 Long-range transport of airborne microorganisms through the free troposphere
27 significantly impacts biological ecosystems, human life, and atmospheric processes in
28 downwind areas. However, microbial communities in the free troposphere have rarely
29 been investigated because the direct collection of microbial cells at high altitudes
30 requires sophisticated sampling techniques. In this study, tropospheric air sampling was
31 performed using a balloon and an aircraft at 800 m and 3000 m, respectively, over the
32 Noto Peninsula in Japan (37.5°N, 137.4°E) where free tropospheric winds carry
33 aerosols from continental areas. The air samples were collected during four different
34 sampling periods when air masses came from desert regions of Asian continent (west
35 samples) and from Siberia of Russia North Asia (north samples). The west samples
36 contained higher levels of aerosols, and bacteria from the west samples grew in culture
37 media containing up to 15% NaCl. In contrast, bacteria from the north samples could
38 not be cultured in the same media. All isolates obtained from the NaCl-amended
39 cultures were similar to *Bacillus subtilis* and classified as *Firmicutes*. A 16S rDNA
40 clone library prepared from the west samples was mainly composed of one phylotype of
41 *Firmicutes* that corresponded to the cultured *B. subtilis* sequence. A clone library
42 prepared from the north samples consisted primarily of two phyla, i.e., *Actinobacteria*
43 and *Proteobacteria*, which are known to dominantly inhabit low-temperature
44 environments of North Asia. Our results suggest that airborne bacterial communities at
45 high altitudes include several species that vary by the direction and interaction of free
46 tropospheric winds.

47

48 **Key words:** Phylogeny, Asian dust, airborne bacteria, bioaerosol, halotolerant bacteria,

49 free troposphere

50 **1. Introduction**

51

52 Bioaerosols, which include bacteria, fungi, and viruses, are transported from
53 marine and terrestrial environments to the free troposphere and are significantly
54 abundant in the organic carbon fraction of atmospheric aerosols (Prospero et al., 2005).
55 Airborne microorganisms increase allergen burden causing increased incidence of
56 asthma (Ichinose et al., 2005) and contribute to dispersion of diseases such as Kawasaki
57 disease in humans (Rodó et al., 2011) and rust diseases in plants (Brown and
58 Hovmøller, 2002). Moreover, bioaerosols are thought to influence atmospheric
59 processes by participating in atmospheric chemical reactions and cloud particle
60 formation (Pratt et al., 2012).

61 The bacterial species composition of the atmosphere should be investigated for
62 understanding the characteristics of bacterial communities that are transported to long
63 distances and influence downwind ecosystems and climates. Aerosol sampling at
64 altitudes of 200–800 m above the ground level has demonstrated that bioaerosols are
65 composed of several species of bacteria (Li et al., 2010). The atmosphere is a
66 heterogeneous environment, and meteorological shifts can alter the bacterial species
67 composition of bioaerosols. The airborne bacterial abundance and species composition
68 at ground level in Asia (Hara and Zhang, 2012) and at 2700 m above sea level on North
69 American mountains (Smith et al., 2012) change significantly depending on Asian dust
70 events. However, few reports have directly investigated bacterial dynamics at high
71 altitudes, such as the free troposphere, where long-range transported bioaerosols are
72 abundant (Griffin 2004).

73 Halotolerant bacteria are tolerant to high salinity and resistant to stressors, such as
74 high pH, extreme temperatures, and desiccation (Lippert and Galinski, 1992). Indeed,
75 using NaCl-amendment culture techniques, viable halotolerant bacteria have been
76 detected from bioaerosols collected at high altitudes (Maki et al., 2008). Halotolerant
77 bacterial communities are typically common to bioaerosols transported hundreds or
78 thousands of kilometers (Echigo et al., 2005). Some halotolerant bacteria isolated from
79 sand dunes in the Gobi Desert were identical to bacterial species isolated in
80 Higashi-Hiroshima, Japan, suggesting their long-range transport (Hua et al., 2007). An
81 experimental design facilitating the isolation and identification of halotolerant bacteria
82 at high altitudes is expected to be useful for analyzing transported bacteria through the
83 free troposphere.

84 To investigate bacterial composition dynamics and the different air mass sources in
85 the free troposphere, we collected air samples at altitudes of 800 m and 3000 m above
86 the ground level over the edge of the Noto Peninsula, Japan. In this region, the air
87 masses moving from continental areas to Japan can be monitored while avoiding
88 aerosol contamination from local areas. We observed the amount of aerosols in air
89 samples microscopically, and estimated the trajectories of air masses during the
90 sampling periods. The viabilities of halotolerant microbial communities in air samples
91 were evaluated using culture media amended with various NaCl concentrations. The
92 bacterial species composition of the air samples was analyzed using clone-library
93 analysis targeting bacterial 16S rRNA genes.

94

95 **2. Materials and Methods**

96

97 **2.1. Sampling**

98 Aerosol samplings were performed over Suzu City (37.5°N, 137.4°E) during four
99 sampling periods. Suzu City is located on the northern coast of the Noto Peninsula,
100 Japan and is the arrival site for aerosols from continental areas. A balloon was used for
101 sampling over Suzu City from 11:00 to 12:00 on May 8, 2008 and from 10:50 to 11:50
102 on April 29, 2009. An aircraft was used for sampling from 14:50 to 16:50 on March 27,
103 2010 and from 11:50 to 13:50 on May 15, 2010. On March 27, 2010, the aircraft
104 traveled westward from Suzu City to a distance of 150 km single way and back (Fig. 1).
105 On May 15, 2010, from Suzu City, the aircraft traveled a distance of 150 km toward
106 northwest, returned to Suzu City, and traveled a distance of 150 km toward northeast.
107 The conditions of the four sampling periods are summarized in Table 1. The four
108 samples collected on May 8, 2008; April 29, 2009; March 27, 2010; and May 15, 2010
109 were named A, B, C, and D, respectively.

110 During the sampling periods on May 8, 2008 and April 29, 2009, the air samples
111 were collected at 800 m above the ground level using a tethered balloon (Maki et al.,
112 2008). An air pump with a sterilized filter holder was carried by the balloon and was
113 switched on at a specific altitude by a signal transmitted from the ground. An air sample
114 (700 l) was collected on a sterilized polycarbonate filter (0.22- μ m pore size; Whatman,
115 Tokyo, Japan) for 1 h. After sampling for an hour, the battery for the air pump failed at
116 800 m in the atmosphere.

117 On April 29, 2009 and May 15, 2010, aerosol samplings were performed at 3000 m
118 above the ground level using an aircraft with a 25-mm-diameter hole at the top. A

119 sterilized sampling tube (1.5 m in length) was inserted into the hole with one end of the
120 tube projecting outside. The other end of the tube was connected to a sterilized filter
121 holder (In-Line Filter Holder, 47 mm; Millipore, Tokyo, Japan) in the sampling device.
122 Given the length and curvature of the sampling tubes used, a loss of particles exceeding
123 0.2 μm in diameter should be considered (Hermann et al., 2001) and less than 5% of
124 particles were lost in this sampling, but the loss could be neglected. Air samples (1400 l)
125 were collected on each sterilized polycarbonate filter for 2 h. The samples were
126 collected on two filters during each sampling period.

127

128 **2.2. Characteristics and trajectories of air masses**

129 Air quality and atmospheric data in the free troposphere were obtained from the
130 Wajima Meteorological Observatory of the Japan Meteorological Agency, which is
131 located at a distance of 100 km from the sampling sites. Environmental data were
132 collected using a radiosonde at 3:00 a.m. At altitudes of approximately 3000 m,
133 information regarding weather conditions, temperatures, relative humidities, wind
134 speeds, and wind directions were obtained for comparative analyses of air masses
135 (Table 1). The potential temperature (PT) on March 27, 2010 suggested the presence of
136 typical free tropospheric air. It also suggested that slight cold air had activated
137 small-scale convection, causing the sky to become cloudy. PT on May 15, 2010
138 indicated that weak anticyclones may be prevalent in this region. Changes in aerosol
139 transportation are primarily controlled by the prevailing air flowing from China, the
140 anticyclonic circulation over the north-central East China Sea, and the subsiding
141 continental outflow air with low-level transport over Korea and Japan. Occasionally, a

142 cyclonic flow originates from the western North Pacific or from the East China Sea
143 contributing to the atmospheric conditions in this region. Therefore, isentropic back
144 trajectory analysis was applied to understand the primary transport patterns affecting
145 aerosols collected on March 27, 2010 and May 15, 2010.

146 To track the transport pathways of air masses, 72-h backward trajectories were
147 calculated using the NOAA Hybrid Single Particle Lagrangian Integrated Trajectory
148 (HYSPLIT) model (<http://www.arl.noaa.gov/HYSPLIT.php>). The location of the
149 backward trajectory start point was used as the sampling location for this study (37.5°N,
150 137.4°E) with altitudes of 2900; 3000; and 3100 m above the ground level for
151 estimating the accurate trajectories of air masses in the free troposphere.

152

153 **2.3. Microscopic analysis of particle abundance**

154 Within 2 h of sampling, aerosols were washed off the filters by shaking with 10 ml
155 of sterilized water containing 0.9% (w/v) NaCl. After washing, aliquots of 8 ml were
156 fixed with paraformaldehyde at a final concentration of 1%. Samples were stained with
157 4, 6-diamidino-2-phenylindole (DAPI) at a final concentration of 0.5 µg/ml for 15 min
158 and filtered through a polycarbonate filter (0.22-µm pore-size; Whatman) stained with
159 Sudan Black (Russell et al., 1974). After the filter was placed on a slide on a drop of
160 low-fluorescence immersion oil, a second drop of oil was added and the coverslip was
161 placed. The prepared slides were observed under an epifluorescence microscope
162 (Olympus, Tokyo, Japan) equipped with a UV excitation system. A filter transect was
163 scanned, and mineral particles, yellow particles and bacterial cells on the transect were
164 counted. The detection limit of aerosols was below 5×10^5 particles/m³ of air.

165

166 **2.4. Physiological experiments**

167 The remaining 2 ml of aerosols obtained after washing the filters with 10 ml of
168 0.9% (w/v) NaCl solution were used as cultivation spike in media containing various
169 NaCl concentrations to assess the viabilities of halotolerant bacteria in the air samples.
170 The washed solution (0.5 ml/sample) was inoculated into 19.5 ml of trypticase
171 soypeptone (TS) liquid medium (17 g trypticase peptone, 5 g phytone peptone, 2.5 g
172 K_2PO_4 , and 2.5 g glucose in 1 l pure water) containing NaCl at final concentrations of
173 0%, 3%, 10%, or 15% (w/v). TS medium has often been used for detecting
174 microorganisms from air samples. Microorganisms in the air samples were cultivated in
175 the media at 20°C in the dark. Microbial growth was estimated every 2 days by
176 measuring the absorbance at 550-nm.

177

178 **2.5. Identification of bacterial isolates by amplifying 16S rRNA sequences**

179 After 12 days of incubation, 1 ml of the microbial culture was used for bacterial
180 isolation by the spread-plate method using TS agar plates. After the bacterial isolates
181 were incubated in 10 ml of TS medium for 3 days, the bacterial cells were collected by
182 centrifugation at $20000 \times g$ for 5 min. Genomic DNA (gDNA) was extracted from the
183 bacterial cell pellets using SDS, proteinase K, and lysozyme, as described previously
184 (Maki et al., 2008). gDNA was purified by phenol–chloroform extraction, chloroform
185 extraction, and ethanol precipitation. Fragments of 16S rDNA (approximately 1450 bp)
186 were amplified from the extracted gDNA by PCR using the following oligonucleotide
187 primers: 27F, 5'-AGA GTT TGA TCM TGG CTC AG-3'; 1492R, 5'-GGY TAC CTT

188 GTT ACG ACT T-3' (Maidak et al., 1997). Thermal cycling was performed using a
189 Program Temp Control System PC-700 under the following conditions: denaturation at
190 94°C for 1 min, annealing at 56°C for 2 min, and extension at 72°C for 2 min for 30
191 cycles. PCR amplicons were purified by phenol–chloroform extraction, chloroform
192 extraction, and ethanol precipitation. The amplicons were sequenced using a Dye
193 DeoxyTM Terminator Cycle Sequencing Kit (ABI, CA, USA) and an ABI Prism 373A
194 DNA Sequencer according to the manufacturer's recommended protocols. The primers
195 27F and 1492R were used as the sequencing primers. The amplicon sequences were
196 searched against the DNA Data Bank of Japan (DDBJ) using BLAST. A phylogenetic
197 tree including all sequences was constructed according to the neighbor-joining
198 algorithm using TreeViewPPC (Saitou and Nei, 1987).

199

200 **2.6. Bacterial 16S rDNA clone libraries**

201 The 10 ml filter wash solution was used to estimate bacterial species composition
202 by clone library analysis targeting 16S rDNA. gDNAs from the solution were extracted
203 and purified as described in Section 2.5. Fragments of 16S rDNA (approximately 1450
204 bp) were amplified from the extracted gDNA by PCR using the following
205 oligonucleotide primers: 27F and 1492R. Thermal cycling was performed using the
206 Program Temp Control System PC-700 under the following conditions: denaturation at
207 95°C for 1 min, annealing at 55°C for 2 min, and extension at 72°C for 2 min for 30
208 cycles. PCR amplicons corresponding to 16S rDNA fragments were cloned into
209 *Escherichia coli* using a commercially available vector plasmid with a TA Cloning Kit
210 (Invitrogen, CA, USA) according to the manufacturer's protocol. Some clones were

211 obtained for each sample, and the sequences were determined as described in Section
212 2.5, except that M13 forward primer was used as the sequencing primer.

213

214 **2.7. Accession numbers**

215 DDBJ accession numbers for the 16S rDNA sequences determined in this study are
216 shown in Table 2.

217

218 **3. Results**

219

220 **3.1. Microscopic observation of aerosols**

221 DAPI-stained mineral particles collected at 3000 m on March 27, 2010 were
222 observed by epifluorescence microscopy as relatively large particles emitting
223 white–blue self-fluorescence with a diameter of $>5 \mu\text{m}$ (Fig. 2). DAPI-stained bacteria
224 were observed as coccoid-like particles with a diameter of $<1 \mu\text{m}$ and bright-blue
225 fluorescence. They were attached to the mineral particles. Yellow fluorescent particles,
226 potentially organic matter, were observed to range in diameter from $0.2 \mu\text{m}$ to $10 \mu\text{m}$.

227 DAPI-stained samples A and C included substantial concentrations of mineral and
228 yellow fluorescent particles (approximately 10^6 particles/ m^3 ; Table 1), as observed by
229 epifluorescence microscopy. The total densities of bacterial cells in samples A and C
230 were $(18.4 \pm 3.2) \times 10^6$ particles/ m^3 and $(2.28 \pm 0.83) \times 10^6$ particles/ m^3 , respectively.
231 In contrast, samples B and D contained particle concentrations below the detection limit
232 (i.e., 5×10^5 particles/ m^3) for microscopic observation.

233

234 **3.2. Physiological cultures**

235 Microbes in samples A and C grew in TS liquid media containing 0%, 3%, and
236 10% NaCl, as indicated by a rapid increase in the absorbance at 550 nm to >95
237 (approximately 4×10^7 cells/ml) within 5 days of incubation and fluctuated in the range
238 10–420 during the experimental period (Fig. 3a, c). Samples A and C, amended with
239 15% NaCl, demonstrated minimal microbial growth from the 8th and 4th day,
240 respectively, and the absorbance gradually increased to >10 during the experimental
241 period. These results indicated that microorganisms capable of tolerating up to 15%
242 NaCl maintained their viabilities in samples A and C. In contrast, no microbial growth
243 was observed from samples B and D in any culture medium during the incubation
244 period (Fig. 3b, d). Uninoculated culture medium in all experiments indicated no
245 microbial growth during the experimental period, suggesting that there was no
246 microbial contamination.

247 Colonies on the agar plates with the NaCl-amended cultures of samples A and C
248 were isolated on the basis of colony formation and colors. Consequently, a total of 8
249 isolates were obtained from each NaCl-amended culture. Sequencing of 16S rDNA
250 indicated that all the 8 isolates belonged to *Firmicutes* and were closely related to
251 *Bacillus subtilis* with >99.9% similarity (Table 2).

252

253 **3.3. Comparison of 16S rDNA clones**

254 The 16S rDNA fragments in the air samples were amplified by PCR using primers
255 targeting eubacterial 16S rDNA. The PCR amplicons were cloned into *E. coli*, and a
256 total of 201 clones including eubacterial 16S rDNA fragments were obtained from the

257 four samples. Sequences of the 16S rDNA clones indicated that the bacterial
258 populations were divided into 10 phylotypes (sequences with >98% similarity; Table 2).
259 The majority of phylotypes recovered from the four samples belonged to the phyla
260 *Firmicutes*, *Bacteroidetes*, *Proteobacteria*, and *Actinobacteria* that are typically well
261 represented in 16S rDNA clone libraries generated from terrestrial and marine
262 environments.

263 More than 85% of clones derived from samples A and C belonged to *Firmicutes*,
264 and all *Firmicutes* sequences corresponded to a single phylotype that was closely
265 related to *B. subtilis* (>99.7% similarity). Isolates obtained from NaCl-amended cultures
266 were identical to *B. subtilis* (Fig. 4). The complete 16S rDNA sequences of isolated *B.*
267 *subtilis* indicated high similarities (>99.7%) with *B. subtilis* detected hundreds of meters
268 above the Taklamakan Desert, China and from dust layers in the snow cover of Mount
269 Tateyama, Japan (2450 m). Sample C also included other bacterial species assigned to
270 the phyla *Bacteroidetes* and *Proteobacteria* (Fig. 4). Of these, a phylotype comprising
271 four clones belonging to *Bacteroidetes* was related to *Owenweeksia hongkongensis* at
272 low similarity (<88.5%). The one remaining clone belonged to the family
273 *Xanthomonadaceae* in *Gammaproteobacteria* and was closely related to
274 *Pseudoxanthomonas byssovorax* (99.3% similarity).

275 All 20 clones derived from sample B were affiliated with *Proteobacteria*, including
276 *Alphaproteobacteria* and *Betaproteobacteria*. The *Alphaproteobacteria* included 10
277 clones (50%) that were identical to *Sphingomonas rhizogenes* (100% similarity; Table 2,
278 Fig. 4). In *Betaproteobacteria*, 7 clones (35%) were closely related to *P. fluorescens*
279 (100% similarity) and 3 clones (15%) were similar to bacterium P618 (100%

280 similarity).

281 Isolates from sample D were dominated by *Proteobacteria* and *Actinobacteria*
282 sequences. Of the 71 clones derived from sample D, 55 belonged to *Proteobacteria*.
283 Forty-two (55%) of the isolates were related to *Variovorax paradoxus* (99.7%–100%
284 similarity), and 13 (18%) were closely related to *Methylobacterium* spp. (99.7%–100%
285 similarity; Table 2, Fig. 4). *Actinobacteria* included 13 clones (18%) that belonged to
286 *Brevibacterium* and were related to *Brevibacterium* sp. SA312 (>99.8% similarity).

287

288 **3.4. Transport trajectory**

289 Backward trajectory analysis indicated that the air-mass sources could be classified
290 into two types across the four sampling periods. The air masses sampled on May 8,
291 2008 and March 27, 2010 from the desert area of Asia had passed over the industrial
292 area in China and across the Sea of Japan (Fig. 5a, c). In contrast, the air masses
293 sampled on April 29, 2009 and May 15, 2010 were from North Asia areas, such as
294 eastern Siberia and the Japanese island, Hokkaido and had passed along the Sea of
295 Okhotsk to Suzu City (Fig. 5b, d).

296

297 **4. Discussion**

298

299 **4.1. Bioaerosols in the free troposphere**

300 Bioaerosols originating from Asia are dispersed in downwind regions such as
301 Korea and Japan by the prevailing westerly winds in the middle latitudes and are
302 sometimes carried to the Pacific Ocean (Iwasaka et al., 2009). Long-range transport of

303 microorganisms contributes to microbial dispersal and significantly impacts ecosystems,
304 human health, agricultural productivity, and climate in downwind areas (Jaenicke,
305 2005 ; Brown and Hovmøller, 2002). In this study, epifluorescence microscopy
306 demonstrated that aerosols collected at 3000 m contained large particles attached with
307 microorganisms, such as bacteria (Fig. 2). The bacterial populations were possibly
308 transported from other regions and dispersed to Japanese environments. Because
309 atmospheric dispersion transports microorganisms to long distances, airborne bacterial
310 composition should be compared among air masses from different continents to better
311 understand bacterial dynamics in downwind regions (e.g., Noto Peninsula, Japan).

312 Samples A and C came from desert regions of Asia, whereas samples B and D
313 came from Siberia and Hokkaido (Fig. 5). Samples A and C contained greater aerosol
314 concentrations than samples B and D and included significant amounts of minerals,
315 potential organic components, and bacterial particles (Fig. 2, Table 1). DAPI-stained
316 particles with yellow fluorescence have been reported to resemble organic materials
317 originating from proteins and other microbial cell components (Mostajir et al., 1995).
318 The large sizes of minerals and organic particles could shelter bacterial cells against
319 environmental stressors such as UV irradiation and desiccation.

320 Dust events have been reported to increase the number of airborne bacteria in
321 correspondence with the number of mineral particles (Hara and Zhang, 2012; Prospero
322 et al., 2005). In East Asia during spring and summer, the prevailing westerly winds
323 constantly carry dust particles, creating weak Asian dust events at 4000 m (Iwasaka et
324 al., 1988). During the May 8, 2008 and March 27, 2010 sampling periods, the westerly
325 wind was believed to carry bioaerosol-associated mineral particles to high altitudes

326 above Suzu City. In fact, the Ozone Monitoring Instrument ([http://jwocky.gsfc.
327 nasa.gov/](http://jwocky.gsfc.nasa.gov/)) and light detection and ranging (lidar) measurements at Toyama City, Japan,
328 revealed that dust particles were transported to Japan on May 8, 2008 (Maki et al.,
329 2010).

330 As described in Section 2.1, sampling losses of particles exceeding 0.2 μm in
331 diameter could be neglected in this sampling. Therefore, our discussions of bacterial
332 species composition are accurate for particles with diameters of 0.2-2.0 μm . Few
333 investigators have examined the ratio between particle size and bacterial species
334 composition. An understanding of this relationship can lend insight to bacterial
335 transportation processes on the global scale, and further investigation is required on this
336 relationship.

337

338 **4.2. Bacterial communities from western and northern areas**

339 Samples A and C included viable halotolerant bacteria that grew in culture media
340 containing up to 15% NaCl (Fig. 3). All isolates obtained from the NaCl-amended
341 cultures were identical to *B. subtilis* and were abundant in the 16S rDNA clone libraries
342 from samples A and C (Table 2). Denaturing gradient gel electrophoresis analysis using
343 PCR products demonstrated that all bacteria grown in the NaCl-amended cultures
344 corresponded to *B. subtilis* (data not shown). *Bacillus* spp. form resistant endospores to
345 enhance their survival in the atmosphere (Nicholson et al., 2000). Halotolerant bacteria
346 identified as *B. subtilis* were dominantly associated with mineral particles collected at
347 high altitudes above the Taklamakan Desert (Maki et al., 2008) and from accumulated
348 aerosols in the snow cover of Mount Tateyama (Maki et al., 2011). Species related to *B.*

349 *subtilis* were isolated from sand samples of the Gobi Desert (Hua et al., 2007) and
350 dominates the surface air of Saul City during Asian dust events (Jeon et al., 2011). From
351 a free-tropospheric sampling on the North American mountains, isolates of *Bacillus* spp.
352 were mainly obtained from air samples carried by Asian dust events (Smith et al., 2012).
353 Presumably, halotolerant *B. subtilis* in samples A and C maintained their viability at
354 high altitudes and were carried from continental desert areas by westerly winds.

355 The clone libraries obtained from samples B and D were mainly dominated by
356 *Proteobacteria* and/or *Actinobacteria* (Table 2, Fig. 4). Terrestrial bacteria in Siberia
357 primarily belonged to *Proteobacteria* (Zhou et al., 1997). In addition, marine bacterial
358 communities in the Antarctic Sea are primarily composed of *Proteobacteria*
359 (Brinkmeyer et al., 2003). The *Proteobacterium*, *V. paradoxus*, that was predominant in
360 sample D is abundant in the snow cover of Mount Tateyama (Segawa et al., 2005) and
361 has been isolated from a Greenland glacier ice core (Sheridan et al., 2003).
362 *Actinobacteria* sequences from sample D were dominated by *Brevibacterium* spp. that
363 originated from soil samples in the Arctic and Antarctica, as confirmed against DDBJ.
364 *Pseudomonas* spp. identified in sample B were 100% similar to *P. fluorescens*
365 originating from polar regions (Berg et al., 2009). *Sphingomonas* sp. from sample B was
366 identical to *S. rhizogenes* and *Sphingomonas* spp. detected from Lake Baikal, Siberia,
367 and Antarctica (Dieser et al., 2010). Members of *Sphingomonas* were often included in
368 marine bacterial communities in North Asia and polar regions (Gloeckner et al., 2000).
369 Several of the bacterial communities in samples B and D could have been carried to
370 high altitudes by the north wind that originated above low-temperature environments in
371 North Asia.

372 The 16S rDNA clone libraries from samples A and C were composed of bacterial
373 species belonging to *Firmicutes* and/or *Bacteroidetes*. In contrast, the clone libraries
374 from samples C and D primarily included sequences of *Proteobacteria* and/or
375 *Actinobacteria* (Fig. 4). Thus, the bacterial species compositions of the four sampling
376 dates were stratified into two different types depending on the air mass sources (Gobi
377 Desert and North Asia). We previously found that when atmospheric wind directions in
378 Kanazawa City, Japan changed from west to north following an Asian dust event during
379 the first week of May 2011, airborne bacterial compositions at 10 m above ground
380 changed from primarily *Firmicutes* (*B. subtilis*) to *Proteobacteria* (data not shown).
381 These results suggest that bacterial communities at high altitudes varied among the four
382 sampling periods during which the air mass sources were from the west and north.
383 However, *Proteobacteria* detected in samples B and D did not overlap (Fig. 4). Since
384 *Proteobacteria* are represented sparsely in the free troposphere, they may be easily
385 affected by migrations of bacterial communities from several terrestrial and marine
386 environments along various air-mass trajectories.

387

388 **4.3. Influences of bacterial communities on ecosystems and human health**

389 The air samples collected from the free troposphere included several bacterial
390 species in the phyla *Firmicutes* and *Proteobacteria* that are often associated with plant
391 growth, human life, and organic matter cycles. Although most bacterial species detected
392 from air samples are non-pathogenic, a dominant *B. subtilis* strain in clinical
393 contaminants has been described as an opportunistic pathogen (Thomas and Whitte,
394 1991). Isolates from sample B comprising a minor phylotype of *Gammaproteobacteria*

395 were closely related to clinical and harmful pathogens, such as *Hafnia* and *Salmonella*
396 spp. (Ridell et al., 1994; Wang et al., 1997). Bacterial species of the genera
397 *Methylobacterium*, *Sphingomonas*, *Variovorax*, and *Bacillus* dominated the air samples
398 and were often found to be associated with leaf surfaces and in the rhizosphere (Idris et
399 al., 2004; Anda et al., 2011). *V. paradoxus* and *Bacillus* spp. have been reported to
400 promote plant growth (Maimaiti et al., 2007; Yadav et al., 2011), whereas *Bacillus*
401 includes several species of plant pathogens (Yoshida et al., 2001). Bacterial populations
402 on leaf surfaces or in the rhizosphere may become aerosolized from grass mowing and
403 can disperse to other regions. The *Variovorax*, *Bacillus*, and *Pseudomonas* sequences
404 dominating the air samples in this study were related or identical to the bacterial species
405 mineralizing organic matters, such as cellulose, and contributed to the carbon cycle in
406 terrestrial environments (Das and Mukherjee, 2007; Ulrich et al., 2008). Some strains of
407 *B. subtilis* ferment organic matters and are useful for the production of Japanese health
408 foods such as *natto* (Ashiuchi et al., 1998). The long-range dispersal of bacteria has
409 positive and negative implications for human societies, plant growth, and microbial
410 ecosystems.

411 Bacteria classified as *Xanthomonadaceae* and *Pseudomonadaceae* that are detected
412 in air samples as minor species facilitate ice-nucleation for cloud formation in the
413 atmosphere (Pratt et al., 2009; Morris et al., 2008). Clone library analyses of air samples
414 over high mountains have revealed that ice nuclei-forming bacteria were minor
415 components in the atmosphere, accounting for <1% of total clones (Bowers et al., 2009).
416 Members of *Xanthomonadaceae* were rare in the clone library from sample C. These
417 results suggest that air masses in the free troposphere contain a low number of bacterial

418 species that possess ice-nucleating activities.

419

420 **5. Conclusion**

421

422 To best of our knowledge, this is the first study to compare bacterial communities
423 in the atmosphere at 800 m and 3000 m over Asia during four sampling periods when
424 the air masses were transported from two different sources (the Gobi Desert and North
425 Asia). The air masses originating from the Gobi Desert included halotolerant bacteria
426 dominated by *B. subtilis* strains that are believed to have been carried by Asian dust
427 events. In contrast, air masses originating from North Asia did not include any
428 halotolerant bacteria and were primarily composed of *Proteobacteria* and
429 *Actinobacteria*. It is possible that bacterial communities at high altitudes exhibit
430 significantly different dynamics depending on the origin of the air mass. Further
431 investigations are required to establish correlations between bacterial species
432 composition and air mass sources. An understanding of the relationships between
433 aerosol sizes and bacterial species and amounts is essential for predicting the dispersal
434 conditions of bioaerosols around downwind environments.

435

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437

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444

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446

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590

591 **Figure Legends**

592

593 Fig. 1 Aircraft flight routes during the sampling periods: 14:50-16:50 on March 27,
594 2010 (solid line) and 11:50-13:50 on May 15, 2010 (dotted line).

595

596 Fig. 2 Epifluorescence micrograph of mineral particles attached to bacterial particles (a)
597 and yellow (organic) particles (b). The bioaerosols were collected on March 27, 2012.
598 Arrows indicate bacterial cells. All photomicrographs were taken at a magnification of
599 $\times 1000$. (scale bar = $5\mu\text{m}$).

600

601 Fig. 3 Microbial growth from samples A–D collected on May 8, 2008 (a), April 29,
602 2009 (b), March 27, 2010 (c), and May 15, 2010 (d), respectively, in media amended
603 with NaCl at concentrations of 0% (squares), 3% (circles), 10% (triangles), and 15%
604 (diamonds). All experiments were performed in five replicate tubes.

605

606 Fig. 4 Phylogenetic tree including the partial sequences of 16S rDNA amplicons
607 obtained from the clone libraries from air samples and from isolates grown in
608 NaCl-amended media. The phylogenetic tree was calculated from a dissimilarity matrix
609 of an approximately 330-bp alignment (*Escherichia coli* numbering 153 to 482) using a
610 neighbor-joining algorithm. The sample information and the accession number of each
611 reference sequence are given in parentheses. Open circles at branch points indicate that
612 bootstrap values obtained by neighbor-joining analysis exceeded 50% (after 1000
613 resamplings).

614

615 Fig. 5 Three-day backward trajectories of aerosols that arrived at Suzu City on May 8,
616 2008 (a), April 29, 2009 (b), March 27, 2010 (c) and May 15, 2010 (d).

617

Table 1 Sampling dat, meteological conditions, and particle concentrations during the sampling periods.

Air sample name		A	B	C	D
Sampring information	Sampling date	May 8th, 2008	Apr 29th, 2009	Mar 27th, 2010	May 15th, 2010
	Collection time	11:00 – 12:00 (1h total)	10:50 – 11:50 (1h total)	14:50 – 16:50 (2h total)	11:50 – 13:50 (2h total)
	Sampling method	baloon	baloon	aircraft	aircraft
	Sampling location ^{*1}	800m	800m	3000m	3000m
Troposheric meteological condition	Observed weather conditions	Clear	Clear	Cloudy skies	Clear
	Temperature (°C)	0	-6.2	-18.4	1.2
	% Relative humidity	100	3	85	4
	Predominat wind direction	W	NNE	W	NNE
Concentrations of particles (10⁶ particles /m³)*²	Mineral particles	8.84 ± 1.94	N.D ^{*3}	1.05 ± 0.79	N.D ^{*3}
	Yellow particles	6.95 ± 1.45	N.D ^{*3}	1.93 ± 0.70	N.D ^{*3}
	Bacterial cells	18.4 ± 3.2	N.D ^{*3}	2.28 ± 0.83	N.D ^{*3}

*1 Height above the ground.

*2 (particles/m³) indicates (particles per m³ of air).

*3 Particles were not detected under microscopic observation.

Table 2 Phylogenetic affiliation of sequences of 16S rDNA clones.

Air sample name	Numbers of Clones or strains ^{*1}	Names of sequences ^{*2}	Conditon	Length (bp)	Category	GenBank accession no.	Closest relative	Similarity (%) ^{*3}
A	65	SzDc-08May-1	directly extracted DNA	1431	<i>Firmicutes</i>	AB749769	<i>Bacillus subtilis</i> (JQ762447)	99.7 – 100
B	10	SzDc-09Apr-1	directly extracted DNA	510	<i>Alphaproteobacteria</i>	AB609064	<i>Sphingomonas rhizogenes</i>	100
	7	SzDc-09Apr-2	directly extracted DNA	465	<i>Gammaproteobacteria</i>	AB609063	<i>Pseudomonas fluorescens</i>	100
	3	SzDc-09Apr-3	directly extracted DNA	500	<i>Gammaproteobacteria</i>	AB609067	Bacterium P618 (JX12010)	100
C	60	SzDc-10March-1	directly extracted DNA	1452	<i>Firmicutes</i>	AB740157	<i>Bacillus subtilis</i>	99.8
	4	SzDc-10March-2	directly extracted DNA	1394	<i>Bacteroidetes</i>	AB740158	<i>Owenweeksia hongkongensis</i>	88.4 – 88.5
	1	SzDc-10March-3	directly extracted DNA	1349	<i>Gammaproteobacteria</i>	AB740159	<i>Pseudoxanthomonas byssovorax</i>	93.6
D	42	SzDc-10May-1	directly extracted DNA	509	<i>Betaproteobacteria</i>	AB769480	<i>Variovorax paradoxus</i>	99.8 – 100
	13	SzDc-10May-2	directly extracted DNA	482	<i>Alphaproteobacteria</i>	AB769478	<i>Methylobacterium</i> sp. SKJH-1	99.8 – 100
	13	SzDc-10May-3	directly extracted DNA	526	<i>Actinobacteria</i>	AB769479	<i>Brevibacterium</i> sp. SA312	99.8 – 100
	3	SzDc-10May-4	directly extracted DNA	476	<i>Firmicutes</i>	AB769477	<i>Streptococcus australis</i>	100
A	4	08Szi-1	<15%NaCl	1426	<i>Firmicutes</i>	AB749540	<i>Bacillus subtilis</i> (AY553094)	100
C	3	10Szi-1	<10%NaCl	1409	<i>Firmicutes</i>	AB740155	<i>Bacillus subtilis</i> (GU826163)	100
	1	10Szi-4	15%NaCl	1426	<i>Firmicutes</i>	AB740156	<i>Bacillus subtilis</i> (HQ425655)	99.9

*1 The numbers of the clones in 16S rDNA clone libraries and the strains of culture isolates.

*2 Isolates from the NaCl amended cultures are named as the Szi serie . Clones of 16S rDNA library were named as the SzDc serie.

*3 Similarity value between each isolate and the closest relative in databases.

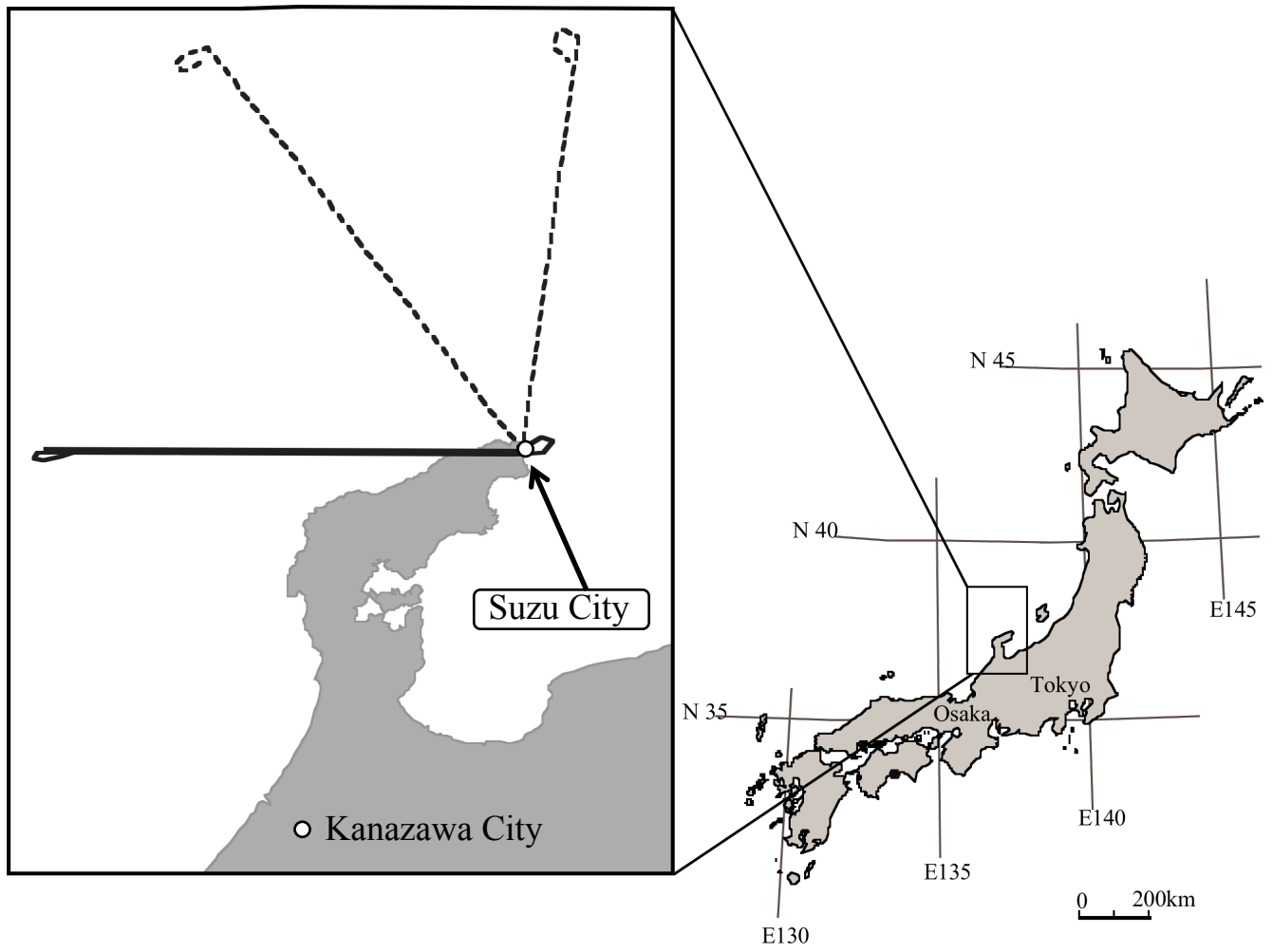


Fig. 1 T.Maki et al.

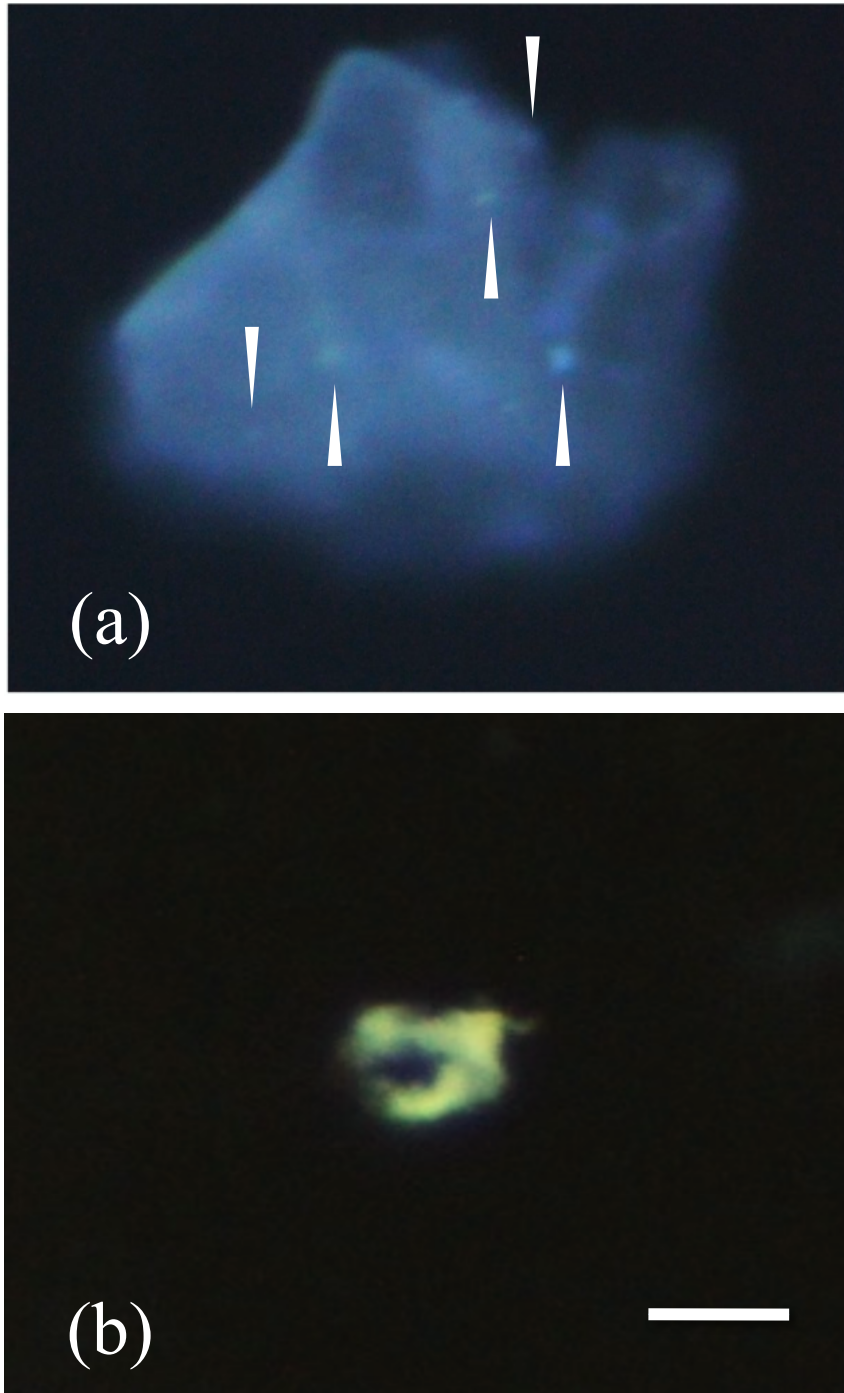


Fig. 2 T.Maki et al.

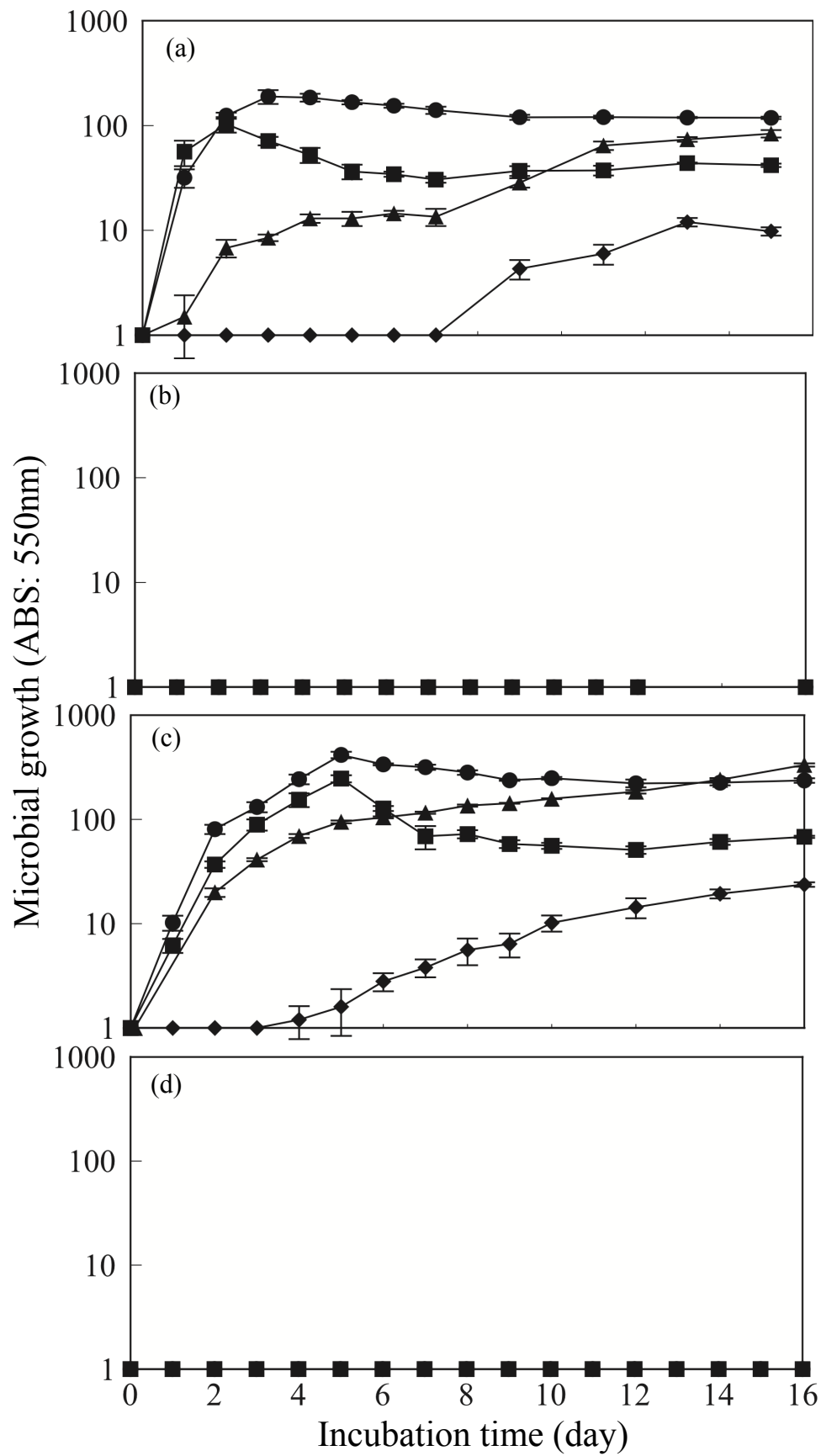


Fig. 3 T.Maki et al.

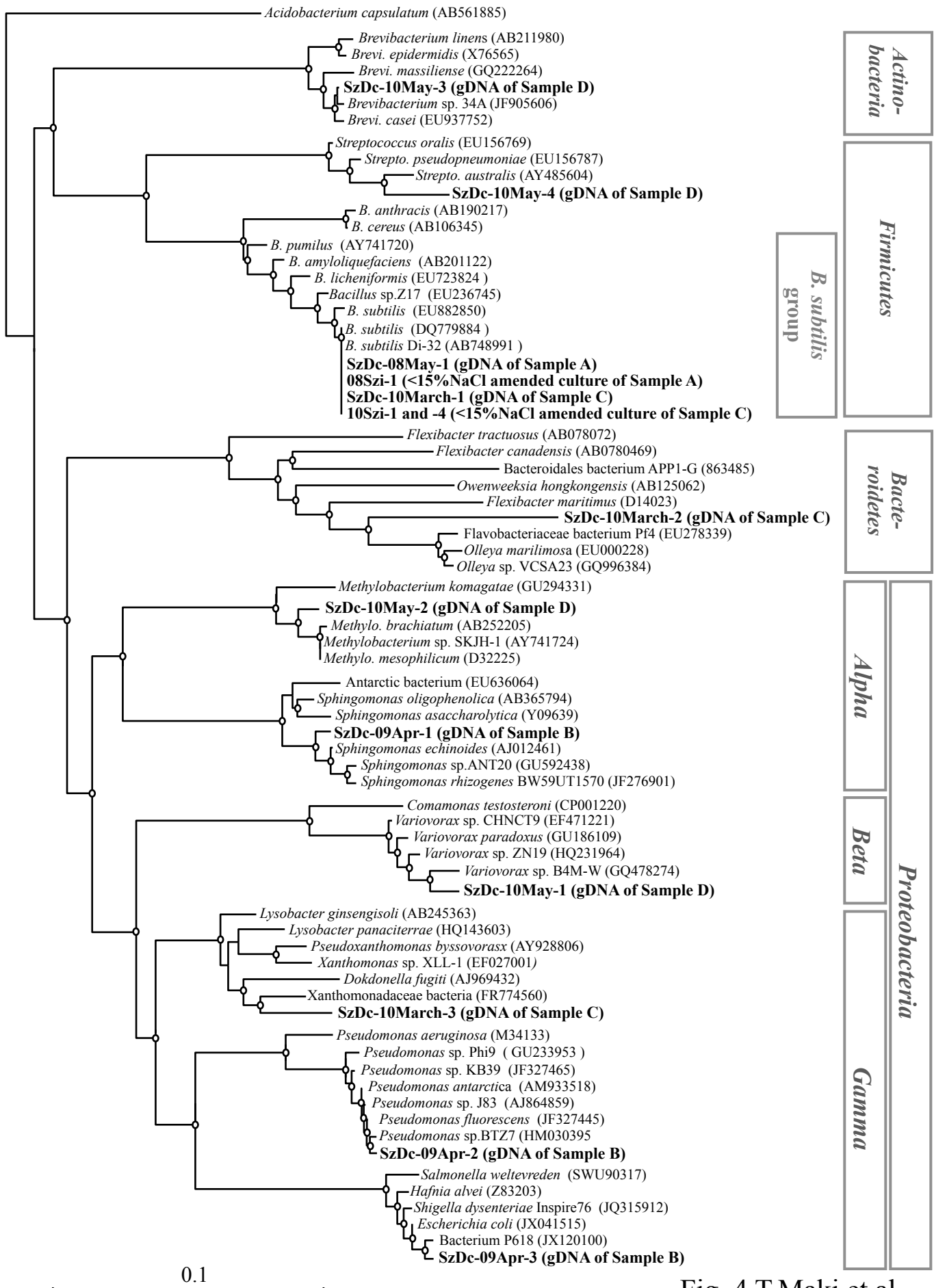
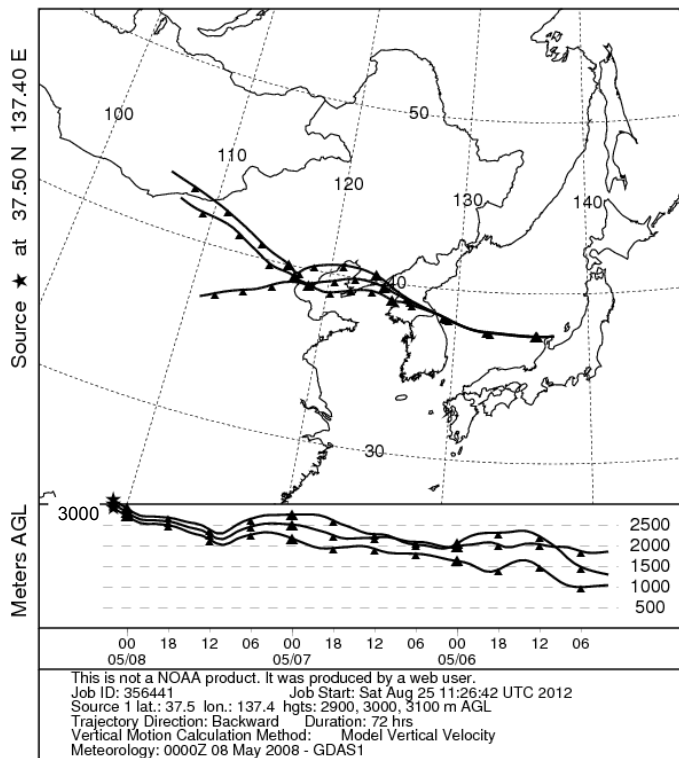
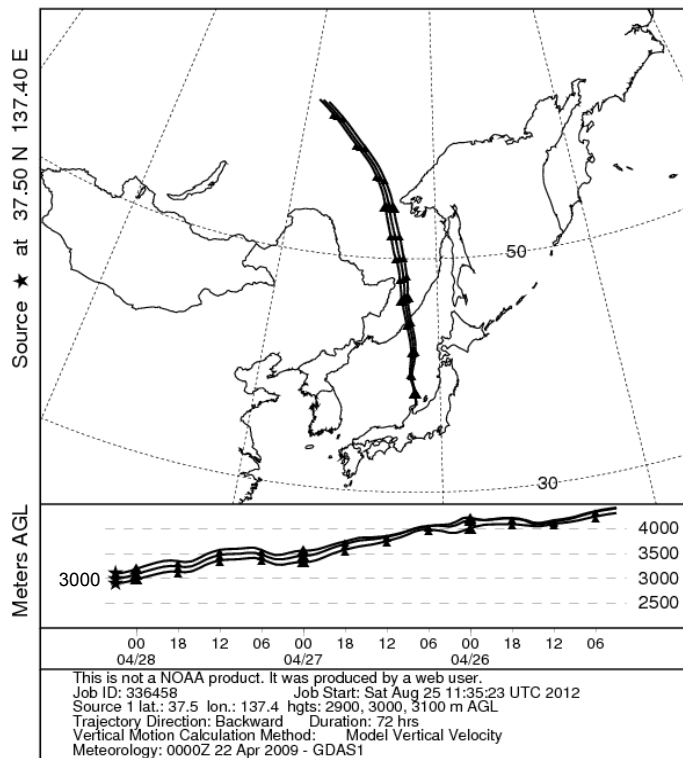


Fig. 4 T.Maki et al.

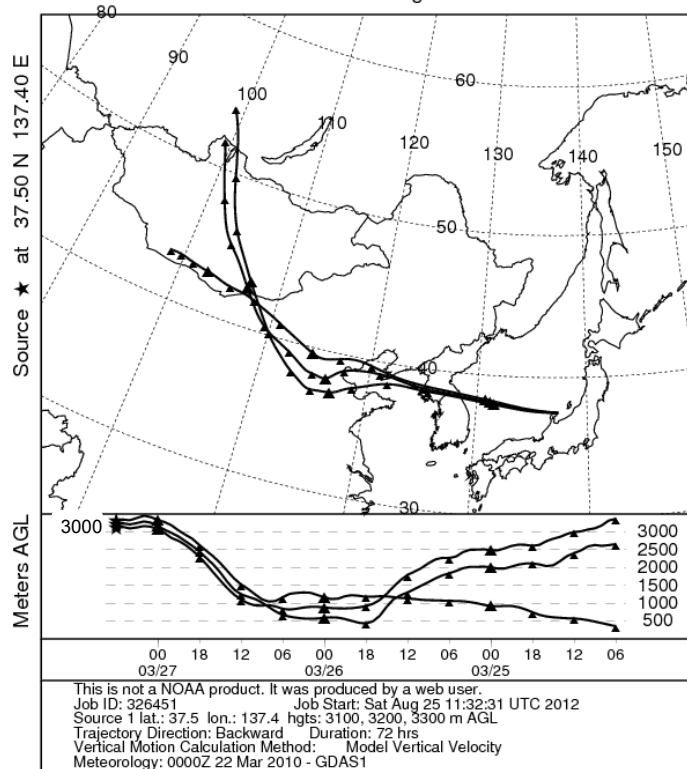
(a) NOAA HYSPLIT MODEL
Backward trajectories ending at 0200 UTC 08 May 08
GDAS Meteorological Data



(b) NOAA HYSPLIT MODEL
Backward trajectories ending at 0300 UTC 28 Apr 09
GDAS Meteorological Data



(c) NOAA HYSPLIT MODEL
Backward trajectories ending at 0600 UTC 27 Mar 10
GDAS Meteorological Data



(d) NOAA HYSPLIT MODEL
Backward trajectories ending at 0300 UTC 15 May 10
GDAS Meteorological Data

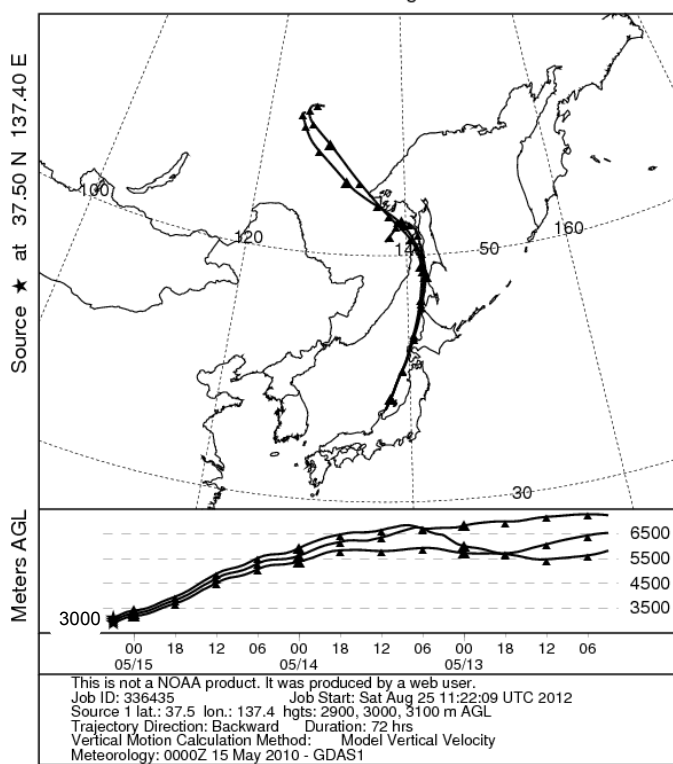


Fig. 5 T.Maki et al.

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