

Genotypic and phenotypic characterization of the food spoilage bacterium brochothrix thermosphacta

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- 1 Genotypic and phenotypic characterization of the food spoilage bacterium *Brochothrix*
- 2 thermosphacta
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27 Abstract (200 words)

Microbial food spoilage is responsible for significant economic losses. Brochothrix 28 thermosphacta is one of the major bacteria involved in the spoilage of meat and seafood. Its 29 30 growth and metabolic activities during food storage result in the production of metabolites associated with off-odors. In this study, we evaluated the genotypic and phenotypic diversity 31 32 of this species. A collection of 161 B. thermosphacta strains isolated from different foods, spoiled or not, and from a slaughterhouse environment was constituted from various 33 laboratory collections and completed with new isolates. A PCR test based on the rpoB gene 34 was developed for a fast screening of *B. thermosphacta* isolates. Strains were typed by 35 36 MALDI-TOF MS, rep-PCR, and PFGE. Each typing method separated strains into distinct groups, revealing significant intra-species diversity. These classifications did not correlate 37 with the ecological origin of strains. The ability to produce acetoin and diacetyl, two 38 molecules associated with B. thermosphacta spoilage, was evaluated in meat and shrimp 39 juices. The production level was variable between strains and the spoilage ability on meat or 40 shrimp juice did not correlate with the substrate origin of strains. Although the B. 41 thermosphacta species encompasses ubiquitous strains, spoiling ability is both strain- and 42 environment-dependent. 43

44 Keywords

45 Meat; spoilage; seafood; off-odors; diversity; acetoin; diacetyl; *rpoB* species-specific PCR

- 46 Highlights
- There is significant diversity between *B. thermosphacta* strains
- Diversity is not related to the ecological origin of the isolates
- The ability to produce acetoin and diacetyl depends on strains and food matrices
- 50

51

52 1. Introduction

Brochothrix thermosphacta is recognized as the dominant food spoiler of meat and seafood 53 products stored under modified atmosphere packaging (Remenant et al., 2015). This 54 ubiquitous microorganism has been isolated from foods of animal origin such as meat, 55 seafood and dairy products (Stackebrandt and Jones, 2006). Moreover, it has been described 56 as widely disseminated along the food chain, from the raw material to the final product, as 57 well as in the food processing environment (Nychas et al., 2008; Stackebrandt and Jones, 58 2006). B. thermosphacta can cause serious economic losses in the food industry due to its 59 60 ability to produce metabolites associated with off-odors. For example in beef meat, it has been shown to produce cheesy and creamy dairy off-odors associated with the production of 3-61 hydroxy-2-butanone (acetoin), 2,3-butanedione (diacetyl), and 3-methyl-1-butanol (Casaburi 62 63 et al., 2014; Dainty and Mackey, 1992). In cold-smoked salmon, B. thermosphacta produces 2-hexanone and 2-heptanone, two compounds responsible for the formation of the blue-64 65 cheese off-odor (Joffraud et al., 2001; Laursen et al., 2006; Mejlholm et al., 2005). Strong butter, buttermilk-like, sour, and nauseous off-odors caused by B. thermosphacta in cooked 66 and peeled shrimp have been associated with the production of 2,3-butanedione (diacetyl), 3-67 methyl-1-butanal, and 3-methyl-1-butanol (Jaffrès et al., 2011; Laursen et al., 2006; Mejlholm 68 et al., 2005). Therefore, the molecules produced by *B. thermosphacta* seem to depend on the 69 70 food matrix. However, the above-mentioned studies used different strains and thus it is possible that the spoilage potential is also strain-dependent. 71

B. thermosphacta is a facultative anaerobe that can grow on chilled meats and fish stored
under low O₂ and under vacuum packaging (Borch et al., 1996; Drosinos and Nychas, 1997;
Ercolini et al., 2006). Although glucose is not present at high concentration in meat, Gill and
Newton (1977) reported that it is the preferred substrate of *B. thermosphacta* when grown in

meat juice. In addition glucose metabolism is greatly affected by the composition of the gas
used for storage. Under aerobic conditions, 3-hydroxy-2-butanone (acetoin) and 2,3butanedione (diacetyl) are the major metabolites produced by the consumption of glucose
while under anaerobic conditions, *B. thermosphacta* produces lactic acid and ethanol (Dainty
et al., 1985; Pin et al., 2002). Thus, spoilage activity may also vary depending on food storage
conditions.

Brochothrix and *Listeria* genera constitute the *Listeriaceae* family, and the *Brochothrix* genus
encompasses two non-pathogenic species: *B. thermosphacta* and *Brochothrix campestris*.
However, little information is available for *B. campestris*, most of the available information
refers mainly to a single strain (ATCC 43754, the type strain) (Gribble and Brightwell, 2013;
Talon et al., 1988).

87 Various molecular techniques have been widely applied to genotype foodborne pathogenic or spoilage bacterial species. Pulsed Field Gel Electrophoresis (PFGE) has been described as 88 89 highly discriminatory, robust and reproducible (Gerner-Smidt et al., 2006; Graves and 90 Swaminathan, 2001; Lukinmaa et al., 2004). It has been used successfully for genotyping B. thermosphacta meat isolates (Papadopoulou et al., 2012). Repetitive-element Palindromic 91 PCR (rep-PCR) has been widely applied for molecular typing and has proven to be a powerful 92 93 tool in environmental and food microbiology (Ishii and Sadowsky, 2009). It can differentiate a wide range of bacterial species at the subspecies or even the strain level (Wolska and 94 95 Szweda, 2012) and has been applied to the differentiation of *B. thermosphacta* (Papadopoulou et al., 2012; Xu et al., 2010). Recently, various studies have also shown the applicability of 96 Matrix Assisted Laser Desorption/Ionization Time Of Flight Mass Spectrometry (MALDI-97 TOF MS) for bacterial identification, taxonomy and strain typing (Singhal et al., 2015). 98 Widely used in clinical microbiology for identification purposes (Carbonnelle et al., 2011), 99 this technique is an effective tool for the intra-specific typing of bacteria from the genera 100

Listeria (Barbuddhe et al., 2008) and *Salmonella* (Dieckmann et al., 2008) and is increasingly
applied for identifying and typing microorganisms associated with food (Böhme et al., 2011;
Kern et al., 2014).

104 Several studies have led to different conclusions about the intra-species diversity of B. thermosphacta (Papadopoulou et al., 2012; Stanborough et al., 2017). This may be due to 105 106 the small number of strains, the low diversity of the ecological origin of the studied 107 collections, or the use of different methods. In addition, the genetic functions involved in spoilage remain little studied. The analysis of draft genome sequences of 13 B. thermosphacta 108 strains pointed out some genes potentially involved in spoilage activity, but did not reveal any 109 110 strong diversity (Stanborough et al., 2017). Therefore, genetic diversity and its potential link with the ecological niches or spoilage ability of *B. thermosphacta* remain unsolved. 111

In order to investigate whether a correlation can be established between the ecological origins of *B. thermosphacta* strains, their diversity and their spoilage potential, we constituted a collection of strains from a wide range of ecological environments. MALDI-TOF MS, PFGE, and rep-PCR were used to assess diversity in the collection while phenotypic diversity was evaluated through the quantification of acetoin and diacetyl production, two molecules associated with spoilage by *B. thermosphacta*.

118 **2.** Materials and methods

119 **2.1.** Bacterial strains and growth conditions.

The 161 *B. thermosphacta* strains isolated during this study (N = 80) or sourced from various
collections (N = 81) are listed in Table 1. In addition, *B. campestris* ATCC 43754 (= DSM
4712), *Listeria innocua* ATCC 33090, *L. innocua* CLIP 11262, *L. monocytogenes* ATCC
35152, *L. monocytogenes* 08-5578 (Gilmour et al., 2010), *Carnobacterium maltaromaticum*ATCC 27865, *Carnobacterium divergens* V41 (Pilet et al., 1995), *Staphylococcus epidermidis*

ATCC 12228, *S. epidermidis* RP62A (Gill et al., 2005), *Serratia liquefaciens* ATCC 27592,
and *Escherichia coli* K12 were used as controls for various purposes. Bacteria were routinely
grown in BHI broth (VWR Chemicals, France) at 25 °C for *Brochothrix* sp., 30 °C for *Listeria* sp., *Carnobacterium* sp., *S. epidermidis*, and *S. liquefaciens*, and at 37 °C for *E. coli*.

Minced beef meat and peeled shrimp juices were used as broth to quantify acetoin and 129 diacetyl production by *B. thermosphacta* strains. Meat juice was prepared by stomaching 130 ground beef, collected frozen from a local supermarket, as previously described by Rantsiou 131 et al. (2012), and filtered through a 0.45 µm membrane filter before sterilization with a 132 0.2 µm membrane filter. Shrimp juice was prepared by crushing frozen raw peeled shrimp 133 from Ecuador (91/100 without sulfite, purchased from industry, Nantes) in sterile distilled 134 water. The shrimp based mixture was heated (100 °C; 2 min), filtered and autoclaved (100 °C; 135 30 min) as previously described by Fall et al. (2010). Five milliliter aliquots were then stored 136 137 frozen at -20 °C in 15 ml tubes until use.

Bacterial enumeration was performed after 48 h of incubation on Plate Count Agar (PCA)
(Biomerieux, France) at 30 °C and *B. thermosphacta* selective STAA agar base containing
STAA selective supplement (Oxoid, France) at 25 °C, to determine the total aerobic and *B. thermosphacta* counts, respectively.

142 **2.2.** Sampling new *B. thermosphacta* isolates

Sampling was carried out in a beef slaughterhouse at five points: (i) the chilling room (walls and floors), (ii) the nacelle receiving the viscera, (iii) the knives used for skinning, (iv) animal skin and (v) cattle barns (floor and walls). About 10 cm² of knife surfaces were sampled by rubbing cotton swabs five times in both vertical and horizontal directions for 30 seconds. Other surfaces (walls, floors, nacelles, skin) were sampled using sterile wipes impregnated with peptone water. Samples were transported from the collection site to the laboratory in a cooler (4 °C) and analyzed immediately. Swab and wipe samples were homogenized by

shaking manually with 10 ml and 25 ml of peptone water (Biokar Diagnostics, France), 150 151 respectively. New isolates from ground beef meat and chicken cuts were also collected in the present study. Chicken cuts were rinsed in peptone water as previously described (Rouger et 152 al., 2017) whereas beef meat was stomached in 0.9% NaCl solution for 3 min. Then, 153 appropriate decimal dilutions were plated on PCA and STAA plates for bacterial enumeration. 154 Three to four colonies were selected from STAA plates, then purified on Brain Heart Infusion 155 (BHI) agar (VWR Chemicals, France) and stored at -80 °C in BHI broth supplemented with 156 20% (v/v) glycerol (VWR Chemicals, France). 157

158 2.3. DNA extraction

DNA was extracted from 2 ml of overnight cultures with the DNeasy blood and tissue kit
(Qiagen, France) according to the manufacturer's instructions. DNA concentration and purity
were estimated after electrophoresis on 1% agarose. DNA extracts were stored at -20 °C.

162 2.4. *rpoB* species-specific primer design and PCR conditions

163 A PCR primer set was designed to amplify a DNA fragment specific to B. thermosphacta, 164 excluding *B. campestris*, other closely related species (such as *Listeria*) and species present in the same environments or reported as growing on STAA medium (such as Carnobacterium, 165 or Staphylococcus sp.). The in silico primer design was based on the multiple alignment of 166 167 rpoB gene sequences available from the GenBank database. The rpoB sequences of B. thermosphacta ATCC 11509, B. campestris ATCC 43754, and the most closely related 168 169 bacterial species were aligned using the BioEdit-ClutalW Sequence Alignment program (Hall, 1999). Specific 170 primers were designed using Primer3 software (http://frodo.wi.mit.edu/primer3/). The in silico specificity of primers was assessed by 171 nucleotide BLAST [National Center for Biotechnology Information (NCBI)] analysis and 172 Primer3. Oligonucleotides with the highest specificity for the *B. thermosphacta* sequence, 173 without hairpin structures or dimers, were selected. This resulted in the design of the forward 174

primer (*rpoB*-Fw1_154-175: 5'- GCGTGCATTAGGTTTCAGTACA-3') and the reverse
primer (*rpoB*-Rev1_525-547: 5'- TCCAAGACCAGACTCTAATTGCT-3') for the specific
amplification of 394 bp of the *B. thermosphacta rpoB* gene. Primer specificity was then
assessed by PCR amplification on the DNA extracted from *B. campestris*, *Listeria* sp, *Carnobacterium* sp, and *Staphylococcus* sp.

Amplifications were performed in a 50 µl reaction volume containing: 1 µl (50-100 ng) of 180 DNA, 5 µl of 10X Taq Buffer (New England Biolabs, France), 0.2 µM of dNTP (New 181 England Biolabs, France), 0.4 µM of each primer and 1.5 U of Taq-polymerase (New England 182 Biolabs, France). PCR reactions were carried out with a PTC-100 Thermocycler (Bio-Rad 183 184 Laboratories, France) using the following amplification conditions: initial denaturation step at 95 °C for 5 min, followed by 25 cycles of [denaturation (95 °C for 30 s), primer annealing 185 (66 °C for 30 s), primer extension (72 °C for 30 sec)], and a final extension step at 72 °C for 186 5 min. Amplicons were separated in a 1.5% (w/v) agarose gel containing 0.05X of Syber Safe 187 (Invitrogen Life Technologies, France) in TAE buffer. The gel was visualized under UV 188 189 transillumination (Bio-Rad Laboratories, France).

190 2.5. 16S rDNA sequencing

The 16S rDNA (about 1500 bp) was amplified by PCR according to Jaffrès et al. (2009). Fragments were partially sequenced (about 800 bp) using the Eurofins Genomics service (Les Ulis, France). The resulting sequences were cleaned then assembled into a unique contig sequence with BioEdit software (Hall, 1999). A BLAST search of partial 16S rRNA gene sequences was performed in the NCBI database (NCBI, Bethesda, USA).

196 **2.6. MALDI-TOF**

For MALDI-TOF MS analysis, fresh cultures of *Brochothrix* sp. incubated at 25 °C were centrifuged (10 min; 3,000 g; 4 °C) and rinsed in 1 ml of molecular biology grade water. One microliter of the bacterial suspension was spotted in a square-form onto the sample target

plate in 8 replicates (Bruker Daltonics, Germany) and allowed to dry in a biosafety cabinet at 200 ambient temperature. Each droplet was overlaid with 1 µl of HCCA matrix solution, a 201 saturated solution of alpha-cyano-4-hydroxy cinnamic acid in 50% acetonitrile with 2.5% 202 trifluoroacetic acid (Bruker Daltonics, Germany), and then dried as above. As a positive 203 control for each run and for calibration purposes, one spot was also covered by the Bacterial 204 Test Standard (BTS) mixture (Bruker Daltonics, Germany). Measurements were made using 205 the manufacturer's recommended settings (linear positive mode, Nitrogen Laser with 60 Hz 206 207 repetition rate, 20-kV acceleration voltage, 18.5-kV IS2 voltage, 250 ns extraction delay, and 2,000 to 20,000 m/z range). For each sample, mass spectra were examined visually using 208 FlexAnalysis (Bruker Daltonics V3.4) to identify large spot-to-spot inconsistent variations. 209 Spectra were then imported into a Matlab (MathWorks) script, which performs smoothing, 210 normalization, baseline subtraction and peak selection automatically. From a selected peak list 211 212 associated to intra-species variations, a dendrogram was generated using the Euclidean distance measure and an average linkage. 213

214 2.7. Rep-PCR

215 DNA was subjected to rep-PCR analysis according to Ouoba et al. (2008) using (GTG)5 primer (5'-GTGGTGGTGGTGGTG-3'). Amplicons were separated in a 2% (w/v) agarose gel 216 in 1 x TAE at 3 V/cm for 3 h. After the run, gels were stained with 0.5 µl/ml Syber Safe 217 (Invitrogen Life Technologies, France) for 1 h and then visualized with UV transillumination 218 (Bio-Rad Laboratories, France). DNA profiles were analyzed with Bionumerics software, 219 version 6.5 (Applied-Maths, Belgium). Isolates were compared using the band-based Dice 220 coefficient (optimization: 0.5%; tolerance: 1%) and UPGMA (unweighted pair-group method 221 using the average approach) cluster analysis. 222

223 **2.8. PFGE**

Genomic DNA from 2 ml of overnight cultures was prepared in low-melting-point agarose 224 225 plugs as described by Doulgeraki et al. (2010), and digested with the endonuclease ApaI (New England Biolabs, France) according to the manufacturer's instructions. Electrophoresis was 226 227 performed on the CHEF-DRIII PFGE system (Bio-Rad Laboratories, France) in 1% (w/v) agarose gels with 0.5 x TBE as the running buffer, at 14 °C. A lambda ladder (Bio-Rad 228 Laboratories, France) was used as the molecular weight marker. Restriction fragments were 229 resolved at a constant voltage of 6 V/cm with switch times of 4-40 s for 18 h and 4-12 s for 230 4 h. Gels were stained with 0.5 mg/ml ethidium bromide, and DNA bands were visualized 231 with UV transillumination (Bio-Rad Laboratories, France). PFGE profiles were analyzed 232 using the BioNumerics Software, version 6.5 (Applied-Maths, Belgium), and then compared 233 using the Pearson coefficient (optimization: 0.5%; curve smoothing: 0%) and UPGMA cluster 234 analysis. 235

236 **2.9. Acetoin/diacetyl production**

237 2.9.1. Voges-Proskauer reaction

Tests were performed in 48-well plates (Falcon, France) on 0.5 ml samples pipetted from 10 ml cultures collected after 48 h of growth at 25 °C and gentle resuspension of cells. A volume of 75 μ l of alpha-naphthol (5% (w/v) in 95% ethanol) and 50 μ l of KOH [40% (w/v) in water] was added. Plates were incubated at room temperature for 1 h. The level of acetoin production was assessed visually using a six-point scale based on the color intensity and noted from (0): not produced or light yellow to (5): very high production or strong red color.

244 2.9.2. Acetoin and diacetyl quantification

Five milliliter aliquots of meat or shrimp juice were gently defrosted at 4 °C and then inoculated (1:100) with overnight cultures grown at 25 °C in BHI broth. After incubation for 48 h at 25 °C, 2 ml aliquots were centrifuged for 10 min at 10,000 g and the supernatant was recovered for acetoin and diacetyl quantification as described by (Nicholson, 2008;

Westerfeld, 1945). The reaction was carried out on 200 µl of culture supernatant by the 249 addition of 140 µl of creatine [0.5% (w/v) in water], 200 µl of alpha-naphthol [5% (w/v) in 250 95% ethanol], and 200 µl of KOH [40% (w/v) in water]. A blank was prepared 251 252 simultaneously with non-inoculated meat and shrimp juices. Absorbance at 560 nm of the samples was measured in a spectrophotometer (Spectronic Genesys 5) after incubation at 253 room temperature for 10 min (for diacetyl) and 1 h (for acetoin). Standard curves were 254 constructed with controls containing increasing acetoin and diacetyl concentrations and used 255 256 to calculate acetoin and diacetyl production. pH was measured at the end of the experiments using a Crison pH-meter (Crison micro pH 2000, Spain). Analyses were performed in 257 triplicate. 258

259 **2.10. Statistical analysis**

260 2.10.1. Hierarchical clustering and multidimensional scaling

Similarities between isolates were first converted into dissimilarities by complement to 1 of the Dice coefficient (for rep-PCR) and the Pearson coefficient (for PFGE). These dissimilarities were analyzed using two different statistical techniques: the hierarchical clustering of isolates and the factorial representation of individuals by Multidimensional Scaling (MDS).

Dendrograms of isolates were obtained by hierarchical clustering of the two matrices of dissimilarities. The agglomerative procedure was the UPGMA, also known as the group average linkage (Everitt et al., 2001).

Multidimensional scaling aims to derive a factorial representation of individuals from a measure of the dissimilarity between them (Borg and Groenen, 2005). The two matrices of dissimilarities between isolates were summed and submitted to MDS in order to produce a spatial configuration representing the distances between isolates. The quality of fit was measured by the stress index, which is the sum of the squared differences between the initial

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dissimilarities and distances in the configuration. The obtained stress value was 0.147 with 6

dimensions, indicating an acceptable goodness of fit of the MDS configuration (Krzanowski,

1990). The final configuration was rotated in order to interpret the results more easily.

277 Data were statistically analyzed with the R packages *cluster* (for hierarchical clustering) and

278 *smacof* (for multidimensional scaling).

279 **2.10.2.** Analysis of variance

Analysis of variance (ANOVA) with R version 3.3.2 (C) 2016 (The R Foundation for Statistical Computing) was performed to determine statistically significant differences between strains grown in meat and shrimp juices. P-values < 0.05 were considered statistically significant.

284 **3. Results**

285 **3.1.** Constitution of a *B. thermosphacta* strain collection

As the purpose of the study was to investigate the genotypic and phenotypic diversity of the 286 species and to determine whether a correlation exists between genotype and ecological origin 287 or spoilage potential, we wanted to constitute as diverse a strain collection as possible. First, 288 79 isolates from spoiled or non-spoiled meat, seafood and milk products were provided by 289 different laboratories. We included DSM 20599 and the type strain ATCC 11509 (= DSM 290 20171), both isolated from pork meat. To complete the collection, chicken cuts and a beef 291 slaughterhouse environment were sampled as no such isolate was represented in the available 292 293 collection. New isolates from non-spoiled ground beef meat were also added for comparison with those provided by other laboratories. More than 200 new isolates were collected and 80 294 were kept for further analysis after removal of putative redundant strains, using a preliminary 295 296 rep-PCR analysis. In total, a collection of 161 B. thermosphacta strains was selected for analysis (Table 1). 297

298 **3.2.** *rpoB* species-specific PCR test

As B. campestris isolates are scarce (only 5 strains have been reported in the literature 299 (Illikoud et al., in press) and references therein), we developed an accurate and reproducible 300 PCR assay for a fast identification of *B. thermosphacta* isolates since both species cannot be 301 302 discriminated through their 16S rDNA sequence. The PCR assay was designed to target a 394 bp region of the *B. thermosphacta rpoB* gene. The specificity of the primer set (*rpoB*-303 Fw1 154-175/rpoB-Rev1 525-547) was tested against a range of DNA from closely related 304 bacterial species, such as B. campestris, and some Listeria or lactic acid bacteria species 305 306 (Fig. 1A).

307 Amplification was only observed from *B. thermosphacta* DNA, showing the specificity of the 308 assay. The 80 new isolates were then tested by this species-specific *rpoB*-PCR assay and all 309 were identified as *B. thermosphacta*.

310 **3.3.** Characterization of genotypic the intra-species diversity

The diversity among the 161 *B. thermosphacta* strains was assessed by MALDI-TOF MS, rep-PCR, and PFGE typing methods. The *B. campestris* type strain ATCC 43754 was included as a control.

314 MALDI-TOF MS spectra were obtained for all strains. Cluster analysis was performed on all the spectra to visualize similarities between those of different strains. This generated a 315 dendrogram composed of 14 groups, named from A to N (Supplementary Fig. S1). Two 316 317 groups (M and N) encompassed about 87% of the collection (N =113 and 26 strains, respectively) with other groups containing only 1 to 4 strains. Strains from different 318 ecological origins were distributed in all groups. Most groups included strains isolated from 319 different ecological origins while groups G, H, J, and L encompassed only 2-4 chicken cut 320 isolates. B. thermosphacta TAP 107 and TAP 104 from group G came from the same sample, 321 thus we cannot exclude that these two isolates are redundant. Conversely, B. thermosphacta 322 TAP 204 and TAP 206 from group H were isolated from different batches. B. campestris 323

ATCC 43754 was the only member of group B. By exploiting only the intra-species spectral variations, typing with MALDI-TOF did not report accuracy the phylogenetic distances between the two species of *Brochothrix* sp. For this reason MALDI-TOF MS did not clearly differentiate *B. thermosphacta* from *B. campestris* since another *B. thermosphacta* isolate (MFPA43A14-06) stayed as an outgroup (Supplementary Fig. S1).

The electrophoresis profiles of rep-PCR products yielded 3 to 11 bands, depending on strains. 329 330 The amplification products were mainly in the range of 0.5 to 3 kb and a quite large diversity between profiles was observed. The UPGMA clustering analysis of the rep-PCR profiles 331 obtained for all strains produced the dendrogram shown in Supplementary Fig. S2. 332 333 B. campestris ATCC 43754 was clearly located in a separate external cluster whereas all B. thermosphacta strains formed a single cluster. By applying a 60% similarity coefficient, 334 the dendrogram analysis generated 12 groups (named A to L) in the B. thermosphacta cluster. 335 336 Each group was composed of at least two isolates from different ecological origins and different laboratory collections, except for group L, which included two isolates (TAP 105 337 and TAP 199) from different chicken cut batches. Some isolates, all from sea bream and the 338 same collection (such as FMCC B-116, FMCC B-118, and FMCC B-119 from group I) had 339 very similar profiles and may be redundant. This was again observed for TAP 107 and 340 TAP 104 in group C. However, these two chicken meat isolates also harbored a profile very 341 close to that of CRE 2333, EBP 3069, EBP 3070, SF 677, and SF 711, all from different 342 seafood products and collected from different laboratories. Using a similarity coefficient of 343 344 80% was not more informative as it generated 50 groups, 20 of which were composed of a single strain. 345

By PFGE, using *Apa*I as the restriction enzyme, *B. thermosphacta* CD 355, MIP 2622, TAP 57, TAP 63 and TAP 78 showed only one band or could not be lysed. These isolates were therefore excluded from the analysis. The UPGMA cluster analysis of the PFGE profiles

obtained for the remaining strains resulted in the dendrogram shown in Supplementary Fig. 349 350 S3. The dendrogram analysis, applying a 35% similarity coefficient, generated 18 groups (A to R) with only 2 single-strain groups (i.e. D and R). Except for these two groups and group 351 352 M composed of two strains isolated from cooked and peeled shrimp, each group comprised at least two strains from different ecological origins. In addition, these groups comprised strains 353 from at least two different laboratory collections. With a 60% similarity coefficient, 75 groups 354 were differentiated including 34 groups consisting of single strains. The dendrogram analysis 355 356 with an 80% similarity coefficient generated 130 groups including 108 single-strain groups. PFGE did not differentiate B. campestris ATCC 43754 from B. thermosphacta strains. In fact, 357 this strain appeared in group I with B. thermosphacta FMCC B-116, FMCC B-118, M1 and 358 VHB2. 359

Multidimensional scaling (MDS), obtained by summing the two matrices of dissimilarities from the rep-PCR and PFGE analyses, produced a spatial configuration representing the distances between isolates (Fig. 2). It revealed significant intra-species diversity within the strain collection. MDS also clearly illustrated the absence of ecotype in *B. thermosphacta*: the intra-species diversity was not related to the ecological origin of the strains as isolates from the various origins were widely distributed and no strain clustering associated with a particular environment was observed.

367 3.4. Acetoin and diacetyl production

All strains were first screened to estimate their ability to produce acetoin in a laboratory medium using the Voges-Proskauer reaction. Levels of acetoin and diacetyl were estimated using a six-point scale, based on the color intensity, noted from (0): no production to (5): very high production. Negative (*E. coli* K12, which does not produce acetoin) and positive (acetoin-producing *S. liquefaciens* ATCC 27592) controls were included (Fig. 3A). The production level varied between strains. For example, 12% of the strains similar to the negative control and noted (0) did not produce acetoin, while only 2% produced very high
levels (noted (5), far above the positive control) (Fig. 3B).

Thirteen strains representing the 6 classes mentioned above (low to high producers) were 376 377 chosen for further analysis to measure acetoin/diacetyl production after growth in meat and shrimp juices. Diverse strains were selected, taking into account the ecological and 378 geographical origins and the diversity assessed by MALDI-TOF MS, rep-PCR and PFGE. 379 The B. thermosphacta and B. campestris type strains and B. thermosphacta DSM 20599 were 380 also included. All tested strains produced both molecules in meat and shrimp juices, and the 381 production level was both strain- and food matrix-dependent (Fig. 4). For all strains, acetoin 382 and diacetyl were produced at higher concentrations in meat juice than in shrimp juice. In 383 meat juice, the production level ranged from $26.87 \pm 8.56 \,\mu$ g/ml to $129.06 \pm 41.32 \,\mu$ g/ml for 384 diacetyl, and from $51.65 \pm 6.32 \,\mu$ g/ml to $111.48 \pm 2.84 \,\mu$ g/ml for acetoin. B. thermosphacta 385 386 EBP 3070, a strain isolated from a spoiled fish product, was the highest producer in shrimp juice, with 22.86 \pm 3.19 µg/ml acetoin and 30.24 \pm 2.78 µg/ml diacetyl. In shrimp juice, the 387 388 lowest production levels were observed with the B. thermosphacta type strain (5.30 389 \pm 1.01 µg/ml acetoin and 5.41 \pm 1.67 µg/ml diacetyl) and *B. thermosphacta* BSAS1 3 (7.05) $\pm 1.20 \,\mu$ g/ml acetoin and 7.94 $\pm 1.71 \,\mu$ g/ml diacetyl), which was isolated from the 390 environment. B. thermosphacta EBP 3033, EBP 3070, and TAP 175 were among the higher 391 392 producers of acetoin and diacetyl in shrimp juice and lower producers in meat juice. Conversely, CD 337 and BSAS1 3 were high producers of acetoin and diacetyl in meat juice 393 and lower producers in shrimp juice. Finally, B. campestris ATCC 43754 was among the 394 lowest acetoin and diacetyl producer, whatever the juice used. The quantitative data obtained 395 396 on shrimp juice correlated with those of the screening test performed in BHI medium. Indeed, 397 the highest producers EBP3033, EBP 3070, and TAP 175 (Fig. 4) were noted (5), (5), and (4), respectively using the six-point scale based color intensity of the preliminary test. On the 398

opposite, the lowest producers ATCC 11509 and BSAS1 3 were noted (2) and (1),respectively.

Bacterial enumeration showed that the bacterial population reached at the end of the experiment was higher in shrimp juice than in meat juice. Counts varied according to strains from $6.06 \pm 0.05 \log \text{CFU/ml}$ to $6.72 \pm 0.20 \log \text{CFU/ml}$ in meat juice and from 7.75 ± 0.11 log CFU/ml to $8.31 \pm 0.12 \log \text{CFU/ml}$ in shrimp juice. After 48 h of incubation, the pH of non-inoculated meat and shrimp juices was 5.54 ± 0.19 and 6.40 ± 0.16 , respectively. For all strains, the final pH reached 5.03 ± 0.062 and 5.66 ± 0.104 in meat and shrimp juices, respectively.

408 **4. Discussion**

The diversity of the microbial populations involved in the spoilage of meat and seafood 409 products has been widely studied and documented (Dainty and Mackey, 1992; Jaffrès et al., 410 411 2011; Koutsoumanis and Nychas, 1999; Nychas et al., 2008; Remenant et al., 2015). These studies have shown that *B. thermosphacta* plays an important role in the spoilage of these 412 products. Although the diversity of its spoilage potential depending on the strains has been 413 414 described, it remains poorly understood (Casaburi et al., 2014). The aims of our study were to evaluate the intra-species diversity in a large and diverse collection of *B. thermosphacta* 415 isolates and to investigate whether this diversity could be correlated with their ecological 416 origin and/or their ability to produce spoilage compounds. 417

Half of the collection was constituted of new isolates collected for this study. Since the two
closely related species *B. campestris* and *B. thermosphacta* cannot be distinguished through
their 16S rDNA sequence, we developed a PCR assay based on the *rpoB* gene to identify *B. thermosphacta* accurately among the new isolates. The *rpoB* housekeeping gene, encoding
the RNA polymerase beta-subunit, has been described as a useful and relevant target for
bacterial identification and phylogenetic studies (Adékambi et al., 2009; Case et al., 2007;

Mollet et al., 1997). The specificity of this PCR assay was confirmed against a range of DNA from closely related bacterial species, such as lactic acid bacteria and *Listeria* species. This enabled the differentiation of both *Brochothrix* species and the identification of new isolates as *B. thermosphacta*. In fact, of 207 chicken cut isolates, 206 were positive to the *rpoB* PCR test, and the other one was identified as *Enterococcus faecalis* by partial 16S rDNA sequencing. Similarly, 11 out of 12 ground beef new isolates were identified as *B. thermosphacta* by this PCR assay (Fig. 1B).

The collection included 161 B. thermosphacta isolates from the environment and various food 431 432 matrices of animal origin (cheese; chicken, pork, beef, horse, and lamb meats; cod, salmon, 433 sea bream, and shrimp). Some were isolated from spoiled food and others were collected from unspoiled products and before the use-by-date. MALDI-TOF MS, rep-PCR, and PFGE typing 434 methods were selected to investigate the diversity in the collection because they are based on 435 different principles. Rep-PCR provides fingerprints related to the presence of small repeats 436 within the genome (Ishii and Sadowsky, 2009) while electrophoretic PFGE profiles are 437 438 generated by the separation of DNA fragments after genomic DNA digestion with rare-cutting restriction enzymes (Li et al., 2009). MALDI-TOF generates peptide mass fingerprinting of 439 proteins, mainly ribosomal ones as they are the most abundant and are constitutively 440 441 synthesized (Rahi et al., 2016).

PFGE and rep-PCR revealed a significant diversity between *B. thermosphacta* isolates. Analysis of rep-PCR profiles by applying 60% or 80% similarity coefficients generated 12 and 50 groups, respectively. With PFGE dendrograms, these similarity coefficients distinguished 75 and 130 groups. Our results thus revealed a greater diversity than reported in previous studies on *B. thermosphacta*. For example, a PFGE analysis performed on 302 *B. thermosphacta* pork isolates distinguished only 8 groups (Papadopoulou et al., 2012), while rep-PCR used on 27 *B. thermosphacta* isolates from meat, poultry and seafood reported only

minor differences between isolates (Xu et al., 2010). The unique pork ecological origin in the 449 450 first study and the small number of isolates in the second one most probably explain the low diversity reported by these authors. Compared to the observations made with the two DNA-451 452 based methods, MALDI-TOF MS typing showed a lower diversity as a major cluster encompassed more than 70% of the strains we tested. MALDI-TOF MS has been shown to 453 discriminate L. monocytogenes (Barbuddhe et al., 2008; Ojima-Kato et al., 2016), 454 Lactobacillus brevis (Kern et al., 2014), and E. coli (Siegrist et al., 2007) strains successfully. 455 However, this method was not suitable for differentiating the two subspecies Lactococcus 456 lactis subsp. cremoris and L. lactis subsp. lactis (Tanigawa et al., 2010). The same failure was 457 reported for species belonging to the genera Bacillus and Pseudomonas (Ghyselinck et al., 458 2011), suggesting that the resolution of MALDI-TOF MS is taxon-dependent (Ghyselinck et 459 al., 2011). To our knowledge, MALDI-TOF MS has been successfully used for identifying 460 461 B. thermosphacta isolates from food (Höll et al., 2016), but never for investigating diversity within this species. Moreover, the available databases contain a limited number of spectra for 462 463 some bacterial species.

Whatever the method used, we observed that the different ecological origins (meat, milk, 464 seafood, or slaughterhouse environment) were widely distributed in all groups. In addition, 465 466 most groups encompassing a reasonable number of strains (more than 2) included isolates from different ecological and geographical origins. Furthermore, most of the groups contained 467 isolates from spoiled and non-spoiled food and from both processed and unprocessed 468 products. This suggests that the strains belonging to the different groups may have a common 469 contamination pattern in the various meat and seafood products. These strains have adapted to 470 471 grow in chilled meat and seafood products. We also noted that the groups comprised isolates recovered from products of different meat animal species as well as from the processing 472 473 environment. This probably reflects the physiological capability of *B. thermosphacta* strains

to grow in various food matrix ecosystems independently of the product type (raw or 474 475 processed), the animal species from which it is derived (beef, pork, salmon, shrimp, etc.), and the packaging conditions (under air, vacuum packaged, modified atmosphere). In other words, 476 no ecotype (strains sharing the same ecological niche) was observed. Of the 80 new isolates, 477 for which information about batch origin was available, a few clustered together 478 systematically, whatever the typing method used. Five pairs of such isolates (TAP 57/TAP 63, 479 TAP 104/TAP 107, TAP 143/TAP 147, TAP 176/TAP 180, and TAP 204/TAP 206) were 480 noticed and were collected from the same batches. Nevertheless, TAP 104, TAP 107, 481 TAP 176, TAP 180, TAP 204, and TAP 206 showed differing acetoin production ability in 482 the Voges-Proskauer test and were therefore not considered redundant. 483

Whether the spoiling potential of *B. thermosphacta* is strain-dependent is unknown. To 484 evaluate this, we focused on acetoin and diacetyl production since both molecules have 485 already been reported as associated with the spoilage of beef and chicken meat and seafood 486 products (Casaburi et al., 2014; Franke and Beauchamp, 2017; Jaffrès et al., 2011). We first 487 488 screened the ability to produce acetoin using the Voges-Proskauer reaction from glucose fermentation in BHI laboratory medium. Our results showed that the production of this 489 molecule was highly variable between strains and did not correlate with their origin (spoiled 490 491 or non-spoiled products, nature of the food) or their genotypic clustering.

The combination of rep-PCR, MALDI-TOF and PFGE clustering and Voges-Proskauer reaction data was used to select a sub-set of thirteen distant strains for acetoin and diacetyl quantification. For this, sterile juices from two food matrices, beef meat and shrimp, were used in this study in order to avoid interference with the endogenous microbiota. Both acetoin and diacetyl were produced by *B. thermosphacta* in both matrices. These results differ from previous studies that reported no acetoin but only diacetyl production in cooked and peeled shrimp packed under modified atmosphere (Jaffrès et al., 2011; Laursen et al., 2006).

Conversely, Casaburi et al. (2014) reported that B. thermosphacta produced acetoin but not 499 diacetyl in beef meat. These apparent contradictions may result from different methods and 500 experimental conditions (shrimp packed under modified atmosphere vs. shrimp juice; pieces 501 502 of beef meat stored aerobically vs. beef juice). Acetoin and diacetyl production was higher in meat juice than in shrimp juice although the bacterial population reached after 48 h of 503 incubation was higher in shrimp juice than in meat juice. These observations could suggest 504 that the production of these molecules is not related to the growth level of *B. thermosphacta*, 505 506 but more probably to the biochemical composition of the food matrix. Moreover, acetoin and diacetyl production levels varied between strains. We noticed that B. thermosphacta 507 508 EBP 3070 (isolated from spoiled salmon) and TAP 175 (isolated from non-spoiled chicken cuts) were among the highest producers of acetoin and diacetyl in shrimp juice but among the 509 lowest ones in meat juice. Conversely, B. thermosphacta CD 337 (isolated from spoiled 510 511 shrimp) and BSAS1 3 (isolated from the slaughterhouse environment) belonged to the highest acetoin and diacetyl producers in meat juice and to the lowest in shrimp juice. This shows that 512 513 the spoilage ability of B. thermosphacta was both strain- and matrix-dependent but was not 514 correlated to the food from which environment strains were isolated. This might result from the regulation of genes involved in the metabolic pathways that produce these molecules, 515 which may vary depending on strains. This was recently suggested by the comparison of 13 516 517 B. thermosphacta draft genomes, which highlighted a large number of transcriptional regulators in these genomes but a small difference between strains (Stanborough et al., 2017). 518 However, the genomes were all sequenced from strains of meat origin and may not represent 519 the diversity of the *B. thermosphacta* species. 520

521 **5.** Conclusion

522 The present study revealed a significant diversity within the strain collection using rep-PCR,523 PFGE and MALDI-TOF typing methods. All these methods showed that there was no ecotype

in this *B. thermosphacta* strain collection. The ability to produce acetoin and diacetyl in meat and shrimp juices varied between strains and did not correlate with the isolation from a spoiled or non-spoiled food product, suggesting that the spoiling ability of *B. thermosphacta* was most probably linked to strain properties rather than to the food environment from which they were isolated. Based on these results four strains were selected for genome sequence comparison, and transcriptomics coupled to volatilome analysis will be performed for better understanding of the *B. thermosphacta* spoilage mechanisms.

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539

540 Figure legends

- 541 Fig. 1. Specificity of the specific PCR assay
- A: Performance of the *B. thermosphacta*-specific PCR assay with DNA obtained from
 representative *Brochothrix* species and closely related bacteria.
- **B:** New isolates from chicken cuts (TAP 125) and beef meat (VHB2, VHB3, VHU1, VHU2,
- 545 VHU3, VHF DLC2 1, VHF DLC2 2, VHF DLC2 3, VHF DLC1 1, VHF DLC1 2, and

546 VHF DLC1 3) were identified as *B. thermosphacta*. Strains from external laboratories:

547 B. thermosphacta ATCC 11509, B. thermosphacta DSM 20599, B. campestris ATCC 43754,

- 548 CD 251, M6, FMCC B-112, and SF 750 were included. VHB1 isolate was negative and thus
- 549 considered as not belonging to the *B. thermosphacta* species.
- 550 A 100 bp ladder (New England Biolabs, France) and a negative (-) control were included.
- 551 Fig. 2. Multidimensional scaling of summed rep-PCR and PFGE distance matrices.
- Each strain is presented with a color code illustrating its ecological origin: (■) beef meat, (■)
 pork meat, (■) lamb meat, (■) horse meat, (■) beef + lamb meat, (■) chicken meat, (■)
 shrimp, (■) salmon, (■) sea bream, (■) cod fillet, (■) cheese, and (■) slaughterhouse
- 555 environment.
- **Fig. 3.** Acetoin production using the Voges-Proskauer test.

557 The level of acetoin production was visually estimated using a six-point scale based on the 558 color intensity and noted from: (0): not produced (light yellow color) to (5): very high 559 production (strong red color)

A: Example of the results of the acetoin production assay for *B. thermosphacta* strains, *S. liquefaciens* (positive control), and *E. coli* (negative control).

562 **B:** Pie-chart representing the overall results for 161 *B. thermosphacta* strains

Fig. 4. Acetoin and diacetyl production by *Brochothrix* sp. strains in meat and shrimp juices. Fifteen *B. thermosphacta* strains and *B. campestris* ATCC 43754 were tested for their ability to produce acetoin and diacetyl after 48 h of culture in meat and shrimp juices. Data are expressed as the mean \pm SD of three biological replicates.

567 Supplementary material

Figure S1. Cluster analysis of the MALDI-TOF MS of 161 *B. thermosphacta* isolates listed in Table 1. The type strain *B. campestris* ATCC 43754 was included as a control. The color code refers to the ecological origin of the strains: (**•**) beef meat, (**•**) pork meat, (**•**) lamb meat, (**•**) horse meat, (**•**) beef + lamb meat, (**•**) chicken meat, (**•**) shrimp, (**•**) salmon, (**•**) sea bream, (**•**) cod fillet, (**•**) cheese, and (**•**) slaughterhouse environment. The distance level indicates the similarity of spectra ranging from 0 (identical spectra) to 1 (maximum variability).

Figure S2. Dendrogram obtained by the cluster analysis of rep-PCR fingerprints of the 161 575 B. thermosphacta isolates listed in Table 1. The type strain B. campestris ATCC 43754 was 576 included as a control. Similarities were calculated using the Dice coefficient and the data 577 clustered using the UPGMA. The color code refers to the ecological origin of the strains: (=) 578 579 beef meat, (\blacksquare) pork meat, (\blacksquare) lamb meat, (\blacksquare) horse meat, (\blacksquare) beef + lamb meat, (\blacksquare) chicken meat, (\blacksquare) shrimp, (\blacksquare) salmon, (\blacksquare) sea bream, (\blacksquare) cod fillet, (\blacksquare) cheese, and (\blacksquare) 580 slaughterhouse environment. The vertical blue dashed line shows the delineation level of 581 60%. Groups from A to L were formed at the similarity level of 60%. 582

Figure S3. Dendrogram obtained by the cluster analysis of PFGE fingerprints of the 161 *B. thermosphacta* isolates listed in Table 1. The type strain *B. campestris* ATCC 43754 was
included as a control. Similarities were calculated using the Pearson coefficient and the data

- clustered using the UPGMA. The color code refers to the ecological origin of the strains: (=)
- 587 beef meat, (\blacksquare) pork meat, (\blacksquare) lamb meat, (\blacksquare) horse meat, (\blacksquare) beef + lamb meat, (\blacksquare) chicken
- 588 meat, (\blacksquare) shrimp, (\blacksquare) salmon, (\blacksquare) sea bream, (\blacksquare) cod fillet, (\blacksquare) cheese, and (\blacksquare)
- slaughterhouse environment. The vertical blue dashed line shows the delineation level of
- 590 35%. Groups from A to R were formed at the similarity level of 35%.

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(19%)



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Table 1 : B. thermosphacta isolates used in this study

| Ecological origin (a) | Strains | Laboratory collection | Reference |
|--|--|--|--|
| Beef slaughterhouse environment (7) | BSAS1 1, BSAS1 3, BSAS2 4, BSBS1 3, BSBS1 6, BSAS2 3, BSK1 3 | INRA-SECALIM | This study |
| Chicken legs (48) | TAP 54, TAP 56, TAP 57, TAP 58, TAP 61, TAP 62, TAP 63, TAP 64, TAP 68, TAP 69, TAP 73, TAP 74, TAP 76, TAP 78, TAP 81, TAP 104, TAP 108, TAP 109, TAP 110, TAP 111, TAP 123, TAP 125, TAP 129, TAP 105, TAP 107, TAP 126, TAP 142, TAP 143, TAP 144, TAP 146, TAP 147, TAP 148, TAP 164, TAP 166, TAP 168, TAP 169, TAP 170, TAP 171, TAP 172, TAP 175, TAP 176, TAP 180, TAP 199, TAP 201, TAP 202, TAP 203, TAP 204, TAP 206, TAP 207 | INRA-SECALIM | This study |
| Beef meat | VHB2, VHB3, VHU1, VHU2, VHU3, VHF DLC1 1, VHF DLC1 2, VHF DLC1 3, VHF DLC2 1, VHF DLC2 2, VHF DLC2 3 | INRA-SECALIM | This study |
| (25) | V2, G8, G6, G7, MFPA17A17-02, MFPA19A15-05, MFPA22A14-04, MFPA22A14-05, MFPA42A14-07, MFPA43A14-06, MFPB17A13-02, MFPB42A12-05, MFPB43D06-02, MFPB43A12-01 | | (Lucquin et al., 2012) |
| Lamb meat (1) | 8727 | INRA-MICALIS/FME | |
| Horse meat | 160X7, 160X8 | | |
| Beef and lamb sausages (4) | M1, M2, M4, M6 | | |
| Pork meat (10) | DSM 20171T = ATCC 11509, DSM 20599 | DSM/ATCC | (McLean and Sulzbacher, 1953; Sneath and Jones, 1976) |
| | FMCC B-427, FMCC B-428, FMCC B-429, FMCC B-430, FMCC B-431, FMCC B-432, FMCC B-433, FMCC B-434 | LFMB/ Agricultural University of Athens | |
| Shrimps (20) | CD 251, CD 252, CD 266, CD 274, CD 280, CD 290, CD 321, CD 322, CD 326, CD 331, CD 337, CD 340, CD 350, CD 352, CD 355, CD 357, CD 358, CD 372, CRE 2330, CRE 2333 | INRA-SECALIM / IFREMER-EM3B | (Jaffrès et al., 2009) |
| Cod fillet (4) | EBP 3017, EBP 3018, EBP 3032, EBP 3033 | IFREMER-EM3B | |
| Salmon (30) | EBP 3069, EBP 3070, EBP 3083, EBP 3084, SF 677, SF 678, SF 711, SF 712, SF 713, SF 746, SF 748, SF 750, SF 779, SF 781, SF 782, SF 1173, SF 1186, SF 1216, SF 1234, SF 1820, SF 1838, SF 1849, SF 1926, SF 1930, SF 1939, MIP 2440, MIP 2490, MIP 2576, MIP 2599, MIP 2622 | INRA-SECALIM / IFREMER-EM3B | (Chaillou et al., 2015) |
| Sea bream (8) | FMCC B-112, FMCC B-113, FMCC B-114, FMCC B-115, FMCC B-116, FMCC B-117, FMCC B-118, FMCC B-119 | LFMB/ Agricultural University of Athens | |
| Cheese rind (1) | cH8.14 | INRA-URF | (Almeida et al., 2014) |
| Unknown (1) | 5X10003 | INRA-MICALIS/FME | |

IFREMER-EM³B: French Research Institute for Exploitation of the Sea, Nantes, France INRA-MICALIS/FME: UMR INRA/AgroParisTech (Microbiologie de l'alimentation au service de la santé), Jouy en Josas, Paris, France INRA-SECALIM: UMR INRA/Oniris (Sécurité des Aliments et Microbiologie), Nantes, France INRA-URF: Unité de Recherches Fromagères, INRA Aurillac, France LFMB: Laboratory of Food Microbiology and Biotechnology, Agricultural University of Athens, Greece