

1 Gene expression plasticity and frontloading promote
2 thermotolerance in Pocilloporid corals

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4 Running title: *Pocillopora* holobiont response to heat stress
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35 **ABSTRACT**

36 Ecosystems worldwide are suffering from climate change. For example, the coral reef ecosystems are
37 globally threatened by increasing sea surface temperatures. However gene expression plasticity provides
38 the potential for organisms to respond rapidly and effectively to environmental change, and would be
39 favored in variable environments. In this study we investigated the thermal stress response of
40 Pocilloporid corals from two contrasting thermal regimes (more stable seawater temperatures in New
41 Caledonia, more variable in Oman) by exposing them to heat stress. We compared the physiological state,
42 bacterial and *Symbiodinium* communities (using 16S and ITS2 metabarcoding), and gene expression levels
43 (using RNA-Seq) between control conditions and heat stress (the temperature just below the first signs of
44 compromised health). Colonies from both thermal regimes remained apparently normal and had open and
45 colored polyps during heat stress, with no change in bacterial and *Symbiodinium* community composition.
46 In contrast, they differed in their transcriptomic responses. The thermotolerant colonies (Oman)
47 presented a more plastic transcriptome (more differentially expressed genes and higher fold-changes
48 between control and heat stress conditions), but some other genes had a higher basal expression level
49 (frontloading) compared to the thermosensitive colonies (New Caledonia). In terms of biological function
50 we observed a trade-off between stress response (including induction of tumor necrosis factor receptors,
51 heat shock proteins, and detoxification of reactive oxygen species) and morpho-anatomical functions.
52 Gene regulation (transcription factors, mobile elements) appeared to be highly enriched, indicating
53 possible epigenetic regulation. These results provide new insights into the balance between plasticity and
54 frontloading, and the origin and evolution of these strategies.

55 **KEYWORDS:** Gene expression plasticity, Frontloading, Coral holobiont, *Pocillopora*, Heat stress.

56

57 INTRODUCTION

58 Earth is undergoing unprecedented global environmental changes with major effects on biodiversity
59 (Barnosky *et al.* 2011). The ongoing erosion of the most vulnerable ecosystems due to current
60 environmental degradation is particularly worrying and is only a premise to what scientists have called
61 the sixth mass extinction (Barnosky *et al.* 2011). In particular, climate change, ocean acidification and the
62 exceptional frequency to which extreme climatic events occur have already resulted in the irreversible
63 degradation of more than 20% of coral reefs worldwide (Bellwood *et al.* 2004; Hoegh-Guldberg *et al.*
64 2007). Scleractinian corals constitute the biological and physical framework for a large diversity of marine
65 organisms [c.a. ~600 coral, ~2000 fish, and ~5000 mollusk species (Veron & Stafford-Smith 2000; Reaka-
66 Kudla 2005). Hence, the extinction or even major decrease of corals would have dramatic repercussions
67 on the overall associated community (Hughes *et al.* 2017a). Natural variation in thermal tolerance exists
68 among coral populations (Oliver & Palumbi 2010; Palumbi *et al.* 2014), especially along a latitudinal
69 gradient (Polato *et al.* 2010; Dixon *et al.* 2015), hence providing some hope for coral survival based on
70 their capacity to acclimatize and/or adapt. More specifically, populations inhabiting in zones with more
71 variable temperature regimes display better tolerance to heat stress. This pattern can be verified from
72 local scale (Kenkel *et al.* 2013; Palumbi *et al.* 2014) to geographical scale (Hughes *et al.* 2003; Riegl *et al.*
73 2011; Coles & Riegl 2013)

74 Understanding the evolutionary processes underlying coral thermo-tolerance at the host level is
75 crucial to better predict the fate of coral populations in response to climate change. In particular, it
76 remains unclear whether thermo-tolerance is acquired via acclimatization (i.e. intra-generational gene
77 expression plasticity; (Barnosky *et al.* 2011; Kenkel & Matz 2016) and/or through Darwinian adaptation
78 (i.e. inter-generational microevolution; (Barnosky *et al.* 2011; Dixon *et al.* 2015). Under the former
79 hypothesis one should expect that the present-day sensitive coral populations could potentially acquire
80 tolerance to heat stress along with the ongoing climate change. According to the latter hypothesis, the
81 persistence of initially thermo-sensitive coral populations would depend on the emergence of better
82 adapted lineages through microevolution and/or on the genetic rescue via gene flow from populations
83 already adapted to heat stress (Bellwood *et al.* 2004; Dixon *et al.* 2015). Actually, some studies strongly

84 suggest that both processes (i.e. acclimatization and adaptation) are likely to co-occur in wild coral
85 populations (Hoegh-Guldberg *et al.* 2007; Reusch 2013; Palumbi *et al.* 2014).

86 The recent advent of high throughput molecular methods have made it possible to go beyond the
87 distinction between the evolutionary processes and focus further on the molecular mechanisms
88 underlying coral response to heat stress. In particular, recent studies clearly demonstrated that coral
89 responses to heat stress involve the fine-tuned regulation of expression levels of some genes/proteins
90 involved in several molecular pathways such as metabolism, stress-response and apoptosis (Veron &
91 Stafford-Smith 2000; Brown *et al.* 2002; Weis 2008; Ainsworth *et al.* 2011; Bellantuono *et al.* 2012a;
92 Barshis *et al.* 2013; Palumbi *et al.* 2014; Kenkel & Matz 2016). In this regard, two main molecular patterns
93 having different temporalities have been put forward. One would be “transcriptional plasticity”, i.e.
94 extensive changes in gene expression levels according to the occurring thermal condition and the other is
95 coined “transcriptional frontloading”, i.e. the elevation of stress related genes baseline expression that
96 precondition organisms to subsequent (recurrent) stresses (Reaka-Kudla 2005; Mayfield *et al.* 2011;
97 Barshis *et al.* 2013; Palumbi *et al.* 2014; Hughes *et al.* 2017a). While such elevated constitutive gene
98 expression levels could reflect local adaptation (i.e. genetically fixed gene expression level; (Oliver &
99 Palumbi 2010; Palumbi *et al.* 2014), it could also reflect an acclimation process via stable epigenetic
100 constitutive gene expression regulation. Surprisingly, frontloading and gene expression plasticity were
101 generally discussed as mutually exclusive patterns (Barshis *et al.* 2013; Dixon *et al.* 2015; Kenkel & Matz
102 2016) although these two molecular processes most likely co-occur during coral responses to heat stress.
103 In particular, one might expect that the regulation strategy of genes (plasticity versus frontloading) will
104 greatly depend on the molecular pathways in which they are involved and the energetic, physiological,
105 and ultimately fitness cost associated with gene expression. So far, frontloading has been detected for
106 stress response genes such as Heat Shock Proteins (HSPs), apoptosis and tumour suppression factors in
107 resilient coral populations under experimentally simulated heat stress inducing bleaching in the common
108 reef-building coral *Acropora hyacinthus* (Polato *et al.* 2010; Barshis *et al.* 2013) and for metabolic genes in
109 populations pre-exposed to warm temperatures in response to long-term heat stress in *Porites astreoides*
110 (Kenkel *et al.* 2013; Palumbi *et al.* 2014). Conversely, in the latter species, plasticity was observed in the
111 expression of environmental stress response genes (Riegl *et al.* 2011; Kenkel & Matz 2016), hence

112 challenging the patterns observed in *A. hyacinthus* (Barshis *et al.* 2013; Coles & Riegl 2013). Although both
113 strategies (i.e. constitutive frontloading versus expression plasticity) undoubtedly exists in wild coral
114 populations, the pre-exposure conditions that foster their induction and their relative effects on coral
115 resistance to heat stress still remain unclear (but see (Hughes *et al.* 2003; Kenkel & Matz 2016)).

116 Importantly, scleractinian corals are composed of several symbiotic organisms including the
117 cnidarian host, the mutualist photosynthetic algae (genus *Symbiodinium*) and bacterial communities. All
118 partners (bionts) involved in a stable symbiosis effectively form the entire organism, and constitute what
119 is referred to as the holobiont (Margulis & Fester 1991). A decade after this term was defined, its use has
120 been particularly popularized in reference to corals (Rohwer *et al.* 2002), and subsequent research has led
121 to the hologenome theory of evolution (Rosenberg *et al.* 2007; Zilber-Rosenberg & Rosenberg 2008). In
122 this context the hologenome is defined as the sum of the genetic information of the host and its symbiotic
123 microorganisms. Phenotypes are thus the product of the collective genomes of the holobiont partners in
124 interaction with the environment, which constitute the unit of biological organization and thus the object
125 of natural selection (Zilber-Rosenberg & Rosenberg 2008; Guerrero *et al.* 2013; McFall-Ngai *et al.* 2013;
126 Bordenstein & Theis 2015; Theis *et al.* 2016). Additionally to the cnidarian host response, the genotype -or
127 association of genotypes- of the photosynthetic mutualist *Symbiodinium* symbionts plays a key role in the
128 thermotolerance of the holobiont (Mayfield *et al.* 2014; Hume *et al.* 2015; Suggett *et al.* 2017; Mayfield *et al.*
129 *et al.* 2018). There is less certainty about the importance of the coral bacterial community in participating to
130 the fitness of the holobiont, although accruing evidences strongly suggest their implication in coral
131 response to environmental conditions (Li *et al.* 2014; Pantos *et al.* 2015; Hernandez-Agreda *et al.* 2016)
132 and in the resistance to disease (Sato *et al.* 2009; Cróquer *et al.* 2013; Meyer *et al.* 2015). Finally, the role
133 of the coral-associated microbes and their potential to modify holobiont adaptability remain so far
134 overlooked (but see (Ziegler *et al.* 2017; Torda *et al.* 2017)). Hence studying how corals respond to stress
135 implies an integrative approach to analyze the response of each symbiotic protagonist.

136 The aim of the present study was to investigate the molecular mechanisms underlying thermo-
137 tolerance of coral holobionts. We analyzed the holobiont response to stress of two coral populations
138 originating from environments with contrasting thermal regimes. We used the scleractinian corals from
139 the genus *Pocillopora* as model species because they have a broad spatial distribution throughout the

140 Indo-Pacific (Veron & Stafford-Smith 2000). These coral species are considered to be one of the most
141 environmentally sensitive (van Woesik *et al.* 2011) but their widespread distribution clearly suggests
142 potential for acclimatization and/or adaptation which may be correlated to specific differences (i.e.
143 different cryptic lineages may be adapted to different environmental conditions). In particular, we focused
144 on *Pocillopora* colonies from two localities with contrasting thermal regimes: colonies from New
145 Caledonia (NC) are exposed to temperate and stable temperatures over the year, while those from Oman
146 are exposed to globally warmer and more seasonal fluctuating temperatures. In order to interpret the
147 results in a precise evolutionary context we first identified the mitochondrial lineage of the colonies. To
148 avoid biases inherent in transplantation-based field experiments resulting from environmental factors
149 other than temperature, we undertook our comparative study in a controlled environment in which we
150 mimicked ecologically realistic heat stress to compare the responses of colonies from both localities. We
151 combined a specific RNA-seq approach to study the cnidarians host response, and metabarcoding analyses
152 using ITS-2 and 16S amplicon sequencing to study the dynamics of the associated algal (*Symbiodinium* sp.)
153 and microbial community compositions, respectively. According to the literature we first expected to
154 detect changes in both symbiotic algal and bacterial communities in corals from both localities when
155 exposed to heat stress. Moreover, since variable environments are expected to promote the evolution of
156 plasticity, we predicted that the cnidarian hosts from Oman will display more gene expression plasticity
157 than those from New Caledonia. However, because frontloading was also found to be an alternative
158 response to recurrent changing conditions, we might also expect some degrees of constitutive high levels
159 of gene expression at least for some molecular pathways and more particularly in Oman corals.

160 MATERIAL AND METHODS

161 CORAL SAMPLING AND MAINTENANCE

162 *Pocillopora damicornis*-like colonies originating from environments characterized by contrasting thermal
163 regimes were sampled in two different localities: (1) in Oman (Om; June 2014), where corals are exposed
164 to a globally warmer and variable thermal environment, and (2) in New Caledonia (NC; November 2014),
165 where corals are subject to more mitigate and stable temperatures (see Table 1 for the temperature
166 regime of locality). At this aim, we sampled colonies from the two geographical locations occupying the
167 same apparent morphological and water depth niche. To account for possible intra-population diversity,

168 three colonies (20 cm in diameter) were collected in each locality, and separated by at least 10 m to avoid
169 the collection of members of the same clone or clonal lineage, as some *Pocillopora* species are able to
170 propagate by asexual reproduction (Adjeroud *et al.* 2013). Immediately following collection, a 1 cm tip of
171 each colony was excised, rinsed three times in filtered seawater (0.22 μm), and placed in RNAlater
172 solution (Sigma Aldrich) for the *in situ* microbiota analysis. The remainder of the colony was fragmented
173 into 20 branches each of 10 cm length, and these were individually placed in plastic bags containing
174 oxygenated seawater, and transported by aircraft to the research aquarium of the Banyuls-sur-Mer
175 oceanographic observatory (France). The coral branches were maintained in artificial seawater (Seachem
176 Reef Salt) at 26°C, and supplied daily with *Artemia* nauplii to satisfy their heterotrophic demand. The
177 conditions in the maintenance tank were controlled to mimic the natural physicochemical parameters of
178 coral reefs (pH: 8.2; salinity: 36 g/L; light Intensity: 150 to 250 photons/m²/s; photoperiod: 12h
179 night/12h day; kH: 6–7.5 dkH; calcium concentration: 410–450 mg/L; inorganic phosphate concentration:
180 < 0.1 mg/L; magnesium concentration: 1300–1400 mg/L; nitrate concentration: < 5 mg/L). After two
181 months of acclimation in the lab, the coral branches were fragmented to produce several clones from each
182 colony (~3 cm). These were individually fixed to a support (here a p1000 tip) using an epoxy adhesive.
183 We waited for complete healing (evident as tissue extending to cover the epoxy adhesive) prior to run the
184 experiment.

185 ECOLOGICALLY REALISTIC HEAT STRESS

186 The aim of this experiment was to compare the response to heat stress of colonies from two localities
187 having the same physiological state, to investigate the patterns of expression of the molecular pathways
188 involved during the stress exposure and the putative modifications of the coral microbiota.

189 The experimental design comprised eight tanks of 53 L (four per locality) in which the seawater was
190 continuously recycled. The water was sterilized using UV (rate 3200 L/h) and renewed twice per hour in
191 each tank (recirculation rate: 100 L/h in each tank). The eight tanks shared the same seawater but their
192 temperature was monitored individually (HOBBY BiothermPro, model 10892; 500W AquaMedic titanium
193 heater; HOBO TidbiT v2 logger). For each population, at least four nubbins per mother colonies were
194 randomly placed in each tank (four tanks per locality) for two weeks at 26°C and the following protocol
195 was applied: three tanks were subjected to a gradual temperature increase (stress treatment) while the

196 fourth (control) was maintained at the control temperature. These controls were used to assess that the
197 stress observed in the stressful treatment was not due to other potential confounding effects (Fig. 1). Both
198 the control and stress temperatures were specific for each sampling locality to mimic their respective
199 natural environment. In particular, we set the control temperature as the mean water temperature for the
200 three warmer months measured at the coral sampling site locality (Table 1): 31°C for the colonies from
201 Om, and 27°C for the colonies from NC. The stress treatment was ecologically realistic, i.e. reflecting a
202 naturally occurring warming anomaly, and consisted in increasing the temperature gradually by 1°C (over
203 5 consecutive hours) each week until physiological collapse of the corals became evident (polyps closure,
204 bleaching or necrosis), as described by (Vidal-Dupiol *et al.* 2009). Sampling was performed each week
205 before the temperature increase and were chosen *a posteriori* for subsequent genetic and transcriptomic
206 analyses. This corresponded to those sampled in each tank just before the first increase of temperature
207 (control samples), and just before the temperature that produced the first signs of physiological collapse
208 (stress temperature samples). Thus, for each condition (control and stress) we obtained three replicates
209 of each colony (three colonies per locality) to reach a total of 36 samples (2 localities × 3 colonies × 2
210 experimental conditions × 3 replicates). The general health of the nubbins was assessed via daily
211 photographic monitoring (at noon prior to feeding) throughout the period of the experiment.

212 DNA EXTRACTION

213 DNA was extracted from each 36 samples as well as coral tips directly collected on the six colonies *in*
214 *natura* for the *in situ* condition (three in Oman, three in New Caledonia), using the DNeasy Blood and
215 Tissue kit (Qiagen) following the manufacturer's instructions. DNA was quantified by spectrophotometry
216 (NanoDrop).

217 HOST HAPLOTYPE

218 As the *corallum* macro-morphology is not a diagnostic criterion in *Pocillopora* genus, the host species was
219 thus identified molecularly. We analyzed the mitochondrial variable open reading frame (ORF) sequence
220 of each colony using the FATP6.1 and RORF primers and following the protocol detailed in (Flot & Tillier
221 2007). The PCR products were sequenced using Sanger sequencing, and the obtained sequences were
222 aligned with the ORF haplotypes from Gélin *et al.* (2017a) using Geneious 8.0 (Kearse *et al.*, 2012) with the
223 MAFFT algorithm (Katoh *et al.* 2005). Each colony used in the experiment was then assigned to Primary

224 Species Hypothesis (PSH) following the nomenclature from G elin *et al.* (2017a). Furthermore, to refine the
225 assignment with the ORF to Secondary Species Hypothesis (SSH) and to identify whether the sampled
226 colonies are members of a same clone, they were genotyped using 13 specific microsatellites as in G elin *et al.*
227 *et al.* (2017a). If relevant, the colonies were assigned to SSH. Meanwhile, the identical multi-locus genotypes
228 were identified as in G elin *et al.* (2017b).

229 MICROBIAL COMMUNITY ANALYSIS USING MISEQ 16S AND ITS2 230 METABARCODING

231 The aim of this analysis was to investigate the composition and the dynamics of the two principal
232 symbiotic coral communities (i.e. bacterial and algal) *in situ* and during heat stress.

233 234 AMPLICON SEQUENCING

235 A bacterial 16S rDNA amplicon library was generated for each of the 42 samples (one *in situ* condition,
236 three control conditions and three stress conditions per colony, three colonies per locality, two localities),
237 using the 341F (CCTACGGGNGGCWGCAG) and 805R (GACTACHVGGGTATCTAATCC) primers, which target
238 the variable V3/V4 loops (Klindworth *et al.* 2012). The *Symbiodinium* assemblages were analyzed using
239 ITS2 (internal transcribed spacer of the ribosomal RNA gene) amplicon libraries and specific primers
240 targeting a sequence of approximately 350 bp (ITS2-F GTGAATTGCAGAACTCCGTG; ITS2-R
241 CCTCCGCTTACTTATATGCTT) (Lajeunesse & Trench 2000; Quigley *et al.* 2014). For both markers, paired-
242 end sequencing using a 250 bp read length was performed on the MiSeq system (Illumina) using the v2
243 chemistry, according to the manufacturer's protocol at the Centre d'Innovation G enome Qu ebec and
244 McGill University, Montreal, Canada.

245 BIOINFORMATIC ANALYSIS:

246 The FROGS pipeline (Find Rapidly OTU with Galaxy Solution) implemented on a Galaxy platform
247 (<http://sigenae-workbench.toulouse.inra.fr/galaxy/>) was used for data processing (Escudi  *et al.* 2015).
248 In brief, paired reads were merged using FLASH (Mago  & Salzberg 2011). After cleaning and removal of
249 primer/adapters using cutadapt (Martin 2011), *de novo* clustering was performed using SWARM (Mah  *et al.*
250 *et al.* 2014). This uses a local clustering threshold with an aggregation distance (d) of 3. Chimeras were
251 removed using VSEARCH (Rognes *et al.* 2016). We filtered the dataset for singletons and performed
252 affiliation using Blast+ against the Silva database (release 128, September 2016) for 16S amplicons

253 (Altschul *et al.* 1990). For ITS2 metabarcoding, the *Symbiodinium* type was assessed using Blast+ against
254 an in-house database of *Symbiodinium* reference sequences built from sequences publicly available. An
255 OTU table in standard BIOM format with taxonomic affiliation was produced for subsequent analyses.

256 For community composition analysis we used the *phyloseq* R package (McMurdie & Holmes 2013) to infer
257 alpha diversity metrics at the OTU level, and beta diversity (between sample similarity) from the OTU
258 table. Community similarity was assessed by Principal Coordinate Analysis (PCoA) using the Bray-Curtis
259 distance matrices.

260 We performed one-way ANOVAs to compare alpha and beta diversity metrics among the groups of
261 samples by sampling locality or by treatment. Corrections based on multiple testing were performed using
262 FDR (Benjamini & Hochberg 1995). For all analyses, the threshold significance level was set at 0.05.

263 TRANSCRIPTOME ANALYSIS

264 The aim of this analysis was to study the transcriptomes of the sampled colonies in response to heat stress
265 compared with controlled conditions.

266 RNA EXTRACTION

267 Total RNA was extracted from each coral sample using TRIzol reagent (Invitrogen), according to the
268 manufacturer's protocol. The quantity and integrity of the total RNA extracted was checked using an
269 Agilent 2100 Bioanalyzer (Agilent Technologies) (mean RIN = 7.5). Paired-end fragment libraries
270 (2 × 100 bp) were constructed and sequenced on an Illumina HiSeq 2000 platform at the Centre
271 d'Innovation Génome Québec at McGill University, Montreal, Canada.

272 BIOINFORMATIC ANALYSES

273 Fastq read files were processed on the Galaxy instance of the IHPE (<http://bioinfo.univ-perp.fr>) (Giardine
274 *et al.* 2005). Quality control and initial cleaning of the reads were performed using the filter by quality
275 program (version 1.0.0) based on the FASTX-toolkit (Blankenberg *et al.* 2010). Reads having fewer than
276 90% of bases having a Phred quality score ≤ 26 were discarded (probability of 2.5 incorrect base call out
277 of 1000, and a base call accuracy of 99.75%). Adaptors used for sequencing were removed using the
278 cutadapt program version 1.6 (Martin 2011). All paired-end reads were aligned using RNASTAR software
279 under default parameters, with at least 66% of the bases being required to align to the reference, and a
280 maximum of ten mismatches per read (Dobin *et al.* 2013). The *Pocillopora damicornis* reference genome

281 used in this study (manuscript in preparation) consisted of a draft assembly of 25,553 contigs (352 Mb
282 total) and N50 = 171,375 bp. The resulting transcriptome served as the reference for reads mapping, and
283 a GTF annotation file was constructed using cufflink/cuffmerge (Trapnell *et al.* 2010). HTseq was used to
284 produce count files for genes (Anders *et al.* 2015). The DESeq2 package was used to estimate the
285 normalized abundances of the transcripts, and to calculate differential gene expression for samples
286 between the control temperature and the stress temperature for each locality (Love *et al.* 2014),
287 considering the different genotypes (3 biological replicates for each genotype) and using default
288 parameters. We next analyzed genes according to their expression patterns among localities and
289 temperature treatments. Genes were clustered manually into six groups according to their differential
290 expression levels: common over-expressed genes, NC-specific over-expressed genes, Om-specific over-
291 expressed genes, common under-expressed genes, NC-specific under-expressed genes, and Om-specific
292 under-expressed genes. Cluster 3.0 (de Hoon *et al.* 2004) and Treeview (Saldanha 2004) were used to
293 build the heatmap.

294 DISCRIMINANT ANALYSIS OF PRINCIPAL COMPONENTS (DAPC):

295 Our aim was to quantify and compare the level of genome-wide transcriptome plasticity between colonies
296 from Om and NC in response to heat stress. To achieve this we performed a discriminant analysis of
297 principal components (DAPC) based on a log-transformed transcript abundance matrix (containing
298 26,600 genes) obtained from the 36 samples (i.e. 9 control and 9 stressed replicates per locality), as
299 described previously (Kenkel & Matz 2016). Specifically, we ran a DAPC analysis using the resulting log₂
300 transformed dataset for the colonies from NC and Om reared in control conditions as predefined groups in
301 the *adegenet* package implemented in R (Jombart *et al.* 2010). Two principal components and a single
302 discriminant function were retained. We then predicted the position of stressed colonies from both
303 localities (Om and NC) onto the unique discriminant function of the DAPC.

304 We next compared the absolute mean DAPC scores between Om and NC colonies for each experimental
305 group (control versus heat stress) using the non-parametric Wilcoxon test. We also ran a general linear
306 model (GLM) using the DAPC scores as dependent variable, and accounted for the locality of origin (NC
307 and Om), the conditions (control and heat stress), and their interaction as explanatory variables. In
308 particular, as an indicator of significant differences in the genome-wide gene expression reaction norms

309 (i.e. differences in DAPC scores between the control and the heat stress treatments) between Om and NC
310 colonies we tested for significant effects of the interaction between the localities and the condition effects.

311 GO ENRICHMENT OF DIFFERENTIALLY EXPRESSED GENES

312 The transcriptome was annotated *de novo* using a translated nucleotide query (blastx (Altschul *et al.*
313 1990)) against the non-redundant protein sequence database (nr). The 25 best hits were then used to
314 search for gene ontology terms using the Blast2Go program (Conesa *et al.* 2005). Lists of significantly up-
315 regulated and down-regulated genes were subjected to GO enrichment analysis by comparison with all
316 expressed gene using a Fischer exact test and a FDR value of 0.05. We used REVIGO to visualize the
317 enriched biological processes (Supek *et al.* 2011).

318 RESULTS

319 HOST HAPLOTYPE

320 Among the three colonies from New Caledonia, colonies NC2 and NC3 presented haplotype ORF18 and
321 were assigned to Primary Species Hypothesis PSH05 and more precisely to Secondary Species Hypothesis
322 SSH05a (Gelin *et al.* 2017a), corresponding to *P. damicornis* type β (Schmidt-Roach *et al.* 2013) also named
323 *P. acuta* (Schmidt-Roach *et al.* 2013) or type 5a (Pinzon *et al.* 2013), while colony NC1 presented ORF09
324 and was assigned to PSH04, *P. damicornis* type α , *P. damicornis* or type 4a, respectively. As for colonies
325 from Oman, they all presented ORF34 and were assigned to PSH12 (Gelin *et al.* 2017a) or type 7a (Pinzon
326 *et al.* 2013)(Additional Table 4). These three PSHs represent three different species.

327 Furthermore, NC2 and NC3 multi-locus genotypes differed from one allele over 26 and were thus part of
328 the same clonal lineage, i.e. the entity that groups together colonies whose multi-locus genotypes slightly
329 differ due to somatic mutations or scoring errors.

330 ECOLOGICALLY REALISTIC HEAT STRESS

331 Our goal was to ensure that our experimental heat stress faithfully reflects a realistic heat stress *in natura*.
332 Following collection from the field, the corals from the different localities were first maintained in the
333 same controlled conditions prior to the experiment. During this period no mortality or signs of
334 degradation/stress were observed for any of the coral colonies. During the experimental heat stress (i.e.
335 gradual temperature increase), visual and photographic monitoring clearly indicated that the first sign of

336 coral stress (i.e. the closure of polyps) occurred at day 30 for both sampling localities, corresponding to
337 30°C and 34°C for the NC and Om colonies, respectively. These temperatures perfectly match the warmest
338 temperature experienced by these populations in the field (Table 1). No signs of physiological collapse
339 were observed in control corals throughout the experiment indicating that all the other parameters were
340 maintained optimal for coral colonies.

341 BACTERIAL COMMUNITIES

342 Among the overall 42 samples analyzed, a total of 5,308,761 16S rDNA sequences were obtained after
343 cleaning and singleton filtering corresponding to 15,211 OTUs. In all samples the class
344 Gammaproteobacteria was dominant (77.7%), particularly the genus *Endozoicomonas* (44.7% of the
345 sequences); this genus is known to be an endosymbiont of numerous scleractinians (Neave *et al.* 2016b)
346 (See Additional Figure 1 for complete bacterial composition in each colony and replicate). The PCoA of
347 Bray-Curtis distances for all colonies showed no evident clusters based on the experimental treatments
348 (Fig. 2). We observed a loose grouping based on localities and colonies, except for colony NC1, which
349 appeared to have a more specific microbiota composition, as it had a different grouping associated with
350 the first axis, which explained 22.3% of the variance. This could be correlated with the different species
351 hypotheses for NC1 compared to NC2 and NC3 (see above). The one-way ANOVA for alpha diversity
352 (Shannon index) revealed significant differences in the microbiota diversity between localities ($P < 0.05$)
353 and colonies ($P < 0.05$), but no differences among the *in situ* control and stress conditions ($P = 0.885$).
354 Similar results were obtained for the beta-diversity (Bray-Curtis distance) (ANOVA between localities: $P <$
355 0.05 ; between colonies: $P < 0.05$; between conditions: $P = 0.554$; the ANOVA results are provided in
356 Additional Table 2). Thus, the bacterial composition appeared to be relatively specific to each colony
357 within each locality, but remained stable during the transition from the natural environment to artificial
358 seawater, and during heat stress exposure.

359 SYMBIODINIUM ASSEMBLAGES

360 Analysis of the *Symbiodinium* composition was performed based on an ITS2 metabarcoding, which
361 facilitated intra-clade resolution.

362 Removal of those OTUs having an abundance of $< 1\%$ left only 4 OTUs among all samples. Two of these
363 corresponded to type C1, while the other two corresponded to type D1a according to (Baker 2003). Type

364 D1a was highly dominant in the colonies originating from Oman, whereas type C1 was almost exclusive to
365 the corals from New Caledonia (Fig. 3). The *Symbiodinium* community composition was very specific to
366 each locality, but remained stable during the transition from the natural environment to artificial
367 seawater, and during heat stress exposure.

368 HOST TRANSCRIPTOME ANALYSIS

369 We generated 36 transcriptomes corresponding to triplicate samples for three colonies of each locality
370 exposed to the control (C) and stress (S) temperatures.

371 Overall, the transcriptome sequencing of these 36 samples yielded 1,970,889,548 high quality Illumina
372 paired reads of 100 bp. Globally, 40–64% of reads obtained for the Om colonies, and 59–70% of reads
373 obtained for NC colonies successfully mapped to the *Pocillopora damicornis* (type β) reference genome.
374 The apparently better alignment of samples from New Caledonia most likely relies on the fact that the
375 New Caledonia colonies used in this study belong to *P. damicornis* types α or 4a (PSH04) and β or 5a
376 (PSH05), which are phylogenetically close to each other and closer from the reference genome, than the
377 Om colonies from type 7a (PSH12) that is phylogenetically more distant from the reference genome. The
378 aligned reads were assembled in 99,571 unique transcripts (TCONS), representing putative splicing
379 variants of 26,600 genes identified as “XLOC” in the genome (FASTA sequences available in Additional File
380 3).

381 The hierarchical clustering analyses clearly grouped together samples belonging to the same locality and
382 species hypothesis according to their genome-wide gene expression patterns, in link with the
383 phylogenetic differences between the NC and Om phylotypes (Fig. 4). Within locality and species
384 hypothesis, whenever pertinent, the transcriptomes also grouped by colony, indicating that the
385 transcriptomes were genotype-specific. For each colony, the transcriptomes then grouped by condition
386 (temperature treatment) excepted for New Caledonia colonies NC2 and NC3 (corresponding to the same
387 MLL) that clustered together when exposed to controlled and heat stress conditions.

388 In total 5,287 genes were differentially expressed between colonies exposed to stress and control
389 conditions (adjusted $P < 0.05$) in Om colonies, and 1,460 genes in NC colonies (full results of the
390 comparisons between stressed and controls (log₂-foldchange and adjusted p -values) are provided in

391 Additional File 5). The heatmap generated to visualize the expression pattern of each gene in Om and NC
392 colonies is shown in Figure 5.

393 Among these genes, 498 were over-expressed and 350 were under-expressed in colonies exposed to
394 stressful conditions compared to those kept under optimal temperatures (i.e. control) irrespective to the
395 locality of origin (NC or Om). Nevertheless, the differential expression level was significantly higher
396 (Wilcoxon test; $P = 2.2 \times 10^{-16}$) for the Om corals (Fig. 6). Among the 498 over-expressed genes, 358 were
397 more induced in Om corals than the NC ones (log₂-fold change Om > NC; Fig. 6), and of the 350 under-
398 expressed genes, 259 were more repressed in Om corals (Additional Table 6). The mean of the log₂-fold
399 change for common over-expressed genes was 0.9 for Om samples vs. 0.6 for NC samples (variance: 0.6 for
400 Om; 0.2 for NC), and for the common under-expressed genes was -1.2 for Om colonies vs. -0.8 for NC
401 colonies (variance: 1 for Om; 0.3 for NC).

402 Additionally, colonies from the two thermal regimes also responded specifically to heat stress. In
403 particular, 272 genes were over-expressed and 294 were under-expressed only in the NC corals and 2,082
404 were over-expressed and 2,311 were under-expressed only in the Om ones when exposed to heat stress.
405 Finally, the colonies from both localities displayed antagonistic transcriptomic responses to heat stress for
406 a subset of genes: 24 were over-expressed in NC colonies but under-expressed in Om ones, while 22 were
407 under-expressed in NC colonies but over-expressed in the Om ones.

408 Altogether these results revealed a greater transcriptomic response to heat stress in colonies originating
409 from Oman compared to those from New Caledonia (4,393 differentially expressed genes for the Om
410 corals vs. 566 genes for the NC ones).

411 DISCRIMINANT ANALYSIS OF PRINCIPAL COMPONENTS (DAPC):

412 The DAPC analysis clearly discriminated the colonies from both localities based on their overall gene
413 expression patterns (Fig. 7). We also found that Om and NC colonies significantly differed in their gene
414 expression plasticity in response to heat stress (Wilcoxon tests; control p -value = 0.73, stress p -value <
415 0.01). This result was corroborated by the significance of the interaction term between localities and
416 temperature effects in the GLM ($P = 0.04$), which indicate that the slope of the reaction norm was different
417 between localities. More particularly, the Om colonies responded to a greater extent than the NC ones, and
418 thus showed significantly higher gene expression plasticity in response to heat stress.

419 ANALYSIS OF GENE FUNCTION:

420 To investigate the functions associated with the differentially expressed genes we performed a blastx
421 annotation of transcripts followed by a Gene Ontology (GO) term annotation to determine to which
422 protein each gene (XLOC) corresponded most closely, and the biological process, molecular function, and
423 cell compartment localization of the protein (Additional File 7). Differentially expressed genes associated
424 with the control and stress conditions (adjusted p -value < 0.05) clustered into six groups according to
425 their expression patterns in corals from both localities (see host transcriptome analysis section).

426 For the 498 common over-expressed genes, 139 biological processes were enriched compared with the
427 entire set of annotated genes. The most significant biological process identified in the REVIGO analysis (i.e.
428 with lowest FDR value: FDR = 2.1×10^{-68}) was stress response (Fig. 8). Following this sequentially, were
429 cellular metabolism (FDR = 3.7×10^{-49}), positive regulation of biological processes (FDR = 2.4×10^{-43}), cell
430 death (FDR = 2.5×10^{-33}), cellular localization (FDR = 8.4×10^{-25}), and pigment metabolism (FDR = 2.1×10^{-21}). Among the 272 genes over-expressed in the NC but not in the Om colonies in response to heat stress,
432 38 biological processes were enriched and the REVIGO analysis showed that organic acid catabolism (FDR
433 = 1.6×10^{-22}), protein transport (FDR = 1.8×10^{-16}), stress response (FDR = 4.8×10^{-13}), and cellular
434 metabolism (FDR = 3×10^{-12}) were the four most significantly enriched biological processes (Fig. 8). Among
435 the 2,082 genes over-expressed in the Om but not in the NC colonies in response to heat stress, the
436 REVIGO analysis showed that 160 enriched biological processes grouped together, with the most
437 significant being ncRNA metabolism (FDR = 8.9×10^{-303}), cellular metabolism (FDR = 4.4×10^{-70}),
438 carbohydrate derivative biosynthesis (FDR = 5.9×10^{-64}), and organic substance transport (FDR = 2×10^{-44}).

439 For the 350 genes that were under-expressed following heat stress irrespective to the locality of origin
440 (Om or NC), 48 biological processes were enriched compared with the entire set of annotated genes (Fig.
441 8). The REVIGO analysis grouped these genes into five biological processes: nitrogen compound transport
442 (FDR = 5.4×10^{-89}), localization (FDR = 8.1×10^{-10}), regulation of neurotransmitter levels (FDR = 1.2×10^{-8}),
443 system development (FDR = 8.8×10^{-6}), and single organism process (FDR = 4×10^{-4}). Among the under-
444 expressed genes in the NC colonies only, a single biological process (anatomical
445 structure/morphogenesis) was found to be enriched (FDR = 9×10^{-3}). Among the under-expressed genes in
446 the Om colonies only the REVIGO analysis grouped 139 enriched biological processes, with the most

447 significant being ion transmembrane transport (FDR = 7.6×10^{-104}), single multicellular organism process
448 (FDR = 7.5×10^{-53}), regulation of biological quality (FDR = 6×10^{-48}), cell-cell signaling (FDR = 1.5×10^{-23}),
449 single organism process (FDR = 1.1×10^{-18}), multicellular organism process (FDR = 1.5×10^{-16}), biological
450 regulation (FDR = 2.3×10^{-15}), response to abiotic stimulus (FDR = 6.2×10^{-13}), and localization (FDR =
451 4.6×10^{-12}).

452 Regarding other Gene Ontology annotations, the common over-expressed genes were correlated with
453 enhancement of 41 cellular components that were associated in the REVIGO analysis with cellular
454 compartments including the mitochondria (FDR = 1.5×10^{-180}), cells (FDR = 1.5×10^{-19}), organelles (FDR =
455 2.5×10^{-15}), endomembrane systems (FDR = 6.5×10^{-7}), the membrane-enclosed lumen (FDR = 2×10^{-4}), the
456 entire membrane (FDR = 0.0019), the membrane raft (FDR = 0.009), and the membrane (FDR = 9×10^{-3}).

457 The genes over-expressed in NC corals were associated in the REVIGO analysis with 23 cellular
458 compartments, with the mitochondria (FDR = 2.5×10^{-82}), cells (FDR = 4.9×10^{-7}), organelles (FDR =
459 6.9×10^{-6}), and the envelope (FDR = 9×10^{-3}) being the four most significant. The genes over-expressed in
460 Om corals were associated in the REVIGO analysis with 70 cellular components, the most significant of
461 which were the intracellular organelle lumen (FDR = 1×10^{-560}), organelles (FDR = 9.9×10^{-48}), cells ($2.7 \times 10^{-$
462 46), the membrane-enclosed lumen (FDR = 1.2×10^{-19}), macromolecular complexes (FDR = 1×10^{-16}), the
463 endomembrane system (FDR = 2.7×10^{-11}), the envelope (FDR = 3×10^{-9}), and the membrane (FDR = $8 \times 10^{-$
464 4).

465 The common under-expressed genes were associated in the REVIGO analysis with 13 cellular components
466 including the plasma membrane (FDR = 1.1×10^{-44}), membrane (FDR = 2.4×10^{-11}), neurons (FDR = 3×10^{-4}),
467 presynapse (FDR = 4×10^{-4}), and synapse (FDR = 7×10^{-3}) (Fig. 8). The genes under-expressed in NC corals
468 were associated in the REVIGO analysis with seven biological processes in cellular components amongst
469 which the plasma membrane region (FDR = 8.5×10^{-9}), cell periphery (FDR = 1×10^{-3}), the membrane (FDR
470 = 2×10^{-3}), and membrane part (FDR = 9×10^{-3}) were the four most significant. The under-expressed genes
471 in Om corals only were associated in the REVIGO analysis with 26 cellular components, the most
472 significant of which included the integral component of the plasma membrane (FDR = 3.1×10^{-81}), the
473 periphery (FDR = 1.2×10^{-22}), the membrane (FDR = 6.2×10^{-17}), the transmembrane transporter complex

474 (FDR = 1.5×10^{-14}), the extracellular region (FDR = 4.5×10^{-6}), the cell (FDR = 9.5×10^{-6}), synapses (FDR =
475 1.9×10^{-5}), and the synapse part (FDR = 4.4×10^{-5}).

476 To investigate whether the supposed thermotolerant colonies in our study (i.e. the colonies from Oman)
477 displayed a frontloading strategy, as previously described in scleractinian corals (Barshis *et al.* 2013), we
478 compared the constitutive expression levels (i.e. in control conditions) between Om and NC colonies for
479 those genes that were over-expressed in NC colonies (Additional File 8). This analysis showed that the
480 basal expression level was often greater in the more thermotolerant colonies (Om). Among the 770 genes
481 that were over-expressed in NC colonies in response to thermal stress (272 specifically and 498 in
482 common with Om), 484 were constitutively (i.e. in the control condition) more expressed in Om than NC
483 control conditions. Among these genes, 301 were over-expressed in the Om colonies, 20 were under-
484 expressed in the Om colonies, and 163 were not differentially expressed between the control and stress
485 temperature in these colonies, reflecting true frontloading based on the definition of Barshis *et al.* (2013).
486 These three categories of genes (over-expressed, under-expressed, and frontloaded) were submitted to
487 GO term enrichment analysis. No significant results were found for the under-expressed genes. The
488 frontloaded genes were enriched in the biological processes cellular respiration (FDR = 4.4×10^{-23}), cellular
489 component organization (FDR = 0.002), homeostatic process (FDR = 0.005), cellular component
490 organization or biogenesis (FDR = 0.007), cofactor metabolism (FDR = 0.009), and stress response (FDR =
491 0.009), and in the cellular compartments including the mitochondrion (FDR = 1.6×10^{-66}), envelope (FDR =
492 0.0002), cell (FDR = 0.0002), and organelle (FDR = 0.0009). Most interestingly, for genes associated with a
493 higher basal expression level and over-expression in the Om colonies, the most enriched biological
494 processes included stress response (FDR = 1.2×10^{-26}), pigment metabolism (FDR = 5.1×10^{-24}), regulation
495 of phosphate metabolism (FDR = 3.2×10^{-15}), cellular metabolism (FDR = 2.7×10^{-11}), and protein folding
496 (FDR = 7.3×10^{-6}). Among the 43 over-expressed genes involved in the response to stress in the NC
497 colonies, 23 were frontloaded in the Om colonies.

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499
500

501 DISCUSSION

502 PHYLOGENETIC CONTEXT OF ADAPTATION

503 Our aim was to study the evolution of adaptive abilities of coral colonies originating from different
504 thermal regimes. As morphology can be misleading for species identification in scleractinians, we used a
505 molecular approach to test the species relationships of our samples. The analysis of mitochondrial
506 sequences indicated that, despite similar morphologies, our samples corresponded to different
507 phylogroups. This agrees well with previous works showing the importance of cryptic lineages and
508 morphological plasticity in the *Pocillopora* genus (Gelin *et al.* 2017a and references herein). Oman
509 corresponds to species hypothesis PSH12 of (Gelin *et al.* 2017a) which is restricted to the Northwestern
510 Indian Ocean. Regarding the two species hypothesis from NC, PSH05 (type β or 5a) is found in the Indian
511 Ocean, the Pacific Ocean and the Red Sea and PSH04 (type α or 4a) is nearly exclusively found in the
512 Pacific Ocean (very rare in the Indian Ocean, and not found yet in Red Sea) (see Gelin *et al.* 2017a). It
513 would be interesting to study if inside each phylogroup, different thermotolerance mechanisms are
514 present, which would be informative on the evolution or conservation of such strategies. Conversely, the
515 observation of a similar response to thermal stress in two different species in NC could indicate either a
516 conserved strategy or a convergence under the same ecological conditions.

517 AN ECOLOGICALLY REALISTIC HEAT STRESS

518 Here we characterized and compared the response to heat stress of coral colonies collected from wild
519 populations which were exposed to two contrasting thermal regimes. The heat stress applied in this study
520 was ecologically realistic, since the first visual response to (i.e. polyp closure) was observed for all
521 colonies when the gradually increasing experimental temperature reached exactly the upper temperature
522 they are subject to *in natura* (30°C and 34°C for NC and Om corals, respectively). From a biological point of
523 view this first result hence clearly supports that these colonies from two localities and different
524 phylogroups within the genus *Pocillopora* are experiencing to different thermal regimes *in natura* and that
525 they display differential ability to deal with heat stress. Moreover, the accurate control of all other
526 seawater parameters allows us to consider that the holobiont response to the thermal treatment is
527 specific to heat stress and not to other possible confounding effects. Any change in the holobiont is
528 therefore a response to the heat stress and not because of disruption of the coral integrity.

529 SYMBIOTIC COMMUNITY: BACTERIAL AND *SYMBIODINIUM* COMPOSITION

530 For the bacterial community we identified significant differences between localities and
531 genotypes. The microbiota composition of all samples was consistent with previous studies, showing a
532 high proportion of Gammaproteobacteria and dominance of the symbiotic *Endozoicomonas* genus (Bourne
533 & Munn 2005; Neave *et al.* 2016a; Peixoto *et al.* 2017). However, our results clearly demonstrate that
534 experimental heat stress did not induce bacterial community changes in coral colonies irrespective to
535 their locality of origin. For the *Symbiodinium* community the ITS2 metabarcoding analysis enabled intra-
536 clade resolution (Quigley *et al.* 2014). Two distinct types of D1a and C1 clades dominated, representing
537 most of the sequences in the Om and NC corals, respectively. Nine ITS types (A to I) have been identified in
538 the genus *Symbiodinium* (Baker 2003). Some *Symbiodinium* strains strongly participate to the overall
539 holobiont fitness, with type D providing tolerance to higher temperatures (Berkelmans & van Oppen
540 2006) and C1 enhancing coral growth rates (Little *et al.* 2004). Interestingly, we found that the
541 *Symbiodinium* type D1a is dominant in the more thermotolerant Om corals, which is consistent with the
542 results of previous works (Berkelmans & van Oppen 2006).

543 Although the microbial community (both bacterial and *Symbiodinium*) differed between the NC
544 and Om corals, the composition did not change during transition from the field to the artificial seawater
545 conditions, and remained stable during the experimental temperature increase. Thus, the coral holobiont
546 assemblage remained stable over the course of the experiment. Such stability of the microbial community
547 during experimental heat stress was previously observed in the scleractinian *Acropora millepora*
548 (Bellantuono *et al.* 2012b) and *A. tenuis* (Littman *et al.* 2010). Thus, our study conforms to the idea that
549 microbial communities associated with scleractinian corals remain unchanged when the holobionts are
550 exposed to stressful temperatures (but see (Ziegler *et al.* 2017)) but further analyses of gene expression
551 level would be needed to assess their functional responses.

552 HOST TRANSCRIPTOMIC RESPONSE

553 Given the observed stability of the microbial symbiotic community during heat stress, we focused
554 more specifically on the responses attributable to the coral host. We thus compared gene expression
555 patterns at the qualitative and quantitative levels in Om and NC colonies in response to heat stress
556 compared to the control condition. Altogether, our results clearly highlight that the Oman thermotolerant

557 colonies exposed to more variable thermal conditions *in natura* also display, in response to heat stress, a
558 greater plasticity in gene expression levels than the NC thermosensitive colonies. In particular, the
559 transcriptomic response of the Oman colonies involved a larger number of genes and among the set of
560 genes that were differentially expressed similarly in both localities, the absolute difference in expression
561 level of 73% of these genes being significantly higher in the Om colonies. These findings are consistent
562 with the theoretical expectations that a more variable environment promotes the evolution of a greater
563 plasticity (Lande 2009). Accordingly, a recent transplantation study conducted *in natura* also identified
564 greater transcriptomic plasticity in a more thermotolerant (in-shore) population compared with an (off-
565 shore) population inhabiting a more stable thermal habitat in the mustard hill coral *Porites astreoides*
566 (Kenkel & Matz 2016) .

567 Importantly however, we also identified several genes whose expression is constitutively higher
568 in the Om colonies compared to the thermosensitive NC colonies (i.e. comparing the expression levels in
569 the control condition); a process recently called “frontloading” (Barshis *et al.* 2013). Frontloading reflects
570 the preemptive expression of stress-response genes hence predisposing organisms to better respond to
571 stress endured in the past. It has been proposed that the occurrence of plasticity vs. frontloading strategies
572 may depend on the frequency of stresses relative to the typical response time of organisms, with frequent
573 stresses promoting frontloading strategies whereas less frequent perturbations result in an increased
574 plasticity (Kenkel & Matz 2016). Surprisingly, plasticity and frontloading are often discussed as exclusive
575 responses (Barshis *et al.* 2013; Kenkel & Matz 2016). We hypothesized that, rather than being exclusive,
576 plasticity and frontloading often co-occur and might also mainly depend on the regulatory pathways in
577 which genes are involved. In particular, each strategy is expected to be selected for, only if it provides a
578 more efficient response and a better fitness despite possible trade-offs. Our results clearly support that
579 plasticity and frontloading indeed co-occur, specifically in the thermotolerant Om colonies experiencing a
580 more variable thermal environment *in natura*. To tease apart the biological processes that are regulated
581 via plasticity or frontloading in *Pocillopora* response to heat stress, we conducted an enrichment analysis.
582 In the next section we detail the response of coral colonies at the molecular level for each main biological
583 process identified, keeping in mind that congruency between gene expression and protein levels should be
584 cautious (Mayfield *et al.* 2016). RNA-sequencing of eukaryotic poly-adenylated mRNA would allow in

585 principle dual analysis of *Symbiodinium* and coral host transcripts (Mayfield *et al.* 2014), but since our
586 RNA extraction methods resulted in very few, we only focused on the host transcriptomic response.

587 **FUNCTIONAL ANALYSES:**

588 **RESPONSE TO STRESS:**

589 **HSPs:**

590 Heat shock proteins (HSPs) are ubiquitous stress-induced chaperone molecules involved in protein
591 folding and protein damage repair in most organisms (Feder & Hofmann 1999). As such, several HSPs
592 were found to be over-expressed in colonies from both localities, whichever their thermal regime, in
593 response to heat stress. Among these, the over-expression of Hsp70 genes was significantly more intense
594 in the Om colonies compared to NC ones. Hsp70 is one of the most documented protein chaperones in
595 coral heat stress response (Barshis *et al.* 2013; Haguenaer *et al.* 2013), but other forms of HSPs such as
596 Hsp60 (Brown *et al.* 2002), Hsp90 (Carpenter *et al.* 2010), and Hsp40 (Maor-Landaw & Levy 2016) are
597 also known to be involved in the response of these organisms. Accordingly we found that Hsp105, Hsp75,
598 Hsp90, and Hsp71 were over-expressed only in Om corals and Hsp60 and Hsp71 showed higher basal
599 expression levels in the Om corals. We also found that chaperones of the Hsp40 family (also known as
600 DNAj) were also regulated in response to heat stress; three of these were over-expressed in colonies from
601 both localities with a higher basal expression level in the Om corals, whereas five were over-expressed
602 only in the Om corals. These proteins seem to be implicated in protein folding during early heat stress in
603 the scleractinian *Stylophora pistillata*, and are highly expressed prior to bleaching and tissue peeling
604 (Maor-Landaw *et al.* 2014). We also found specific Om over-expression of the pyrexia gene, which codes
605 for a transient receptor-activated cation channel, and had been shown to protect flies from high
606 temperatures (Lee *et al.* 2005). Together these results indicate that most genes involved in the heat shock
607 protein pathways are plastically regulated by corals when exposed to stress with a greater response in
608 colonies inhabiting thermally variable environment. However few HSP genes are also frontloaded in these
609 pre-exposed colonies, in particular Hsp60 which gene is encoded in the mitochondria. This pattern is
610 likely to reflect a more general higher mitochondrial activity in colonies accustomed to more variable
611 and/or warmer temperatures.

612

613 REACTIVE OXYGEN SPECIES (ROS) DETOXIFICATION AND DNA REPAIR:

614 One of the first consequences of heat stress is the production of reactive oxygen species (ROS), which
615 might cause protein, membrane and DNA damages (Kowaltowski & Vercesi 1999; Yakovleva *et al.* 2009;
616 Jena 2012). These oxidative molecules are produced in excess by the symbionts and mitochondria during
617 the photosynthesis and the respiration processes, respectively. Cells produce several detoxification
618 enzymes to limit cellular damages (Weis 2008). In particular, under-expression of the calcium-binding
619 messenger protein calmodulin is a sign of response to oxidative stress (Schallreuter *et al.* 2007; Voolstra
620 *et al.* 2009). This protein had more isoforms that were differentially expressed in response to heat stress
621 in Om compared to NC corals, together with one calcium calmodulin-dependent kinase. The Quinone
622 oxidoreductases proteins are involved in the generation of ROS (Porté *et al.* 2009) and were also
623 specifically under-expressed in Om corals in response to heat stress. Conversely, the thioredoxin gene was
624 over-expressed in colonies from both localities, but with a higher fold change in Om corals. This enzyme
625 detoxifies oxidized molecules and is often implicated in coral response to heat stress (DeSalvo *et al.* 2010;
626 Maor-Landaw *et al.* 2014). Finally, polyamine oxidase is a key component of the oxidative burst in plants,
627 and is involved in the induction of apoptosis (Yoda *et al.* 2006). Two genes, one under-expressed in Om
628 corals and one over-expressed in NC corals, were found to be regulated in our experiment. Overall, the
629 molecular pathway underlying ROS detoxification hence appears to be plastically regulated, with greater
630 plasticity in the thermo-tolerant Om colonies than in the thermo-sensitive NC colonies.

631 Several genes coding for DNA repair enzymes including DNA excision repair, DNA ligase, and DNA
632 polymerase were over-expressed in colonies from both localities when exposed to heat stress. However,
633 among these genes, three showed higher basal expression levels and two were frontloaded in the Om
634 colonies.

635 APOPTOSIS:

636 Among the genes differentially regulated in colonies when exposed to heat stress, many are also involved
637 in apoptosis, a process that has been recurrently associated with coral responses to heat stress
638 (Ainsworth *et al.* 2011; Pratlong *et al.* 2015). Among these, TNFR and TRAF genes code for receptors and
639 receptor-associated components of the tumor necrosis factor (TNF), the latter being central in the
640 apoptosis pathway (Ashkenazi & Dixit 1998). Five TNFR and 11 TRAF genes were specifically over-

641 expressed in the Om stressed colonies, indicating a more intense regulation of the apoptosis pathway in
642 these thermotolerant colonies. Furthermore, four TNFs were found to be specifically under-expressed in
643 the Om corals, whereas TNFAIP3 was specifically over-expressed in response to heat stress. It is worth
644 stressing that the TNFAIP3 gene (TNF alpha-induced protein3) is known to inhibit the NFkB process
645 (inflammatory response) together with apoptosis (Opipari *et al.* 1990). Moreover, Caspase8 and Caspase3
646 were specifically over-expressed in the Om stressed corals. Caspases constitute the effector core of the
647 apoptotic process (Nicholson & Thornberry 1997), and Caspase3 has commonly been implicated in the
648 heat stress response in corals (Maor-Landaw & Levy 2016). This further suggests that the thermotolerant
649 Om colonies are capable of greater regulation of apoptosis than the thermosensitive NC ones when
650 exposed to warm temperature. Another set of 10 TNFR genes and 3 TRAF genes were found to be over-
651 expressed in colonies from both localities, but with a higher expression level in the Om corals both in the
652 control and the stress conditions for 10 of the 13 genes irrespective to the treatment (i.e. stress and
653 control temperature). Fem-1 homolog B, which is involved in the apoptotic process as a death receptor-
654 associated protein, was also constitutively higher expressed in the Om compared to the NC colonies.
655 Together these results indicate that part of the apoptosis pathway is frontloaded in the thermotolerant
656 Om colonies. These results echoes with previous studies showing that TNFR genes are frontloaded in the
657 common reef-building coral *Acropora hyacinthus* (Barshis *et al.* 2013) and hence strengthen the idea that
658 frontloading such biological pathway is an adaptive response to variable and/or warmer temperatures in
659 corals.

660 **ENERGETIC METABOLISM:**

661 **MITOCHONDRIAL FUNCTIONS:**

662 The mitochondria are central to many important biological processes of their cell hosts including energy-
663 generating respiration and can be the source of several oxidative molecules involved in apoptosis. As such,
664 heat stress is expected to promote severe changes in the expression levels of genes involved in the
665 mitochondrial function in all living organisms (Guderley & St-Pierre 2002). Accordingly, our enrichment
666 analysis highlighted several mitochondrial genes (either mitochondria- or nuclear-encoded). Among
667 those, some of them were specifically or more intensively regulated in Om corals, including genes coding
668 for 2-oxoglutarate dehydrogenase (which catalyzes the conversion of 2-oxoglutarate to succinyl-CoA and

669 CO₂ in the Krebs cycle), and several monocarboxylate transporters for lactate or pyruvate, the latter being
670 essential for the regulation of energy metabolism in the thermosensitive sea anemone *Aiptasia* (Halestrap
671 & Meredith 2004; Lehnert *et al.* 2014). These results corroborate with the general higher gene expression
672 plasticity observed in colonies from a population experiencing a more variable thermal regime (Om)
673 compared to colonies from a population exposed to narrower thermal amplitudes (NC).

674 Nevertheless, several other genes that participate in energy metabolism displayed a frontloaded
675 pattern in the Om colonies such as the succinate dehydrogenase [ubiquinone] flavo mitochondrial-like and
676 the citrate synthase genes. Interestingly, the expression of the latter gene is often used as a reliable
677 indication of organisms' aerobic capacity and mitochondrial density (e.g. (Rivest & Hofmann 2014;
678 Hawkins & Warner 2017)). Thus, it is likely that the observed frontloading pattern of these mitochondrial
679 genes (encoded in the mitochondrial genome) reflects a greater mitochondrial density in the Om
680 compared to the NC colonies. According to the literature related to non-phototrophic symbiotic
681 ectotherms such increase in mitochondrial density as an adaptive response to heat stress appears
682 counterintuitive since the opposite pattern is generally observed (Pörtner 2002). However, elevated
683 mitochondrial activity on cnidarian hosts have already been found to be associated with thermal-
684 tolerance in corals and sea anemones (Dixon *et al.* 2015; Hawkins & Warner 2017). Hawkins & Warner
685 (2016) suggest that such specific pattern might be at least partly explained by a decrease in symbiotic
686 algae density resulting from heat stress. Energy metabolism in scleractinian corals is highly dependent of
687 their symbiotic algae, which are the major source of glucose generated by photosynthesis. A partial loss of
688 their symbiotic algal may force cnidarian hosts to increase their energy metabolism, hence requiring an
689 increase in their mitochondrial contents, and to use heterotrophic sources of energy that can be obtained
690 from feeding or from the use of metabolic reserve such as lipids. In this context we also observed a high
691 level of regulation of lipid metabolism in the Om corals. The apolipo A (a constituent of lipoprotein
692 associated with lipid dissolution) and dLp HDL-BGBP precursor (which has lipid transporter activity)
693 genes were specifically over-expressed in the Om corals, and could facilitate the transport and
694 assimilation of lipids.

695

696 SYMBIONT REGULATION:

697 Perturbation of energy metabolism and a decrease in the efficiency of energy production by symbiotic
698 algae can be linked to the modulation of genes associated with symbiont maintenance. In our experiment
699 three Rab-11 isoforms were found to be over-expressed in colonies when exposed to stress, including: one
700 being over-expressed in colonies from both localities but with a higher fold change in the Om corals; one
701 that occurred in NC corals while showing frontloading expression in Om corals; and one induced only in
702 the Om corals. This recycling regulatory protein has been shown to regulate phagosomes containing
703 *Symbiodinium* cells in the *Aiptasia-Symbiodinium* model system (Chen *et al.* 2005). Calumenin was also
704 over-expressed in both Om and NC colonies in response to heat stress, but had a higher fold change in Om
705 corals. This protein is known to be a signal intermediate for sym32, a signaling protein involved in
706 symbiont recognition by the coral host (Reynolds *et al.* 2000; Schwarz & Weis 2003). In addition, three
707 lectins exhibited differential expression in the Om corals: one was over-expressed (with a log₂-fold
708 change of 4.1) and two were under-expressed (log₂-fold change of -2.8 and -2.5). It is worth stressing that
709 lectins are key proteins involved in symbiont recognition (Wood-Charlson *et al.* 2006) and that a
710 breakdown of this recognition may lead to the disruption of the symbiosis (Vidal-Dupiol *et al.* 2009). Thus
711 although we did not expressly quantify the density of *Symbiodinium* algae in colonies during our
712 experiment, our results strongly suggest that *Symbiodinium* were at least partly excluded as a response to
713 heat stress prior to bleaching process. These results hence corroborate with those found relative to a shift
714 in both the greater involvement in energy metabolism and resources use shift.

715 CNIDOCYTES:

716 We observed a higher level of regulation of genes involved in the functioning of cnidocytes (also known as
717 nematocytes) in Om corals. These ectodermal cells are specific to cnidarians, and are involved in
718 environment sensing, defense, and predation. Three cytosolic phospholipase A2 (cPLA2) genes were
719 specifically over-expressed in Om corals. In cnidocytes, the corresponding proteins are involved in venom
720 efficacy by promoting prey lysis (Argiolas & Pisano 1983) (Nevalainen *et al.* 2004). One MAC perforin
721 domain-containing gene had a higher basal expression level in Om corals. This gene encodes for a
722 membrane attack complex/perforin-domain containing protein, and is found in gland cells and
723 nematocytes in the sea anemone *Nematostella*, suggesting a potential role in prey killing (Miller *et al.*
724 2007). Conversely, we found under-expression in Om corals of a gigantoxin homolog, which is a cytolysin

725 (actinoporin) found abundantly in cnidocytes (Hu *et al.* 2011) (Frazão *et al.* 2012). Several polycystic
726 kidney disease gene (PKD1 and 2) isoforms were also found to be under-expressed in the Om corals.
727 These proteins are localized in tentacles in Hydra, and seem to be involved in cnidocyte discharge
728 (McLaughlin 2017). These changes could have been promoted by notch signaling (with over-expression of
729 one notch homolog 2); in cnidarians, this pathway is particularly important for cnidogenesis as shown in
730 *Nematostella vectensis* (Marlow *et al.* 2012). Together these results clearly indicate that the Om colonies
731 are better adapted to switch to heterotrophic nutrition in response to heat stress. Such switch could be
732 linked to higher levels of resistance to heat stress via a decrease in *Symbiodinium* cell numbers, as
733 described in other scleractinian corals (Hughes & Grottoli 2013; Aichelman *et al.* 2016).

734 To summarize, we found that Om colonies displayed a more plastic transcriptomic response for many
735 genes involved in fundamental biological pathways such as stress-response pathways (e.g. HSPs,
736 apoptosis, DNA repair, ROS management) in response to heat stress. Moreover, several genes were found
737 to be frontloaded especially in the apoptosis as well as in the energy metabolic pathway. The latter pattern
738 could be explained by a higher regulation of metabolic genes but could also reflect in part an increase in
739 mitochondrial activity as already documented in other symbiotic cnidarians in response to heat stress
740 (Rivest & Hofmann 2014). Higher plasticity and frontloading are two strategies that are costly. As
741 expected, we found strong evidence for possible trade-offs with other critical life history traits such as
742 growth and reproduction. In the next sections we will discuss the transcriptomic response of both Om and
743 NC colonies in response to stress at molecular pathways associated with central fitness traits.

744 **MORPHO-ANATOMIC PROCESSES AND REPRODUCTION:**

745 **SKELETON AND MUSCLE CELLS:**

746 Some enzymes potentially involved in the calcification and muscle contraction were under-expressed in
747 response to heat stress in Om corals, in particular, a single chain carbonic anhydrase and four carbonic
748 anhydrase genes (1, 2, 7, and 12). Carbonic anhydrases catalyze the interconversion of CO₂ and water to
749 bicarbonate and protons, enabling maintenance of the acid-base balance (Tashian 1989). In scleractinian
750 corals, carbonic anhydrases are also involved in biomineralization processes involved in skeleton
751 formation (Bertucci *et al.* 2013). Two hephaestin genes were also under-expressed following heat stress in
752 Om corals. These copper-dependent ferroxidases were identified in the skeleton of *Acropora millepora*

753 (Vulpe *et al.* 1999), where they can be involved in the incorporation and regulation of iron in the
754 carbonate skeleton (Ramos-Silva *et al.* 2013). Several myosin related proteins similar to myosin light
755 chain kinase or myosin regulatory light polypeptide 9 were found to be specifically under-expressed in the
756 Om corals. Actin-binding LIM partial, potentially involved in Z-disk activity of muscles, and several genes
757 having the LIM domain were also found to be under-expressed in Om corals. In eumetazoans including
758 cnidarians, these proteins are involved in regulation of muscle activity (for a review, see (Leclère &
759 Röttinger 2016). In addition, the genes for the acetyl choline receptor “43 kDa receptor-associated of the
760 synapse-like” and the “calcium-activated potassium channel subunit beta-like” proteins, both of which
761 down-regulate muscle contraction, were over-expressed in Om colonies, which is in accordance with a
762 global decrease in functioning of the contractile apparatus. The observed down-regulation of skeleton
763 formation and muscle contraction strongly suggest that mounting a higher –and most likely better–
764 transcriptomic response to heat stress is associated with a decrease energy allocation to crucial fitness
765 traits and particularly to growth in scleractinian corals.

766 NERVOUS SYSTEM:

767 Many genes involved in nervous system functioning were differentially regulated, with some being over-
768 expressed (e.g. neuronal acetylcholine receptor and VWFA, which regulate the calcium dependent voltage
769 chain), and others being under-expressed, including a proton myo-inositol cotransporter, SCO-spondin
770 (axon guidance), sodium- and chloride-dependent GABA and taurine transporters (neurotransmission),
771 sodium-dependent phosphate transport 2B, and synaptotagmin. Many transcription factors that could be
772 involved in morpho-anatomic integrity, including numerous forkhead box homologs implicated in cell
773 growth, proliferation, and differentiation were over-expressed in Om corals. All these may be involved
774 more globally in the regulation of morpho-anatomic processes (muscle contraction, cnidocytes).

775 CIRCADIAN CLOCK / REPRODUCTION:

776 Several photoreceptors were differentially expressed in the corals. A cryptochrome gene was frontloaded
777 in Om colonies. These flavoproteins are found in plants, animals, insects, and cnidarians, and in corals it
778 has been shown that cryptochrome can synchronize the circadian clock and the reproductive system
779 (Reitzel *et al.* 2013). Three melatonin receptors were also found to be regulated in response to heat stress
780 in Om corals (two over-expressed and one under-expressed). As in vertebrates, in which melatonin is well
781 characterized, in cnidarians it is also involved in the circadian clock (Peres *et al.* 2014). Similarly, two

782 under-expressed and three over-expressed opsin-like genes were identified. The protein is a homolog of
783 the photoreceptor melanopsin, and in *Acropora* is involved in circadian cycle regulation (Vize 2009).

784 Taken together, changes in morpho-anatomic regulation and perturbations in circadian cycles and
785 reproduction could also reflect a trade-off mechanism in the stress response of Om colonies, which is
786 consistent with greater plasticity, and intensive and not fully specific gene regulation activity. We believe
787 that our study contribute to a better comprehension of the biological processes involved in coral
788 thermotolerance. In particular we highlighted the molecular pathways that were regulated either via
789 frontloading or plasticity or both. We also detected clear transcriptomic evidence that allocating energy in
790 heat stress response to better cope with higher temperatures is at the expense of other crucial biological
791 processes such as growth and reproductive functions. However, the molecular mechanisms underlying
792 such global response to heat stress are still partly unresolved. Interestingly we also found specific gene
793 expression patterns linked with epigenetic regulation that could give new insights into our
794 comprehension of such mechanisms and these are exposed and discussed in the next and last section.

795 **EPIGENETIC PROCESSES:**

796 Two histone acetyltransferase genes were over-expressed in Om corals. These enzymes, which
797 are also found in *Nematostella*, are involved in epigenetic gene regulation by modifying the nucleosome
798 structure, and thus the transcription of genes (Sterner & Berger 2000; Karmodiya *et al.* 2014). Several
799 histone-lysine N-methyltransferases were also identified, five of which were under-expressed (four
800 PRDM6 and one setd3 isoform) and three of which were over-expressed in Om corals. As with histone
801 acetyltransferase, these enzymes are involved in gene expression regulation through their influence on
802 chromatin structure (Huang 2002; Vervoort *et al.* 2016). One lymphoid-specific helicase-like (HELLS)
803 gene was frontloaded in Om corals. This protein is known to participate in epigenetic processes, and
804 interacts with DNMT1 (cytosine methyl-transferase) for the maintenance of *de novo* methylation (Myant
805 *et al.* 2011; Dabe *et al.* 2015). We identified under-expression of two SID-1 transmembrane family
806 member genes involved in dsRNA regulation (Li *et al.* 2015). Many reverse transcriptase homologues,
807 including RNA-directed DNA polymerase from a jockey-like mobile element, were differentially expressed
808 or more intensely regulated during heat stress in the Om corals. These reverse transcriptases are typical
809 of the retro-transposon activity known to be activated or less controlled during general stress (Wessler

810 1996). The loss of inhibition of retro-transposon activity may reflect a trade-off with more specific
811 functions, but could also be a “SOS-like” survival mechanism. Indeed, in a threatening environment a high
812 transposition activity may induce mutations in polyps with potential genomic changes, which could fuel
813 rapid adaptive evolution (Maumus *et al.* 2009).

814 CONCLUSION:

815 Comparison of the response to an ecologically realistic heat stress of corals from the same genus but
816 pertaining to different phylogroups thriving in two contrasting thermal environments sheds light on the
817 molecular basis of thermotolerance. We found that during heat exposure the symbiotic community
818 composition was stable in colonies from both thermal environments, but we identified in coral hosts
819 major differences in gene regulation processes. The more thermotolerant colonies displayed (i) a more
820 plastic transcriptome response involving more differentially expressed genes and higher fold expression
821 changes; and (ii) a constitutive and higher level of expression for a range of genes (frontloaded genes). In
822 the context of climate change, which is predicted to cause abnormal and rapid temperature increase
823 (Change 2014), phenotypic plasticity and the capacity for rapid adaptation through epigenetic regulation
824 and/or genetic assimilation could increase the probability of coral survival. Previous studies highlighted
825 the importance of reef managements measures (Rogers *et al.* 2015) and assisted evolution (van Oppen *et*
826 *al.* 2015), but also underlined the importance of preserving standing genetic/epigenetic variation in wild
827 coral populations (Matz *et al.* 2017). Our results also suggest that management measures must include
828 protection of natural thermotolerant populations having the potential to resist increasing thermal
829 anomalies. Although the molecular mechanisms we described are most likely largely shared in this group
830 of scleractinians, the question remains as to whether these more thermotolerant genotypes are secured in
831 different phylogroups as was the case here, or whether they also exist inside more closely related
832 phylogroups where they could circulate via long range gene flow. (Mumby *et al.* 2011). It is however
833 essential to keep in mind that even the most thermotolerant reefs may bleach if they are exposed to
834 temperature significantly higher to their own norm (Hughes *et al.* 2017b; Le Nohaïc *et al.* 2017).

835

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849 **DATA ACCESSIBILITY**

850 The datasets generated and analyzed during the current study have been submitted to the SRA repository
851 under bioproject number PRJNA399069 (to be released upon publication).

852 **AUTHORS' CONTRIBUTIONS**

853 JVD, MA, DA, GM, and ET were involved in the study concept and design. KBR, LF, MC, MA, PR and JVD
854 were involved in the collection of samples. All authors were involved in data acquisition and analyses.
855 KBR, JVD, GM, OR and ET drafted the manuscript, and all authors contributed to critical revisions and
856 approved the final manuscript.

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863 **TABLES**

864 Table 1: Sea Surface Temperature (SST) regimes to which the colonies sampled in this study (i.e. Oman
865 and New Caledonia) are exposed in their natural environments. Thermal regime descriptors were
866 compiled from weekly mean sea surface temperature data collected from the Integrated Global Ocean
867 Services System Products Bulletin (i.e. IGOSS: <http://iridl.ldeo.columbia.edu/SOURCES/IGOSS/>) for
868 quadrats of 1° longitude X 1° latitude and from 1982 to the year of sampling (2013-2014).

| | New Caledonia | Oman |
|----------------------------------|------------------|------|
| Mean SST (°C) | 24.8 | 27.9 |
| Variance (°C) | 2.7 | 9.5 |
| Min SST (°C) | 22.6 | 22.1 |
| Max SST (°C) | 27.1 | 33.2 |
| Mean SST of 3 warmer months (°C) | 26.8 | 31.3 |
| Mean SST of 3 cooler months (°C) | 22.8 | 23.8 |

869

870 **FIGURE LEGENDS**

871 FIGURE 1: The ecologically realistic heat stress experiment: from mean temperatures of the warmer
872 months *in natura* to a pre-bleaching physiological state. The arrows represent points at which nubbins
873 were collected for analyzing the microbial composition and the transcriptomic response of the host.

874 FIGURE 2: Principal coordinate analysis plot for Bray-Curtis distances of the bacterial composition of each
875 colony in each experimental condition. Different colors represent different colonies, the circles represent
876 the *in situ* conditions, the triangles represent the control conditions, and the squares represent the stress
877 conditions.

878 FIGURE 3: Composition of the *Symbiodinium* community in each colony *in situ* and in controlled and
879 stressful experimental conditions.

880 FIGURE 4: Hierarchical clustering analyses performed using DESeq2 rlog-normalized RNA-seq data for the
881 36 transcriptomes: two conditions (control and heat stress); three replicates per condition for each
882 colony; three colonies per locality; and two localities [Oman (Om, i.e. thermotolerant) and New Caledonia
883 (NC; thermosensitive)]. The color (from white to dark blue) indicates the distance metric used for
884 clustering (dark blue corresponds to the maximum correlation values).

885 FIGURE 5: Heatmap for significant differential gene expression in at least one comparison. Each gene is
886 represented by a line.

887 FIGURE 6: Scatterplot of the log₂-fold changes in gene expression in response to heat stress in the Om
888 colonies (y-axis) vs. the NC colonies (x-axis) for the 848 genes that were over-expressed (498 genes) or
889 under-expressed (350 genes) in colonies from both localities. The line represents the y=x line depicting
890 similar responses between colonies.

891 FIGURE 7: Colony level gene expression variation in response to heat stress, based on DAPC analysis. The
892 x axis is the first discriminant function of the DAPC along which the overall gene expression difference
893 between colonies at both experimental conditions (stress and control) and from both localities (NC and
894 Om) was maximized. This indicates the degree of similarity between the transcriptomes. The density plots
895 obtained for NC and Om colonies are represented in blue and green, respectively. Dark and light density
896 plots correspond to samples from the control and stress experimental conditions. The arrows above the
897 density plots represent the direction of the mean change in the gene expression profiles.

898 FIGURE 8: Summary of the GO enrichment analysis following REVIGO synthesis. Each enriched biological
899 process is represented by a bar proportional to the log₁₀(FDR). The colors correspond to the three
900 categories of genes (common: black; Om-specific: grey; NC-specific: white) that were over-expressed (left
901 panel) or under-expressed (right panel).

902 ADDITIONAL FILES

903 Additional Figure 1. Bacterial class composition (for the 24 most abundant) within each replicate for the
904 Om and NC colonies, the three colonies of each locality, and three experimental conditions per colony. *In*
905 *situ* (dark arrows); control temperature (green arrows); stress temperature (red arrows).

906 Additional Table 2: ANOVA results for alpha diversity (Shannon index) and beta diversity (Bray-Curtis
907 distance) between localities, colonies, or experimental conditions.

908 Additional File 3: List and sequences of the 26,600 genes (XLOC) generated during RNAseq alignment.

909 Additional Table 4: Haplotype analysis of the six sampled colonies with microsatellite genotyping for the
910 colonies from New Caledonia.

911 Additional File 5: DEseq2 results for the log₂-foldchanges, and adjusted *p* values between stress and
912 control conditions for each locality.

913 Additional Table 6: Comparison between the log₂-foldchange in Om and NC colonies for genes
914 differentially under-expressed or over-expressed in the same way in colonies from both localities.

915 Additional File 7: GO enrichment results for biological processes, molecular functions, and cellular
916 compartments for common, New Caledonia-specific, or Oman-specific over-expressed and under-
917 expressed genes.

918 Additional File 8: Frontloaded genes in Oman corals among genes over-expressed in New Caledonia
919 corals.

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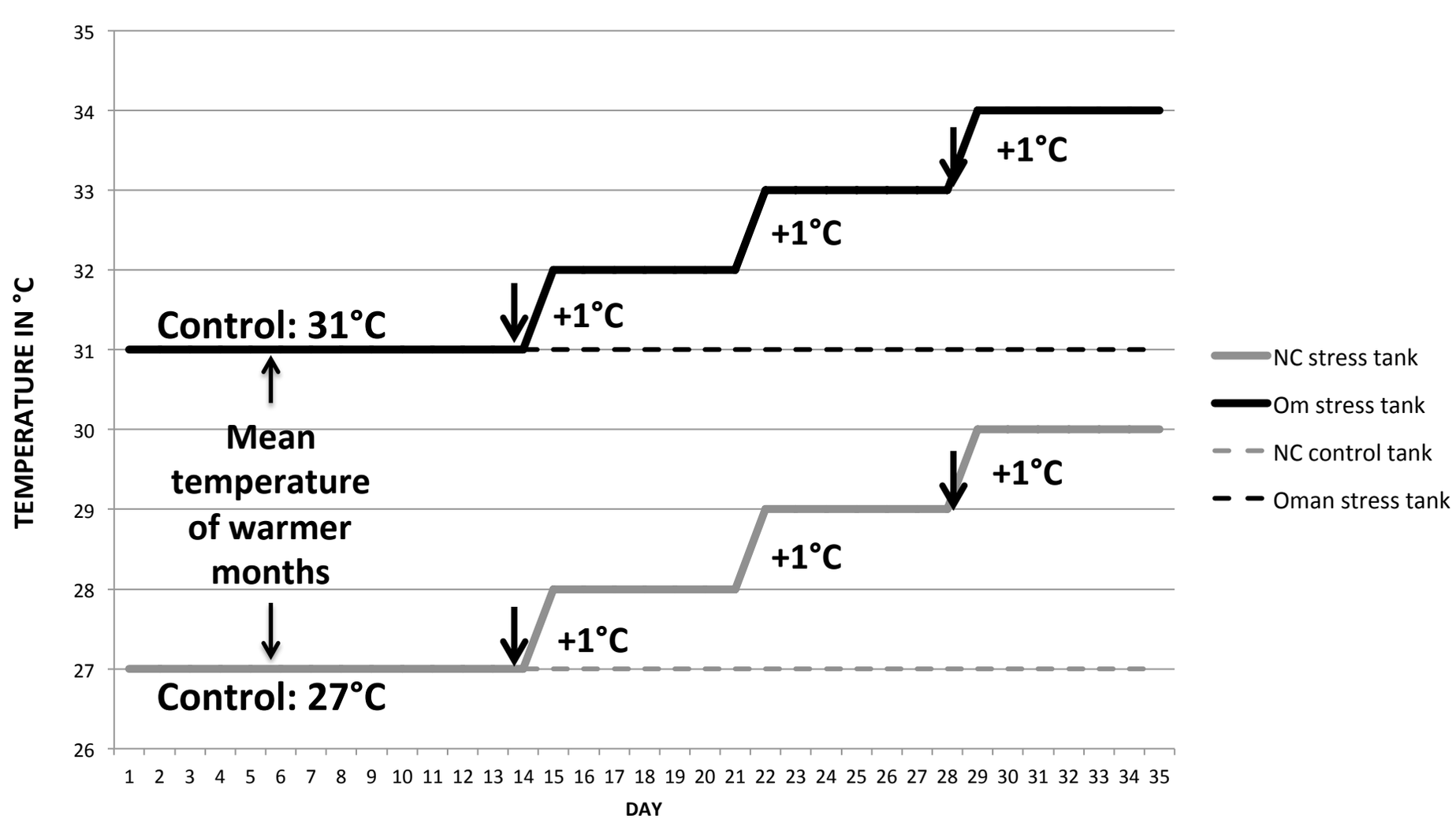
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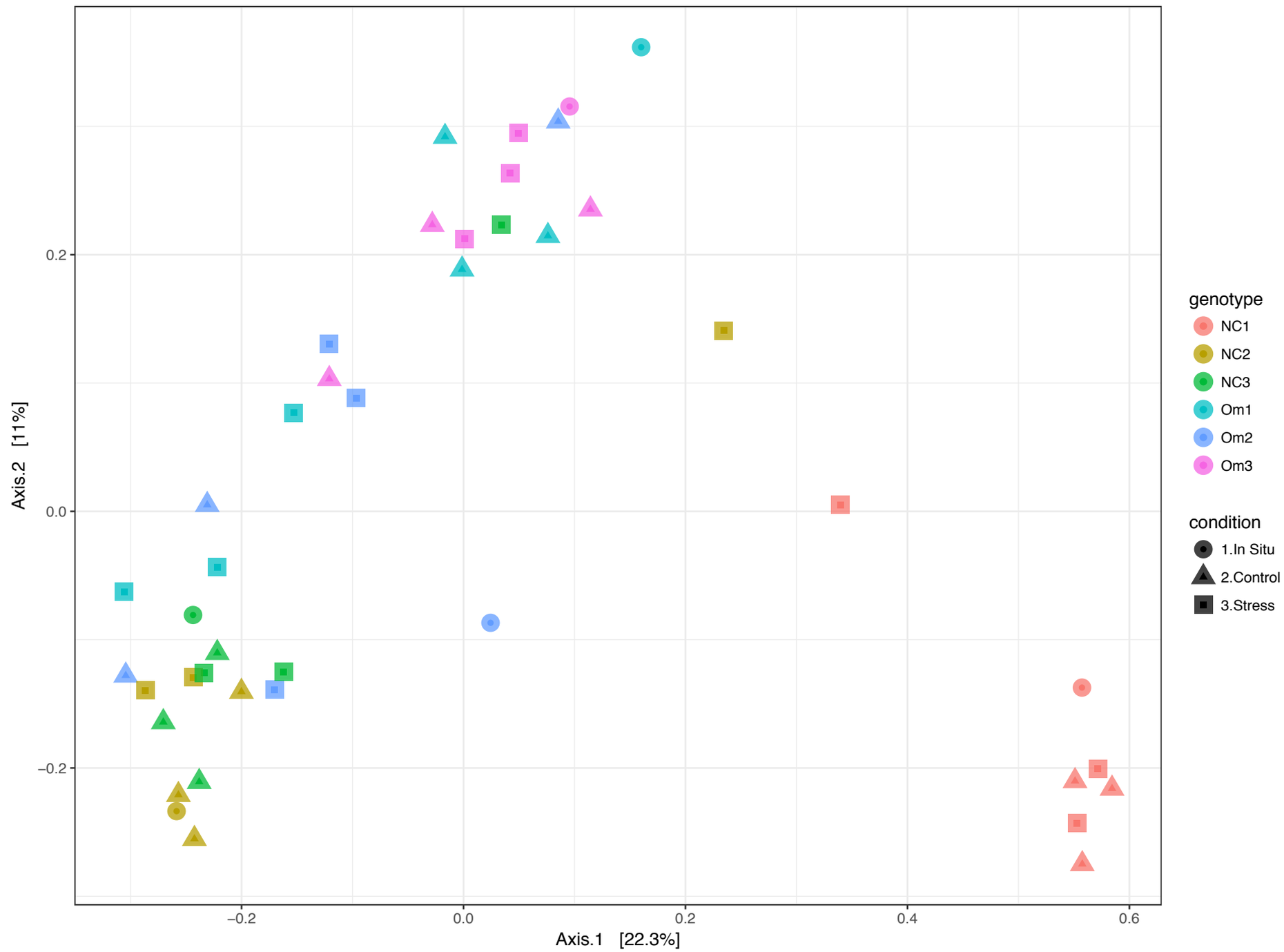
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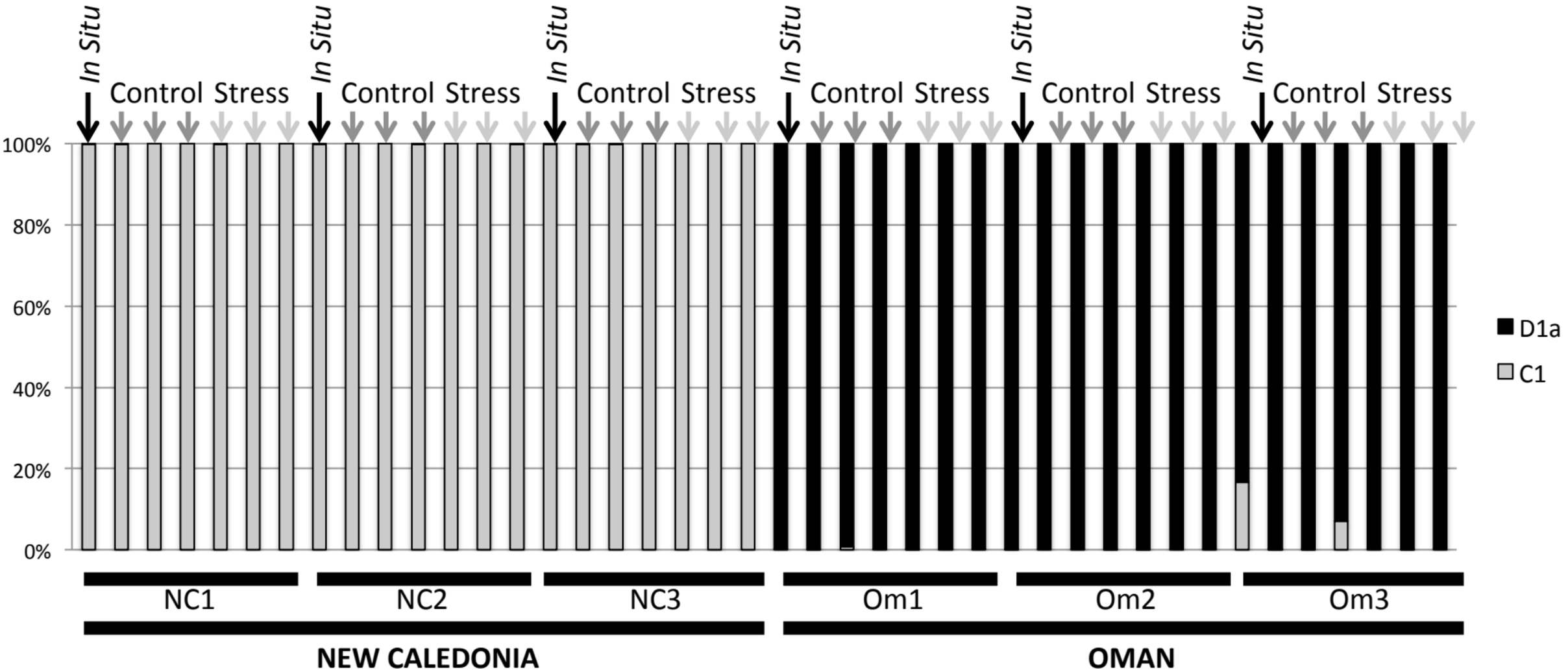
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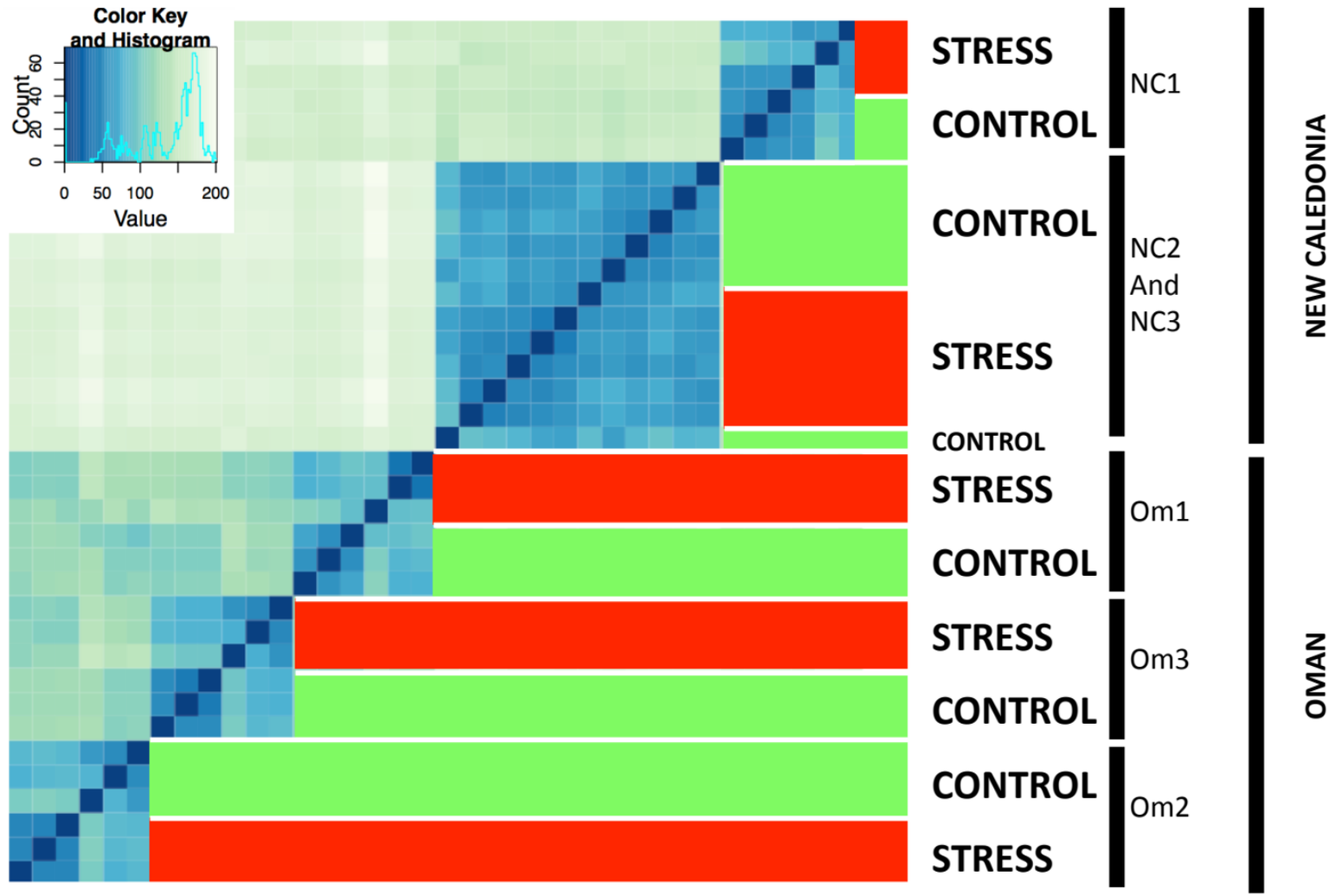
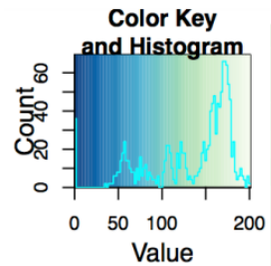
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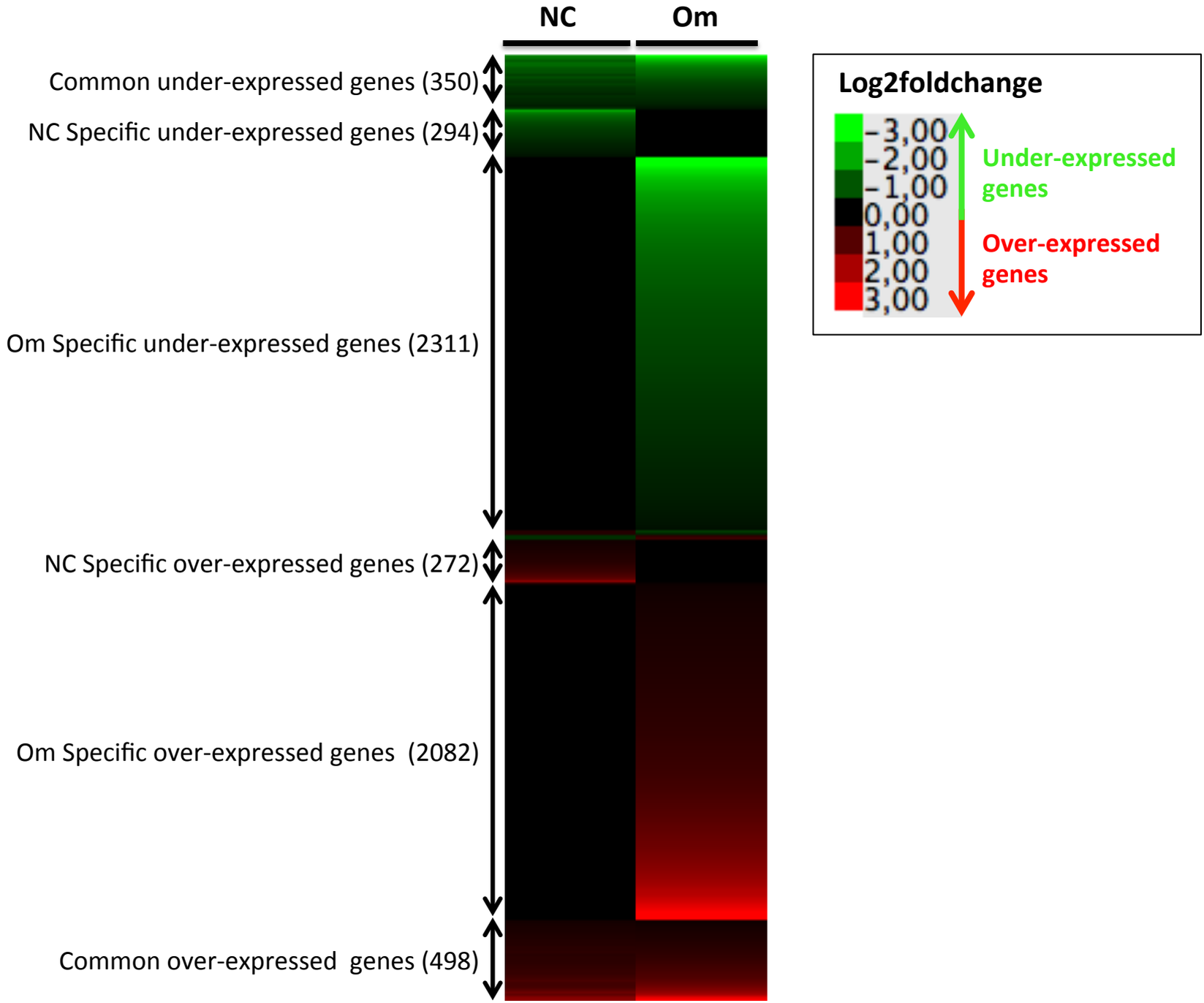


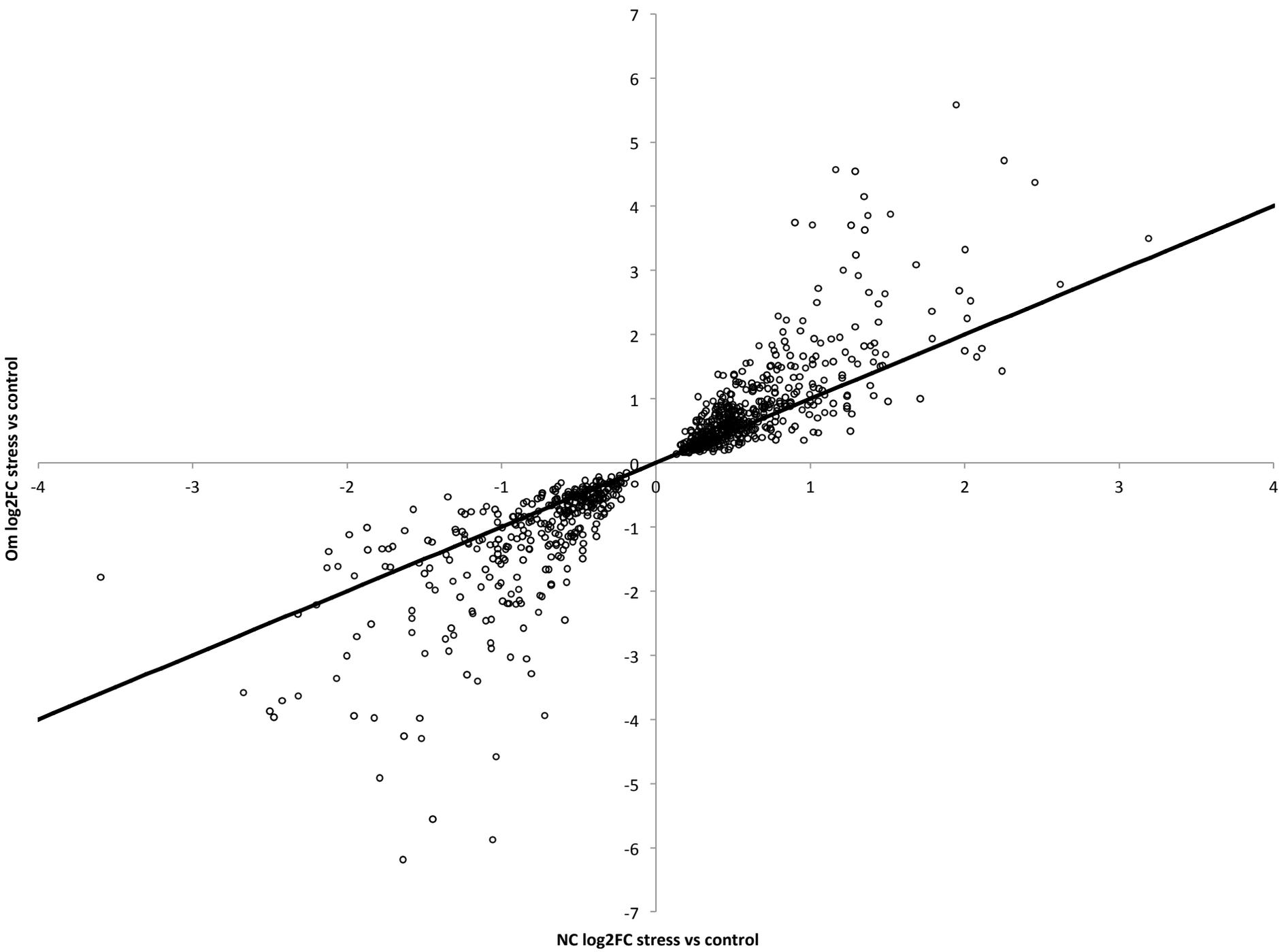
MDS + BC

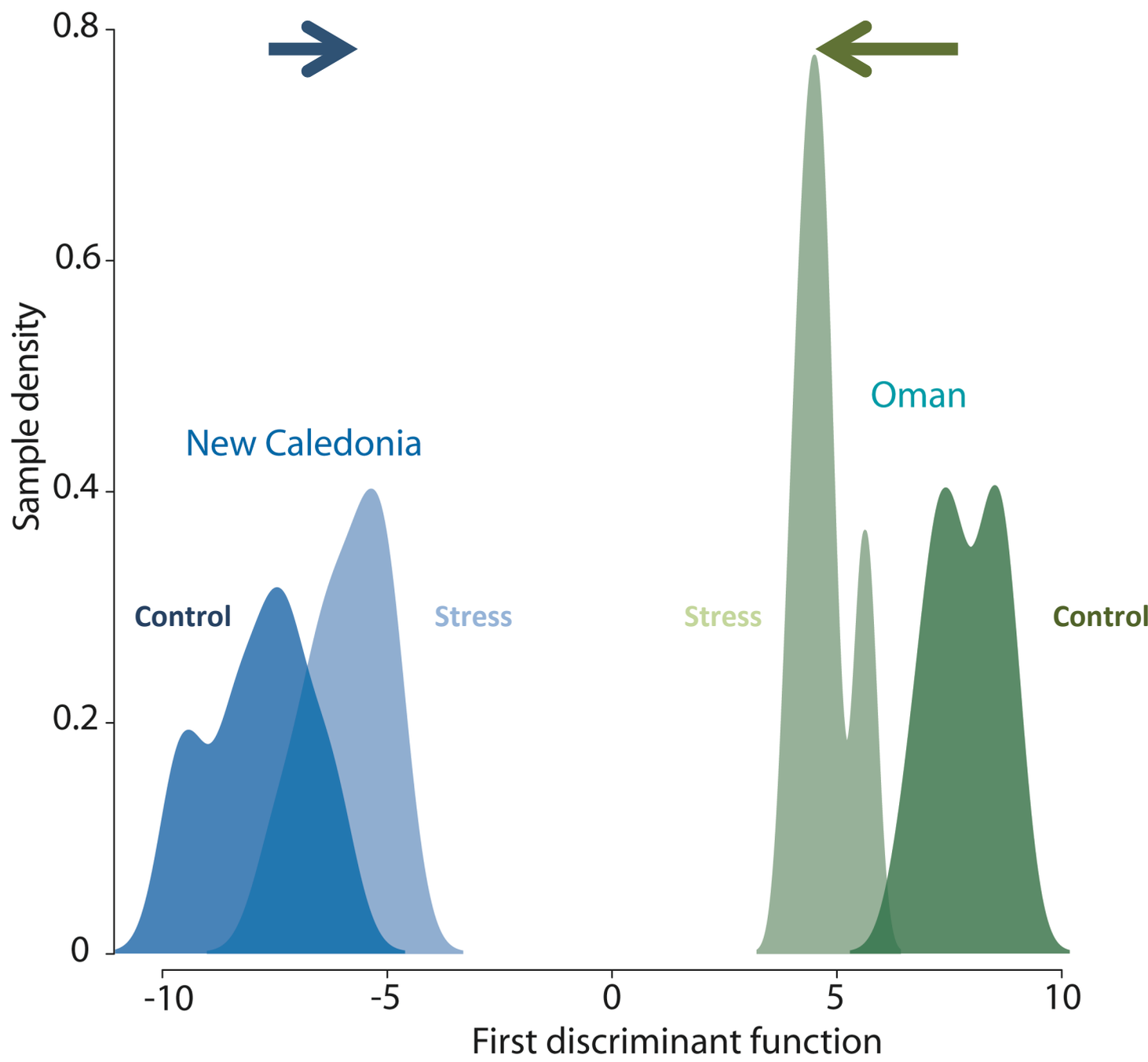




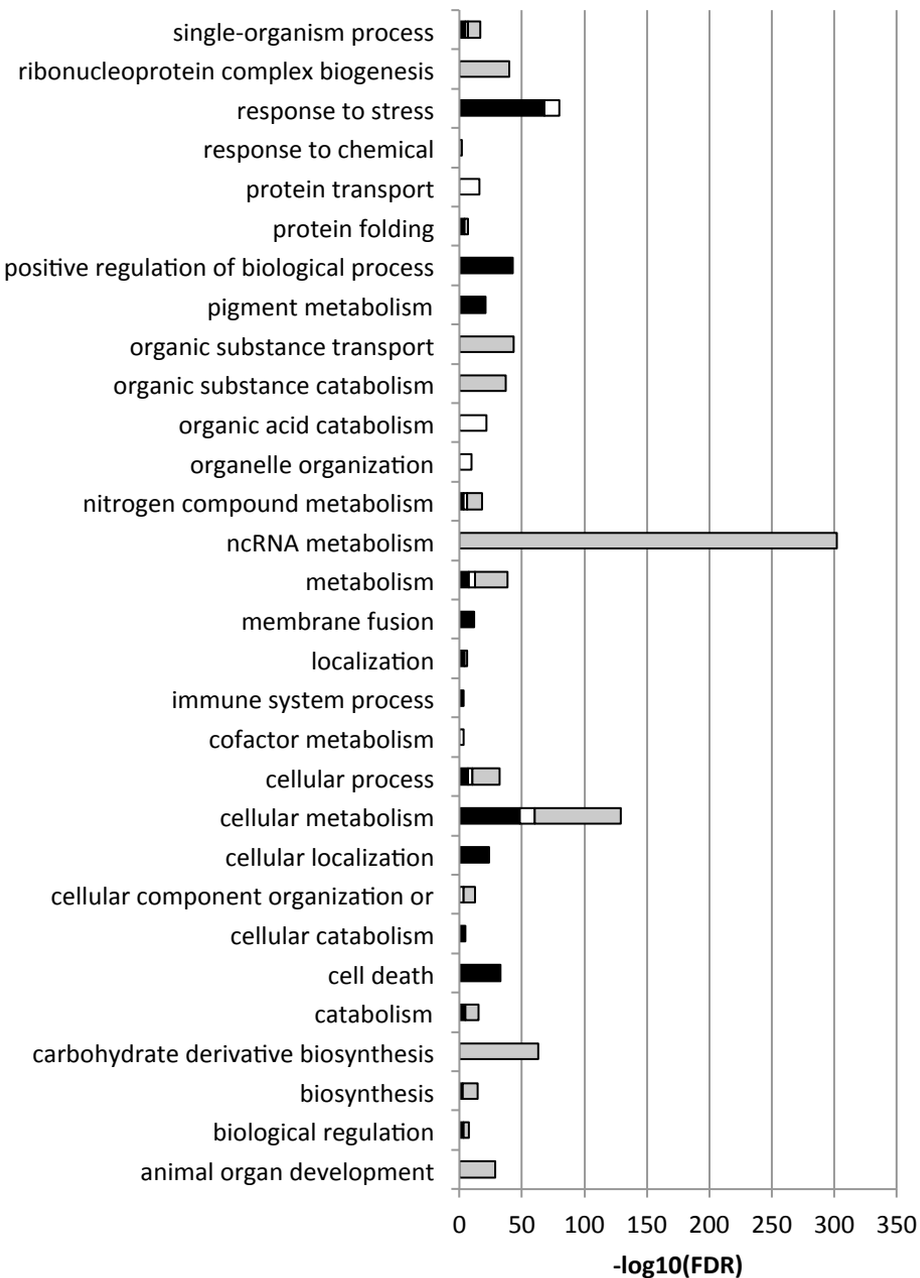








Over-expressed genes



Under-expressed genes

