1	Identification of novel genes involved in high hydrostatic pressure resistance
2	of Escherichia coli
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### 21 Abstract

22 High hydrostatic pressure (HHP) is an interesting hurdle in minimal food processing that aims to 23 synergistically combine different stresses to improve food microbiological safety and stability 24 without compromising quality. For a proper understanding and design of hurdle technology, the 25 cellular impact of the applied stresses on foodborne pathogens should be well-established. To study the mechanism of HHP-mediated cell injury and death, we screened for loss-of-function 26 27 mutations in E. coli MG1655 that affected HHP sensitivity. More specifically, ca. 6,000 random 28 transposon insertion mutants were individually exposed to HHP, after which the phenotype of the most resistant or sensitive mutations was confirmed by *de novo* gene deletions in the parental 29 30 strain. We found that disruption of *rbsK*, *rbsR*, *hdfR* and *crl* decreased HHP resistance, while disruption of sucC and sucD (encoding subunits of the succinyl-CoA synthetase) increased HHP 31 resistance. More detailed study of the tricarboxylic acid cycle enzymes encoded by the 32 sdhCDAB-sucABCD operon surprisingly showed that disruption of the sucA or sucB gene 33 34 (encoding subunits of the 2-oxoglutarate dehydrogenase complex) notably decreased HHP 35 survival. We also found that the increased HHP resistance of a  $\Delta sucC$  and  $\Delta sucD$  mutant was mediated by increased basal RpoS activity levels, although it did not correlate with their heat 36 37 resistance. Our results reveal that compromising TCA cycle enzymes can profoundly affect HHP 38 resistance in E. coli.

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40 Key-words: *E. coli*; high hydrostatic pressure; heat; resistance; TCA cycle

#### 42 **1. Introduction**

Consumers' tendency for healthy fresh-like foods requires the development of mild food 43 preservation methods that minimally affect the sensorial and nutritional properties of the food 44 45 product while at the same time ensuring food safety. To avoid intensive heat processing, a number of non-thermal food processing techniques capable of inactivating pathogenic and 46 spoilage microorganisms have emerged, including high hydrostatic pressure (HHP), pulsed 47 48 electric fields, ultrasound and UV-based irradiation (Barba et al., 2017). HHP processing is one 49 of the most accepted technologies, being already used in the food industry to improve safety and extend shelf life of a variety of products (Huang et al., 2017). Unfortunately, full exploitation of 50 51 HHP pasteurization is hampered in part by the considerable HHP resistance that some pathogenic bacteria tend to display naturally (Bruschi et al., 2017; Liu et al., 2015) or can readily 52 acquire upon recurrent exposure to HHP processing (Vanlint et al. 2012). In addition, the high 53 capital investment in HHP equipment and the need to combine pressure and thermal treatment to 54 inactivate bacterial endospores limit the use of this technology in certain type of foods (Elamin et 55 56 al., 2015; Khan et al., 2017). Rather than using intense HHP processing to ensure inactivation of 57 resistant bacteria at the expense of food quality, HHP can be combined with other stresses in a hurdle-type approach. In fact, bacteria surviving HHP exposure tend to be sublethally injured 58 59 (Sokołowska et al., 2014; Somolinos et al., 2008) and therefore sensitized to a number of subsequent stresses (Koseki et al., 2008; Somolinos et al., 2008). As such, combining mild HHP 60 treatment with other preservation methods at moderate intensity could be a promising strategy to 61 62 adequately control microorganisms while reducing the loss of food quality and processing costs (Khan et al., 2017; Oliveira et al., 2015). Indeed, the combination of HHP with mild heat, low 63 pH, oxidative stress and certain natural antimicrobial compounds, such as reuterin,  $\alpha$ , $\beta$ -64

unsaturated aldehydes or isothiocyanates, has been demonstrated to synergistically enhance
inactivation (Feyaerts et al., 2015; Gao et al., 2006; Montiel et al., 2015; Muñoz et al., 2006;
Wang et al., 2012). However, the combination of HHP with other compounds, such as carvacrol,
linalool or eugenol, only exerted an additive or even an antagonistic lethal effect (Feyaerts et al.,
2015).

Paramount to the proper design of effective hurdle approaches is a better understanding of the 70 71 cellular impact of available preservation stresses. With respect to HHP processing, current 72 knowledge on its effects on bacteria is mainly coined from the study of (i) HHP on biomolecules (Gayán et al. 2017a; Rivalain et al., 2010), (ii) cellular responses to HHP stresses (Bowman et 73 74 al., 2008; Malone et al., 2006), and (iii) the genetic analysis of bacterial mutants that were 75 selected to become HHP resistant through directed evolution approaches (Gayán et al. 2017b; Vanlint et al., 2013a, 2013b). As such, HHP was shown to hamper the functionality of several 76 cellular components including phospholipid bilayers, proteins, and ribosomes, which in turn 77 pleiotropically compromises cellular homeostasis (Gayán et al. 2017a; Rivalain et al., 2010). The 78 79 RpoS general stress response has proven to be important for HHP resistance in E. coli, and the 80 absence of this sigma factor renders *E. coli* hypersensitive to HHP (Charoenwong et al., 2011; Gayán et al. 2017b). Furthermore, basal RpoS activity levels correlate well with HHP resistance 81 82 variability among Shiga-toxin producing E. coli isolates (Álvarez-Ordóñez et al., 2013; Robey et al., 2001), while mutants with increased RpoS activity are rapidly selected for after iteratively 83 exposing E. coli O157:H7 to HHP stress (Vanlint et al., 2013a). However, other response 84 85 pathways were shown to be important as well, and E. coli MG1655 mutants lacking the RpoS sigma factor can restore HHP resistance by acquiring mutations causing cAMP/CRP (cyclic 86 adenosine monophosphate/cAMP receptor protein) downregulation (Gayán et al. 2017b). 87

88	Furthermore, upregulation of the heat shock response has been reported in E. coli after HHP
89	stress (Aertsen et al., 2004; Malone et al., 2006), and HHP-resistant mutants of E. coli and L.
90	monocytogenes with increased basal levels of heat shock proteins have been isolated (Aertsen et
91	al., 2004; Gayán et al., 2016a; Van Boeijen et al., 2010).
92	Nevertheless, our view on the cellular impact of HHP still remains fragmentary and needs to be
93	further refined with alternative complementary experimental approaches. Whereas the above
94	mentioned directed evolution approach specifically revealed genes and functions that are altered
95	in mutants selected to acquire HHP resistance, we decided to embark in assessing the
96	contribution of individual E. coli genes to HHP sensitivity or resistance. More specifically, ca.
97	6,000 single-gene disruption mutants of a random E. coli transposon insertion library were
98	examined for their ability to survive a HHP shock.

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# 101 2. Material and Methods

# 102 **2.1. Bacterial strains, mutant construction and growth conditions**

103 *E. coli* K-12 MG1655 (Blattner et al., 1997) and its derivatives described in Table 1 and 2 were 104 used throughout this study. Transposon insertion mutagenesis was performed using the 105 Tn*10*dCm transposon system and  $\lambda$ NK1324 as described by Kleckner *et al.* (1991). In-frame 106 gene deletions were performed according to the method of Datsenko and Wanner (2000), using 107 the MG1655 parental strain equipped with the plasmid pKD46 (encoding the  $\lambda$  red recombinase

108	genes behind the araBAD promoter) and an amplicon prepared on pKD13 (containing the
109	kanamycin resistance cassette) using the primers listed in Baba et al. (2006). The kanamycin
110	marker was flanked by FRT sites and was further excised by transiently equipping the strain with
111	the plasmid pCP20 (expressing the Flp site-specific recombinase (Cherepanov and Wackernagel,
112	1995)) to obtain the desired deletion mutant. Deletion constructs were verified by PCR and
113	sequencing (Macrogen; Amsterdam, The Netherlands) using the locus specific primers listed in
114	Table S1. Where indicated, the strains were transformed with pFPV-P <sub>bolA</sub> -gfp (encoding the E.
115	coli MG1655 bolA promoter upstream of gfp (Gayán et al., 2016a)) or pFPV-P <sub>dnaK</sub> -gfp (encoding
116	the E. coli MG1655 dnaK promoter upstream of gfp (Aertsen et al., 2004)) by electroporation.
117	Strains were grown in Lysogeny Broth (LB) medium (Miller, 1992) and when necessary, a final
118	concentration of 50 $\mu$ g/ml of kanamycin (Km; Panreac-AppliChem, Darmstadt, Germany), 100
119	$\mu$ g/ml of ampicillin (Ap; Thermo Fisher Scientific, Waltham, MA, USA) or 30 $\mu$ g/ml of
120	chloramphenicol (Cm; Sigma-Aldrich, St. Louis, MO, USA) was added to select for the presence
121	of recombined amplicons (Datsenko and Wanner, 2000) or pKD46 and pCP20 vectors
122	(Cherepanov and Wackernagel, 1995; Datsenko and Wanner, 2000) or Tn10dCm transposon
123	insertions (Kleckner et al., 1991), respectively. For inactivation experiments, test tubes
124	containing 4 ml of LB were inoculated with a single colony and then incubated aerobically with
125	shaking (300 rpm) for 18 h at 37°C to obtain stationary phase cultures containing $ca$ . 10 <sup>9</sup> Colony
126	Forming Units per milliliter (CFU/ml). For pFPV-P <sub>bolA</sub> -gfp and pFPV-P <sub>dnaK</sub> -gfp reporter assays,
127	cells were grown in LB supplemented with Ap to select for the presence of the plasmids (Aertsen
128	et al., 2004; Gayán et al., 2016a) and for the $\beta$ -galactosidase assay, a final concentration of 1 mM
129	of isopropyl $\beta$ -D-1-thiogalactopyranoside (IPTG; Acros Organics, Morris Plains, NJ, USA) was
130	added to induce the <i>lac</i> operon.

### 132 2.2. HHP and heat treatment

Cells from a stationary phase culture were harvested by centrifugation  $(4000 \times g, 5 \text{ min})$  and 133 resuspended in an equal volume of 10 mM potassium phosphate buffer of pH 7.0. For HHP 134 treatment, 200  $\mu$ l of the cell suspension was heat-sealed in a sterile polyethylene bag after 135 exclusion of the air bubbles and subjected to 300 MPa or 400 MPa for 15 min in an 8-ml 136 pressure vessel (HPIU-10000, 95/1994; Resato, Roden, The Netherlands), held at 20°C with an 137 external water jacket connected to a cryostat. Both the slow pressure increase (100 MPa/min) 138 and the external water jacket attenuated adiabatic heating during pressure build-up. Finally, 139 140 decompression was almost instantaneous. For heat treatment, three sterile PCR tubes were 141 aseptically filled with a 65 µl portion of resuspended cells and subjected to 57.0°C for 15 min using a PCR apparatus (T-personal 48; Biometra GmbH, Goettingen, Germany). After HHP or 142 143 heat treatment, samples were aseptically retrieved from the polyethylene bags or PCR tubes, and survival was determined as described below. 144

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#### 146 **2.3. Determination of viability**

Samples were serially diluted in 10 mM potassium phosphate buffer, and subsequently a 5-µl
sample of each dilution was spotted onto LB agar in triplicate, as previously described
(Sieuwerts et al., 2008). After 24 h of incubation at 37°C, spots containing between 5–50
colonies were counted, so that the quantification limit was 1,000 CFU/ml. The logarithmic

reduction factor was calculated as  $\log_{10} (N_0/N)$ , in which  $N_0$  and N represent the number of survivors in CFU/ml prior and after treatment, respectively.

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# 154 2.4. Screening for transposon insertion mutants of *E. coli* MG1655 with altered HHP 155 resistance

To screen for changes in HHP resistance, ca. 6,000 transposon insertion mutants of E. coli 156 MG1655 were individually grown in LB in 24-well microtiter plates and individually exposed to 157 a HHP shock of 300 MPa (15 min, 20°C). Mutants that presented at least a 2-log higher or lower 158 HHP inactivation compared to the WT strain were kept for further study. To confirm that the 159 transposon insertion was involved in the HHP resistance change of selected mutants, the locus 160 161 with the transposon was transferred by P1-transduction (Sambrook and Russell, 2001) to the WT strain and the transductant was then subjected to 400 MPa (15 min, 20°C) in potassium 162 phosphate buffer (10 mM, pH 7.0). The transposon location in the bacterial genome was mapped 163 164 according to the method of Kwon and Ricke (2000). In brief, mutants' genomic DNA was digested with NlaIII (Thermo Fisher Scientific) and ligated to the Y-shaped linker. Subsequently, 165 the transposon flanking region was amplified using the Y-linker primer (specific to Y-linker 166 167 sequence) and a transposon specific primer (NK\_Cm\_DWN; Table S1) and sequenced. Sequences were compared against the *E. coli* MG1655 genome published at the GenBank 168 169 database (Blattner et al., 1997) using the NCBI BLASTn (Nucleotide Basic Local Alignment Search Tool, blast.ncbi.nlm.nih.gov). 170

#### 172 **2.5.** Measurement of RpoS, RpoH and cAMP/CRP activity reporter

The basal levels of RpoS sigma factor (directing expression of the general stress response) were 173 174 quantified by the activity of the *bolA* promoter (P<sub>bolA</sub>) using the pFPV-P<sub>bolA</sub>-gfp construct (Gayán et al., 2016a), while RpoH levels (directing expression of the heat shock response) were 175 176 measured by the *dnaK* promoter activity ( $P_{dnaK}$ ) using the pFPV- $P_{dnaK}$ -gfp vector (Aertsen et al., 177 2004). To determine the fluorescence derived from the reporter, 200  $\mu$ l of the stationary phase cultures equipped with pFPV-PbolA-gfp or pFPV-PdnaK-gfp were transferred to microplate wells 178 and placed in a Fluoroscan Ascent FL (Thermo 180 Labsystems, Brussels, Belgium). The basal 179 GFP fluorescence was measured at an excitation wavelength of 480 nm and an emission 180 wavelength of 520 nm. The obtained fluorescence values were subsequently divided by the 181 optical density at 600 nm ( $OD_{600}$ ) of the same sample to obtain the relative fluorescence units. 182 Differences in RpoS and RpoH activity are expressed as fold change with respect to the parental 183 strain. 184

The cAMP/CRP activity was indirectly tested by measuring β-galactosidase activity of
permeabilized cells using ortho-nitrophenyl-β-galactoside (ONPG; Acros Organics) as a
substrate in a Multiskan RC (Thermo Labsystems, Vantaa, Finland). Enzyme activity was
expressed in Miller units (MU) (Miller, 1992).

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# 190 **2.5. Sensitivity to hydrogen peroxide**

The susceptibility of *E. coli* to hydrogen peroxide was evaluated by the size of the growth
inhibition zone. An aliquot of 100 µl of stationary phase cultures was plated into 15 ml of LB

193	soft agar. After drying, a volume of 5 $\mu$ l of a 30% (w/w) hydrogen peroxide solution (Acros
194	Organics) was dropped on the center of the plate. The diameter of the inhibition zone was
195	measured after overnight incubation at 37°C.

#### 197 **2.6. Statistical analysis**

198	Statistical analyse	s (ANOVA, Bonferron	i multiple compariso	on test) were carrie	d out using the
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software GraphPad PRISM 5.0 (GraphPad Software Inc., San Diego, CA, USA), and differences

200 were regarded as significant when P was  $\leq 0.05$ . All microbial inactivation, fluorescence and

201 enzymatic activity data shown in figures correspond to averages and standard deviations

202 calculated from three replicates performed in different working days.

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# 205 **3. Results and Discussion**

### **3.1.** Screening for mutants of *E. coli* MG1655 with HHP resistance change

207 To identify novel genes involved in stationary phase HHP resistance, *ca*. 6,000 random

transposon insertion mutants of *E. coli* MG1655 were individually examined for their ability to

- survive a HHP shock. From this library, six and two mutants showed a confirmed  $\geq 2.0 \log$  cycle
- higher ( $P \le 0.05$ ) sensitivity and resistance to a 400 MPa (15 min, 20°C) shock than the wild-
- type (WT) strain, respectively (Table 2). The exact position of the transposon in each mutant

with the function of the gene disrupted is included in Table 2. The transposon insertion in the
two resistant mutants was located in the *sucD* gene and in the intergenic region between *sucB*and *sucC*, whereas in the six sensitive mutants the transposon was mapped in either the *rbsK*, *rbsR*, *hdfR* or *crl* gene (with *crl* and *hdfR* disruptions independently found twice).

216 In order to confirm the role of the identified genes in pressure resistance, each gene was deleted 217 de novo in the WT strain, and the corresponding mutants were exposed to a 400 MPa (15 min, 218 20°C) shock that caused ca. 3.2 log reductions of the WT strain (Fig. 1). As shown in Fig. 1, 219 deletion of the transcriptional regulator HdfR, which positively controls the glutamate-dependent 220 acid resistance response (Krin et al., 2010; Reynolds et al., 2017) and negatively controls the 221 FlhDC regulator for flagellum biogenesis (Ko and Park, 2000), caused the highest HHP sensitivity reaching 5.9 log cycles of inactivation. Mutants  $\Delta rbsR$ ,  $\Delta rbsK$  and  $\Delta crl$  showed ca. 2 222 223 log cycles lower ( $P \le 0.05$ ) HHP resistance than the WT strain. The *rbsR* and *rbsK* genes form part of the *rbs* operon involved in ribose catabolism and transport, and encode the ribokinase and 224 225 the transcriptional repressor of the operon, respectively (Shimada et al., 2013). On the other hand, Crl enhances RpoS activity by facilitating the assembly of  $\sigma^{S}$  to the RNA polymerase core 226 in detriment to  $\sigma^{70}$  (involved in the transcription of most genes during exponential phase) and 227 protecting  $\sigma^{S}$  from proteolysis, and it specifically promotes the activity of certain  $\sigma^{S}$ -dependent 228 229 promoters as well (Dudin et al., 2013; Robbe-Saule et al., 2007; Typas et al., 2007). Indeed, 230 disruption of *crl* was previously shown to decrease the resistance to thermal, oxidative and acidic stress in Salmonella enterica subsp. enterica serovar Typhimurium (Robbe-Saule et al., 2008; 231 232 Robbe-Saule et al., 2007).

233	Regarding HHP-resistant mutants, the absence of either of the succinyl-CoA synthetase (SCS)
234	subunits (encoded by <i>sucC</i> and <i>sucD</i> genes) significantly ( $P \le 0.05$ ) increased the HHP
235	resistance of the WT strain by ca. 2 log cycles (Fig. 1). SCS catalyzes the reversible conversion
236	of succinyl-CoA to succinate, accompanied by the generation of a nucleoside triphosphate
237	molecule and coenzyme A (Park et al., 1997). The <i>sucC</i> and <i>sucD</i> genes form part of the
238	sdhCDAB-sucABCD operon (Cunningham and Guest, 1998; Park et al., 1997), which contributes
239	to two additional tricarboxylic acid (TCA) cycle enzymes: the succinate-quinone oxidoreductase
240	complex (SQR; encoded by <i>sdhCDAB</i> ) and the 2-oxoglutarate dehydrogenase complex (OGDH),
241	with the last consisting of the oxoglutarate decarboxylase (E1; encoded by <i>sucA</i> ), the
242	dihydrolipoyltranssuccinylase (E2; encoded by <i>sucB</i> ) and the lipoamide dehydrogenase (E3;
243	encoded by <i>lpd</i> ) subunit. OGDH converts 2-oxoglutarate to succinyl-CoA and CO <sub>2</sub> coupled to
244	NADH formation (Park et al., 1997), while SQR oxidates succinate to fumarate with the
245	reduction of ubiquinone to ubiquinol (Hagerhall, 1997).
246	Although some disruptions independently appeared twice in our screening (cfr. hdfR and crl), the
247	screening of ca. 6,000 random transposon mutants was likely not sufficient to redundantly cover

all non-essential genes of E. coli, since some gene disruptions previously known to significantly 248

modulate HHP resistance (such as rpoS, crp or cyaA (Gayán et al., 2017b; Vanlint et al., 2013a, 249

2013b) did not appear in the screening. 250

Because of their resistance conferring ability, the impact of sucC and sucD disruptions were the 251 subject of further scrutiny. 252

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#### 3.2. Deletion of *sucA* and *sucB* sensitizes *E. coli* to HHP treatment

Since the *sucCD* genes are part of a larger *sdhCDAB-sucABCD* operon, we decided to assess the 255 256 impact of the upstream genes on E. coli's HHP resistance. Fig. 1 correspondingly shows the inactivation (400 MPa, 15 min, 20°C) of MG1655 harboring an individual deletion of each gene 257 within the operon. While the lack of any of the SQR subunits (*i.e.* SdhC/D/A/B) did not 258 significantly (P > 0.05) affect HHP resistance, the deletion of *sucA* or *sucB* surprisingly 259 260 sensitized ( $P \le 0.05$ ) the parental strain up to a 1,800-fold at 400 MPa. Please note that it was likely impossible to isolate the corresponding *sucA* nor *sucB* transposon insertion mutants in our 261 screening because either *sucAB* or *sucCD* are essential for cell viability (Yu et al., 2006), while 262 263 insertion of the transposon in the promoter or *sucAB* region would likely have (polarly) affected the expression of *sucABCD* as a whole. 264

Li et al. (2006a) described the different metabolic impact of  $\Delta sucA$  and  $\Delta sucC$  on E. coli 265 266 BW25113. As such, the blockage of OGDH in the  $\Delta sucA$  mutant decreased the carbon flow through isocitrate dehydrogenase (involved in the reversible reaction of isocitrate to 2-oxoglutare 267 268 producing CO<sub>2</sub> and NADPH) and activated the glyoxylate shunt to provide oxaloacetate, while 269 the  $\Delta sucC$  mutant refilled the TCA cycle intermediates without activating the glyoxylate cycle 270 (Li et al., 2006a). Surprisingly, deletion of the *lpd* gene (encoding the lipoamide dehydrogenase 271 subunit), which likewise compromises the OGDH complex and activates the glyoxylate shunt (Li 272 et al., 2006b), did not significantly (P > 0.05) alter HHP resistance (Fig. 1). Moreover, 273 upregulating the glyoxylate shunt by deleting the isocitrate dehydrogenase (*icd*) gene (Kabir and Shimizu, 2004) also failed (P > 0.05) to confer HHP sensitivity, suggesting that the shortcut of 274 275 the TCA cycle itself cannot explain the increased HHP sensitivity of  $\Delta sucA$  and  $\Delta sucB$  mutants.

# 3.3. HHP resistance of *∆sucA/B/C/D* correlates with RpoS levels but not with RpoH or cAMP/CRP activity

279	In order to obtain more mechanistic insights into the opposing HHP tolerance encountered in
280	$\Delta sucAB$ and $\Delta sucCD$ mutants, we wondered to which extent their phenotype could stem from
281	underlying alterations in the global regulatory pathways (such as cAMP/CRP, RpoH or RpoS
282	regulation (Aertsen et al., 2004; Gayán et al., 2017b, 2016a; Vanlint et al., 2013a, 2013b) that
283	have previously been found to affect HHP resistance in E. coli.
284	More specifically, since attenuated cAMP/CRP regulation was shown to coincide with increased
285	HHP resistance (Gayán et al., 2017b; Vanlint et al. 2013b), the $\beta$ -galactosidase activity in
286	$\Delta sucA/B/C/D$ mutants, and for comparison in a $\Delta crp$ and $\Delta cyaA$ mutant (lacking the cAMP)
287	receptor protein and the cAMP synthase, respectively), was evaluated as a proxy of cAMP/CRP
288	activity (Fig. 2). Compared to the wild-type strain, the $\Delta sucC$ and $\Delta sucD$ mutants displayed
289	similar ( $P > 0.05$ ) $\beta$ -galactosidase levels, indicating an unaltered cAMP/CRP regulation that
290	therefore cannot explain their increased HHP resistance. The observation that $\Delta sucA$ and $\Delta sucB$
291	mutants actually displayed slightly attenuated ( $P \le 0.05$ ) levels of $\beta$ -galactosidase activity
292	compared to the parental strain, seems likewise to rule out involvement of cAMP/CRP
293	upregulation in their HHP hypersensitive phenotype. Indeed, the lower $\beta$ -galactosidase activity
294	tends to agree with the fact that accumulation of intracellular 2-oxoglutarate (as occurs in a
295	$\Delta$ sucA mutant (Li et al., 2006a)) decreases cAMP levels (Doucette et al., 2011).

Increased expression of heat shock proteins has previously also been shown to contribute to HHP 296 resistance in E. coli (Aertsen et al., 2004; Gayán et al., 2016a). However, when examining 297 298 expression of the *dnaK* promoter (*i.e.* P<sub>*dnaK*</sub>, using the pPFV-P<sub>*dnaK*</sub>-gfp reporter plasmid (Aertsen et al., 2004)) as a proxy of the heat shock response governed by the RpoH sigma factor, the 299  $\Delta sucC$  and  $\Delta sucD$  mutants showed lower ( $P \leq 0.05$ )  $P_{dnaK}$  activity than the WT strain, while no 300 301 significant (P > 0.05) differences were observed among WT and its  $\Delta sucA/B$  mutants (Fig. 3). This indicates that decreased or increased levels of the heat shock response are not the 302 underlying cause of the HHP sensitive  $\Delta sucA/B$  or the HHP resistant  $\Delta sucC/D$  phenotypes, 303 304 respectively.

305 Finally, expression of the RpoS-dependent *bolA* promoter (*i.e.* P<sub>bolA</sub>, which can be fluorescently monitored using the pFPV-PbolA-gfp reporter plasmid (Lange and Hengge-Aronis, 1991; Gayán et 306 al., 2016a)) was quantified as a proxy of basal cellular RpoS activity (Fig. 4A). For proper 307 comparison,  $P_{bolA}$  expression in a  $\Delta rssB$  mutant with attenuated RpoS quenching (since RssB is 308 309 the RpoS anti-sigma factor (Battesti et al., 2011; Gayán et al., 2017b)), a  $\Delta rpoS$  mutant lacking the  $\sigma^{s}$  factor, and the  $\Delta crl$  mutant (also originating from this screen) lacking the RpoS 310 holoenzyme assembly factor (Typas et al., 2007) were included (Fig. 4A). This revealed that, 311 compared to the WT parent, the  $\Delta sucA$  mutant exhibited a decreased ( $P \le 0.05$ ) P<sub>bolA</sub> expression 312 313 similar to that of the  $\Delta crl$  mutant, while the  $\Delta sucB$  mutant displayed even lower ( $P \le 0.05$ ) levels than the  $\Delta sucA$  and  $\Delta crl$  mutants (Fig. 4A). However,  $\Delta sucA/B$  mutants were shown to display a 314 slower growth and a later entrance into stationary phase (Fig. S1), which could potentially 315 316 explain their attenuated RpoS activity levels. Interestingly, the survival of the  $\Delta sucA$  and  $\Delta sucB$ mutants to a 300 MPa (15 min, 20°C) shock was still significantly lower ( $P \le 0.05$ ) than that of 317

the  $\Delta rpoS$  and  $\Delta crl$  mutants (Fig. 4B), indicating that besides an attenuated RpoS activity additional factors might contribute to their HHP sensitivity.

320	Interestingly, the $\Delta sucC$ and $\Delta sucD$ mutants displayed no growth defects (Fig. S1) but a
321	significantly ( $P \le 0.05$ ) increased P <sub>bolA</sub> expression compared to the WT strain, with their P <sub>bolA</sub>
322	expression being equal ( $P > 0.05$ ) to that of the $\Delta rssB$ strain (Fig. 4A). Importantly, this increase
323	was shown to be truly RpoS dependent since additionally deleting the <i>rpoS</i> gene in the $\Delta sucC$
324	and $\Delta sucD$ mutants (resulting in MG1655 $\Delta sucC \Delta rpoS$ and $\Delta sucD \Delta rpoS$ , respectively)
325	abolished P <sub>bolA</sub> expression to the same extent ( $P > 0.05$ ) as in a $\Delta rpoS$ mutant (Fig. 4A).
326	Moreover, since lack of RpoS activity also reduced the HHP resistance of the $\Delta sucC \Delta rpoS$ and
327	$\Delta sucD \Delta rpoS$ double mutants to the same level ( $P > 0.05$ ) as a single $\Delta rpoS$ mutant (Fig. 4B), the
328	increased basal RpoS activity seems to be causally involved in the HHP resistance of the $\Delta sucC$
329	and $\Delta sucD$ mutants.

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# 331 **3.4.** Heat and hydrogen peroxide resistance of *sucA/B/C/D* mutants

Since our data suggested that the HHP resistance of  $\Delta sucC/D$  mutants (and partially of  $\Delta sucA/B$ mutants) was modulated by varying basal RpoS activity levels, we examined the resistance of these mutants to other stresses such as hydrogen peroxide and heat. Like the  $\Delta rpoS$  strain, the  $\Delta sucA$  and  $\Delta sucB$  mutants showed larger ( $P \le 0.05$ ) zones of growth inhibition by hydrogen peroxide than the WT strain (Fig. 5A), indicating the higher susceptibility of these mutants to oxidative stress. In agreement, Ma *et al.* (2010) revealed the higher sensitivity of a BW25113 *sucB* deletion mutant to hydrogen peroxide and also to low pH, salicylate acid and various

antibiotics than its parent. In contrast, the deletion of neither the *sucC/D* or *rssB* genes reduced 339 340 (P > 0.05) the growth inhibition zone of hydrogen peroxide, indicating that further upregulation 341 of the RpoS response does not necessarily improve hydrogen peroxide protection in MG1655. However, the  $\Delta sucA/B/C/D$  mutants showed a peculiar behavior against thermal stress (Fig. 5B). 342 Contrary to their HHP resistance, the  $\Delta sucC$  and  $\Delta sucD$  mutants were on average 1.3 log cycles 343 344 more sensitive ( $P \le 0.05$ ) to a heat shock (57°C, 15 min) than the parental strain, while the  $\Delta sucA$ mutant was ca. 1.2 log cycles more resistant ( $P \le 0.05$ ) and the  $\Delta sucB$  mutant equally resistant (P 345 346 > 0.05) to the WT strain. Therefore, while disruption of the sucA/B/C/D genes seems to differentially modulate the general stress response, which correlates with HHP and oxidative 347 stress resistance of  $\Delta sucA/B/C/D$  mutants, the lack of these genes has a lower and different 348 impact on heat survival suggesting that they might affect thermal tolerance by an additional 349 RpoS-independent mechanism. 350

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# 352 3.5. *sucC* and *sucD* genes are unaffected in spontaneous HHP-resistant mutants of *E. coli* 353 MG1655

Finally, since we identified that loss-of-function mutations in *sucC* or *sucD* gene could cause
significant HHP resistance in *E. coli* MG1655 and could hence support evolutionary routes
towards acquisition of HHP resistance, the corresponding *sucABCD* loci of five HHP-resistant
mutants of MG1655 previously obtained by directed evolution (*i.e.* LMM1010, LMM1020,
LMM1030, DVL1 and DVL20 (Hauben et al, 1997; Vanlint et al., 2012, 2011)) were sequenced.
In fact, we recently found that some of these mutants (*i.e.* LMM1010, LMM1020 and DVL1)
incurred loss-of-function mutations in the *crp* or *cyaA* gene, which confers HHP resistance but

361 could still not completely explain their extreme piezoresistance (Gayán et al., 2017b). However,
362 none of these mutants harboured any mutation in *sucABCD* genes, although the fact that the
363 *sucABCD* operon as a whole cannot be compromised could limit the adaptive evolution through
364 loss-of-function mutations in this locus.

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# 367 4. Conclusions

In this study, the screening of a transposon insertion library of E. coli MG1655 for gene 368 369 disruptions affecting HHP inactivation revealed a set of genes and operons (*hdfR*, *rbs* and *suc*) that were previously not recognized to be important for HHP survival. Upon further scrutiny of 370 371 the *sucABCD* operon, we found that *sucA/B* or *sucC/D* deletions had opposing effects on HHP 372 resistance. In fact, compromising SCS functionality by deleting *sucC/D* significantly improved HHP resistance through a mechanism relying on increased RpoS activity rather than an increased 373 374 heat shock (RpoH) or decreased cAMP/CRP response. Although RpoS activity is known to be 375 regulated at many levels, including alterations in carbon metabolism (Battesti et al., 2011, 2015), 376 the input of the SCS complex and/or its metabolic network on quenching RpoS activity was previously unreported and deserves further attention. In contrast, compromising OGDH 377 functionality by deleting *sucA/B* tremendously sensitized cells against HHP, and this phenotype 378 379 could only partially be attributed to decreased RpoS activity levels.

The considerable impact of disruptions of the *hdfR* gene and within the *rbs* and *suc* operons also
underscores a close link between cellular metabolism and HHP susceptibility. In fact, such a link

382	was also recently suggested by directed evolution of an MG1655 $\Delta rpoS$ strain towards HHP
383	resistance, which selected for a number of HHP-resistant mutants harboring loss-of-function
384	mutations in the crp gene (Gayán et al., 2017b). Interestingly, the observed lack of heat (cross-
385	)resistance in the HHP resistant $\Delta cyaA$ or $\Delta crp$ mutants (Gayán et al., 2017b) agrees with the
386	current observation that the HHP resistant $\Delta suc C/D$ mutants did not display an improved heat
387	resistance. This clearly suggests that HHP and heat resistance mechanisms are not necessarily
388	functionally equivalent. In fact, the cellular impact of HHP and heat stress is likely to be very
389	different, as was also recently evidenced by the observation that heat injured E. coli cells proved
390	to be heat sensitive but at the same time remarkably HHP resistant compared to non-injured
391	control cells (Gayán et al., 2016b).
392	Such insights into differential bacterial stress response and adaptation mechanisms will become
393	important for the proper design and prediction of successful hurdle technology approaches.
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395	
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Bacteria and plasmids	Characteristics	Reference
Bacteria		
Escherichia coli		
MG1655 (WT)	Parental wild-type strain	Blattner et al. (1997)
MG1655 $\Delta crl$	MG1655 carrying in frame deletion of <i>crl</i>	This study
MG1655 $\Delta crp$	MG1655 carrying in frame deletion of <i>crp</i>	Gayán et al. (2017b)
MG1655 ΔcyaA	MG1655 carrying in frame deletion of <i>cyaA</i>	Gayán et al. (2017b)
MG1655 $\Delta h df R$	MG1655 carrying in frame deletion of <i>hdfR</i>	This study
MG1655 $\Delta icd$	MG1655 carrying in frame deletion of <i>icd</i>	This study
MG1655 $\Delta lpd$	MG1655 carrying in frame deletion of <i>lpd</i>	This study
MG1655 $\Delta sdhA$	MG1655 carrying in frame deletion of <i>sdhA</i>	This study
MG1655 $\Delta sdhB$	MG1655 carrying in frame deletion of <i>sdhB</i>	This study
MG1655 $\Delta sdhC$	MG1655 carrying in frame deletion of <i>sdhC</i>	This study
MG1655 ΔsdhD	MG1655 carrying in frame deletion of <i>sdhD</i>	This study
MG1655 ΔsucA	MG1655 carrying in frame deletion of <i>sucA</i>	This study
MG1655 $\Delta sucB$	MG1655 carrying in frame deletion of <i>sucB</i>	This study
MG1655 $\Delta sucC$	MG1655 carrying in frame deletion of <i>sucC</i>	This study
MG1655 $\Delta sucC \Delta rpoS$	MG1655 carrying in frame deletion of <i>sucC</i> and <i>rpoS</i>	This study
MG1655 $\Delta sucD$	MG1655 carrying in frame deletion of <i>sucD</i>	This study
MG1655 ΔsucD ΔrpoS	MG1655 carrying in frame deletion of <i>sucD</i> and <i>rpoS</i>	This study
MG1655 $\Delta rbsK$	MG1655 carrying in frame deletion of <i>rbsK</i>	This study
MG1655 $\Delta rbsR$	MG1655 carrying in frame deletion of <i>rbsR</i>	This study
MG1655 $\Delta rpoS$	MG1655 carrying in frame deletion of <i>rpoS</i>	Gayán et al. (2017b)
MG1655 $\Delta rssB$	MG1655 carrying in frame deletion of <i>rssB</i>	Gayán et al. (2017b)
LMM1010	HHP-resistant derivative of MG1655	Hauben et al. (1997)
LMM1020	HHP-resistant derivative of MG1655	Hauben et al. (1997)
LMM1030	HHP-resistant derivative of MG1655	Hauben et al. (1997)
DVL20	HHP-resistant derivative of MG1655	Vanlint et al. (2012)
DVL1	HHP-resistant derivative of MG1655	Vanlint et al. (2011)
Plasmids		
pKD46	Expression of $\gamma$ , $\beta$ and <i>exo</i> recombination genes of phage $\lambda$ under the control of <i>araBAD</i> promoter, temperature-sensitive, Ap <sup>R</sup> .	Datsenko and Wanner (2000)
pKD13	Template plasmid containing <i>kan</i> gene flanked by FRT sites, Ap <sup>R</sup> and Km <sup>R</sup> .	Datsenko and Wanner, (2000)
pCP20	Expression of Flp recombinase, temperature-sensitive, $Ap^{R}$ and $Cm^{R}$ .	Cherepanov and Wackernagel (1995)
pFPV-P <sub>bolA</sub> -gfp	bolA promoter of MG1655 upstream of gfp	Gayán et al. (2016a)
pFPV-P <sub>dnaK</sub> -gfp	dnaK promoter of MG16559upstream of gfp	Aertsen et al. (2004)

**Table 2.** Overview of transposon insertion mutants with altered resistance ( $\geq 2 \log$  cycles in comparison to the WT strain) to a 400 MPa (15 min, 20°C) shock in potassium phosphate buffer (10 mM, pH 7.0), position of the transposon insertion site and corresponding function of the genes affected. Inactivation data correspond to the strains obtained after transduction of the transposon from mutants screened for HHP-resistance change to the WT background.

Mutant	Log reduction	Gene	Transposon position	Function
WT	3.59 (0.16) <sup>a</sup>			
95/1	≥ 6.5*	hdfR	+371	DNA-binding transcriptional dual regulator HdfR (H-NS-dependent <i>flhDC</i> regulator). Repressor for <i>flhDC</i> operon (encoding the master regulator for flagellar biosynthesis and swarming migration) and activator of the <i>gltBDF</i> operon (encoding the glutamate synthase)
101/19	6.38 (0.11) <sup>b</sup>	hdfR	+356	
107/16	6.14 (0.45) <sup>b</sup>	crl	+222	$\sigma^{38}$ (RpoS) RNA polymerase holoenzyme assembly factor Crl
189/16	6.27 (0.25) <sup>b</sup>	crl	+231	
76/37	5.78 (0.49) <sup>b</sup>	rbsK	+748	Ribokinase (involved in ribose catabolism)
31/5	6.04 (0.24) <sup>b</sup>	rbsR	+36	DNA-binding transcriptional dual regulator RbsR. Repressor for <i>rbs</i> operon (involved in ribose transport and catabolism)
126/27	1.93 (0.23) <sup>c</sup>	sucD	+540	Succinyl-CoA synthetase subunit $\alpha$ (TCA cycle)
2/39	1.71 (0.49) <sup>c</sup>	sucC	-102	Succinyl-CoA synthetase subunit $\beta$ (TCA cycle)

The position of the transposon gives the nucleotide after which the transposon was inserted, starting from the first base of the start codon. In the case of *sucC*, the insertion was 102 bp upstream of the *sucC* open reading frame.

Letters indicate statistically significant differences ( $P \le 0.05$ ) among the inactivation of all the strains.

\*Survival counts fell below de quantification limit (1,000 CFU/mL)

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#### 585 Figure legend

**Figure 1.** Logarithmic reduction factor of *E. coli* MG1655 WT and indicated mutants by a HHP treatment at 400 MPa (15 min, 20°C). The dotted line represents the quantification limit (1,000 CFU/ml). Letters indicate statistically significant differences ( $P \le 0.05$ ) among the inactivation of all strains.

Figure 2. β-Galactosidase activity of stationary phase cultures of *E. coli* MG1655 WT and its indicated mutants. Letters indicate statistically significant differences ( $P \le 0.05$ ) among the Miller Units (MU) calculated for each strain.

**Figure 3.** Fluorescence derived from pFPV-P<sub>*dnaK*</sub>-gfp (encoding the *E. coli* MG1655 *dnaK* promoter upstream of gfp) in the indicated strains. Values are expressed as fold change with respect to the average value of the parental strain. Letters indicate statistically significant differences ( $P \le 0.05$ ) among the fluorescence values of all strains.

**Figure 4.** (A) Fluorescence derived from pFPV-P<sub>bolA</sub>-gfp (encoding the *E. coli* MG1655 bolA promoter upstream of gfp) in the indicated strains. Values are expressed as fold change with respect to the average value of the parental strain. (B) Logarithmic reduction factor of *E. coli* MG1655 WT and indicated mutants by a HHP treatment at 300 MPa (15 min, 20°C). The dotted line represents the quantification limit (1,000 CFU/ml). Letters indicate statistically significant differences ( $P \le 0.05$ ) among the inactivation and fluorescence values of all strains.

**Figure 5.** Resistance of *E. coli* MG1655 WT and indicated mutants to (A) hydrogen peroxide

and (B) heat. The resistance to hydrogen peroxide was measured by the inhibition growth

diameter of a 5-µl aliquot of the chemical (30% w/w) on stationary phase cells plated in 15 ml of

606	LB soft agar, while heat resistance was assayed by the logarithmic reduction factor by a heat
607	treatment at 57.0°C for 15 min. The dotted line represents the quantification limit (1,000
608	CFU/ml). Letters indicate statistically significant differences ( $P \le 0.05$ ) among the inhibition
609	growth diameter and inactivation of all the strains.
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