

# Quantifying the effects of heating temperature, and combined effects of heating medium pH and recovery medium pH on the heat resistance of Salmonella typhimurium

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2	Quantifying the effects of heating temperature, and combined effects of heating medium
3	pH and recovery medium pH on the heat resistance of Salmonella typhimurium
4	
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6	
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12	Abstract
13	The influence of heating treatment temperature, pH of heating and recovery medium on the
14	survival kinetics of Salmonella typhimurium ATCC 13311 are studied and quantified. From
15	each non log linear survival curve, Weibull model parameters were estimated. An average
16	shape parameter value of 1.67 was found, which is characteristic of downward concavity
17	curves and is in agreement with values estimated from other Salmonella typhimurium strains.
18	Bigelow type models quantifying the heating temperature, heating and recovery medium pH
19	influences, are fitted on scale parameter $\delta$ data, (time of first decimal reduction), which
20	reflects the bacterial heat resistance. The estimate of $z_T$ (4.64°C) is in the range of values
21	given in literature for this species. The influence of pH of the heating medium on the scale
22	parameter ( $z_{pH}$ : 8.25) is lower than that of the recovery pH medium influence ( $z'_{pH}$ : 3.65)
23	Key words: Weibull, heating medium, recovery medium, pH, Salmonella typhimurium

#### 24 Nomenclature

- $N_{0:}$  initial number of cells.
- *N*: number of surviving cells after the heating time t.
- $\delta$ : scale parameter : first decimal reduction of surviving spores or cells from  $N_0$  to  $N_0/10$ .
- *p:* shape parameter.
- *T*: heating temperature.
- pH: heating medium pH.
- pH': recovery medium pH.
- $T^*$ : reference temperature fixed at 60°C.
- $pH^*$ : reference pH of the heating medium fixed to 7.
- $PH'_{opt}$ : recovery medium: pH corresponding to the maximal apparent bacterial heat resistance.
- $z_T$ : distance of T from  $T^*$  which leads to a ten fold reduction in  $\delta$ -value  $z_T$  quantifies the
- 36 influence of the heating temperature on the bacterial heat resistance.
- $z_{pH}$ : distance of pH from  $pH^*$  which leads to a ten fold reduction in  $\delta$ -value  $z_{pH}$  quantifies the
- influence of the pH of the heating medium on the bacterial heat resistance.
- $z'_{pH}$ : distance of pH' from  $pH'_{opt}$ , which leads a ten fold reduction in apparent  $\delta$  -value.  $z'_{pH}$ 40 characterises the influence of the pH on the recovery of the micro-organism after a heat 41 treatment.
- $\delta^*$  :estimated  $\delta$  value corresponding to T\*, *pH*\* and *pH*'<sub>opt</sub> conditions.

#### 45 Introduction

Salmonellae continue to be a major public health setting a great problem to the food industry. 46 These bacterial species appear in a wide variety of foods and food ingredients. Various heat 47 treatments implemented by the food processing industry are generally effective for destroying 48 the vegetative bacteria. Taking into account not only temperature but other environmental 49 factors is likely to allow significant reduction of heat treatment intensity with the same 50 microbial safety and minimize damage of heat sensitive food. It is commonly agreed that the 51 pH decrease of the heating medium is the main environmental factor after temperature, which 52 53 reduces the bacterial heat resistance of spores (Bigelow and Esty, 1920) or vegetative form (White, 1963; Blackbrun, Curtis, Humpheson, Billon & McClure, 1997). The same effect was 54 observed for Salmonella enteritidis, (Casadei, Ingram, Hitchings, Archer & Gaze, 2001) and 55 56 for Salmonella typhimurium (Mazzotta, 2001).

Moreover the pH of the recovery medium highly influences the apparent heat resistance of 57 bacterial spores for the same heat treatment condition. The D values decrease when pH values 58 shift from an optimum (Cook and Brown, 1965). However, as far as we know, the influence 59 of the recovery medium pH on the apparent heat resistance of Salmonella species has never 60 been described. A decrease of pH of the heating medium or of the recovery medium both 61 reduces the bacterial heat resistance (Couvert, Leguérinel & Mafart, 1999). Theses influences 62 can be taken into account for reducing heat treatments. Such a cumulative effect is more or 63 64 less specific of pH and is not observed for water activity where, oppositely, the protective effect of a low water activity of the heating medium tends to be balanced by the selectivity of 65 a low water activity of the recovery medium towards injured cells (Coroller, Leguérinel & 66 67 Mafart, 2001).

68 The aim of this study was to quantitatively characterise the impact of heating temperature, pH69 of the heating medium and pH of the recovery medium on the heat resistance of *Salmonella*70 *typhimurium* from relevant parameters of appropriated primary and secondary models

71

#### 72 Materials and methods

#### 73 Strain and culturing conditions

The studied strain was *Salmonella typhimurium* ATCC 13311 (NCTC 74). Cultures were stored in cryotube in mixing nutrient broth 50% glycerol 50% at -70°C.

The basic heating medium was tryptone salt broth (10g/l tryptone USP (Biokar Diagnostics, A1401HA) and 10g/l NaCl). The pH was adjusted with addition of  $H_2SO_4$  and sterilised by filtration through 0.22µm porosity filter.

The recovery medium was nutrient agar (Biokar Diagnostics, BK021HA). The pH were adjusted with H<sub>2</sub>SO<sub>4</sub> prior autoclaving at 121°C for 15 minutes. The pH values were checked after autoclaving.

82 Preparation of cells suspension

Nutrient broth (Biokar Diagnostics, BK003HA) in 200ml flask was inoculated with *Salmonella typhimurium* and incubated at 37°C for 24 hours under agitation (150 rpm). The culture (40ml) was centrifuged (2000g 15min at 20°C) and re-suspended in 3 ml of heating medium.

87 Thermal treatment of bacterial suspension and recovery conditions

Capillary tubes of 200 µl (Ringcaps® Duran®) were filled with 100µl of sample and submitted to a thermal treatment in a thermostated water bath. After heating, the tubes were cooled in water/ice bath. After rinsing, the ends were flamed with ethanol. The capillary tubes were broken at both ends and their contents poured into a tube containing 9 ml sterile tryptone salt broth (Biokar Diagnostics, BK014HA) by rinsing with 0.9 ml tryptone salt broth. 93 Viable cells were counted by duplicate plating in ajusted pH nutrient agar (Biokar
94 Diagnostics, BK021HA) and incubated at 37°C for 48h.

95 Experimental design

96 To determine survival kinetic parameters, bacteria, for each sample corresponding to different
97 heating times, were counted on nutrient agar plates.

- Heating temperatures applied were 53, 55, 57 and 59°C (heating and recovery media pH equal
  7). For studying the effect of pH, a complete factorial design was implemented according to
  the following levels of pH of the heating medium: (7, 6.5, 6, 5.3, 5, 4.4 and 3.8) and to the
  following levels of pH of the recovery medium (denoted pH'):7, 6.5, 6, 5.5 and 5 (controlled
  pH values are given table3) at 55°C.
- 103 *Data from literature*
- Data taken from figures in literature were scanned and digitized using the software program
  DigXY 1.2 (Thunderhead Engineering, Manhattan, USA).
- 106 Primary and secondary models

Different authors considered the survival curve as a cumulative form of temporary
distribution of lethality event frequency (Cunhan, Oliveira & Oliveira, 1998; Peleg & Cole,
109 1998; Fernandez, Salmeron, Fernandez & Martinez, 1999). In 2002, Mafart, Couvert,
Gaillard, and Leguerinel proposed a new presentation of the Weibull frequency distribution
model.

112 
$$\log N = \log N_0 - \left(\frac{t}{\delta}\right)^p \operatorname{Eq1}$$

Parameter *p* characterises the shape of the curve: concave curves p<1, convex curves p>1 and linear curves p=1, in this case  $\delta$  value corresponds to classical *D* value. This equation was taken up during the IFT summit in January 2003 (Heldman and Newsome, 2003), was used by different authors (Mafart et al., 2002; Gómez, García, Álvarez, Raso & Condón, 2005; Geeraerd, Valdramidis & Van Impe, 2005; Virto, Sanz, Álvarez, Condón & Raso, 2005;
Carlin et al., 2006) and was implemented in this work as primary model.

The influences of environmental factors on the Weibull model parameter estimated from survival kinetics related to bacterial spores or vegetative cells, were studied (Fernandez et al., 1999; Fernandez, Collado, Cunhan, Ocio & Martinez, 2002; van Boekel, 2002; Couvert, Gaillard, Savy, Mafart & Leguerinel 2005).These studies showed that the shape parameter was practically independent of the heating temperature and the pH of the heating medium. The conclusion of these studies leads us to determine a single average shape parameter value for a set of kinetics.

126 The  $\delta$  value, first decimal reduction time, is highly influenced by heating temperature. The 127 classical Bigelow model was used to describe the influence of heating temperature on the  $\delta$ 128 with the conventional  $z_T$  value.

The effect of the pH of the heating medium and the pH' of the recovery medium on the heat
resistance was described according to the following equation (Leguérinel, Spegagne, Couvert,
Gaillard & Mafart, 2005) :

132 
$$\log \delta = \log \delta^* - \left| \frac{pH - pH^*}{z_{pH}} \right| - \left( \frac{pH' - pH'_{opt}}{z'_{pH}} \right)^2 \text{ Eq } 2$$

133 *Curve fitting*.

134 In a first time,  $N_0$ ,  $\delta$  and p values are estimated from each survival curve to assess the 135 influence of environmental factors on these parameters.

136 In a second time a single shape parameter p value was estimated from the corresponding 137 whole set of experimental kinetics and from set of kinetics taken from the literature. Scale 138 parameter  $\delta$  values and logN<sub>0</sub> values were determined for each curve.

Bigelow parameter  $z_T$  was estimated from scale parameter  $\delta$  values obtained from the temperature mono factorial design. Equation 2 parameter values:  $z_{pH}$  and  $z'_{pH}$  were estimated from scale parameter  $\delta$  values obtained from pH factorial design.

143 The parameter values and their associated confidence interval were estimated by using a non-144 linear module (nlinfit and nlparci Matlab 6.1, Statistical Toolbox, The Mathworks).

145

#### 146 **Results and discussion**

Survival kinetics curves of *Salmonella typhimurium* showed a clear downward concavity. The
same pattern of curves was elsewhere observed for different *Salmonella typhimurium* strains
(Garibaldi, Ljichi & Bayne, 1969; Mackey and Derrick, 1986; Jäckle, Geiges & SchmidtLorenz, 1987). Such non linear curves were fitted according to the Weibull model.

151 The influences of the heating temperature, the pH of the heating and of the recovery medium on the shape parameter p are shown Figure 1. p values appear to be not clearly influenced by 152 these environmental factors. This observation is in agreement with those of Fernandez et al. 153 154 (2002), Collado, Fernandez, Rodrigo, Camats and Martinez Lopez. (2003) regarding Bacillus 155 cereus, Couvert et al. (2005) and Bacillus pumilus and van Boekel's (2002). These authors did not observe any significant influence of the temperature on the shape parameter p. The 156 presented results (Figure 1) show no clear influence of the heating medium or the recovery pH 157 on estimates of p for the lower pH. However structural correlation between parameters p and 158  $\delta$  could explain the variability of p values (Couvert et al., 2005). 159

160 Then, a single average p value was estimated, regardless of the heating temperature, the pH of 161 the heating and of the recovery medium. The three parameters ( $N_0$ ,  $\delta$  and p) were globally 162 estimated from the whole set of data by using the least square regression method (nlinfit 163 Matlab 6.1). The single p value estimated from our set of data (1.677 ± 0.065) is close to the p 164 value estimated from other sets of data from literature for the same Salmonellae species:

 $1.648 \pm 0.313$  (Jäckle *et al.*, 1987),  $1.538 \pm 0.187$  (Garibaldi *et al.*, 1969),  $1.429 \pm 0.295$ 165 (Mackey & Derrick, 1986). log  $N_0$ , the scale parameters  $\delta$  and their confidence interval 166 coefficients as functions of different pH and temperature conditions, are presented in Table 1 167 and 2. The scale parameter  $\delta$  values are influenced by environmental factors: heating 168 temperature, pH of the heating and the recovery medium. Within the investigated temperature 169 range (53°C -59°C) the classical Bigelow relationship was kept to quantify the effect of 170 temperature on  $\delta$  value. The corresponding  $z_T$  value 4.64°C (Table 3) is lower than  $z_T$  values 171 172 determined from Jäckle et al. (1987) data for the same bacterial strain but are in agreement with  $z_T$  value estimated from Mackey and Derrick data (1986) and with other values found in 173 literature. Doyle and Mazzotta (2000) reviewed  $z_T$  values concerning different Salmonella 174 175 *typhimurium* strains heated in different media. These  $z_T$  values, ranging from 3.24°C to 9.5°C with a mean of 5.56°C, illustrates the large variability of z<sub>T</sub> values reported in literature. For 176 the same Salmonella typhimurium strain ATCC13311 Casadei et al. (2001) reported a z<sub>T</sub>-177 value of 4.6°C (Table 4). 178

Both the pH of the heating and the recovery medium affect the heat resistance of *Salmonella typhimurium* as shown Figure 2. A decrease of the pH of the heating medium reduces the salmonella heat resistance. This observation has been reported by Blackburn *et al.* (1997) and Casadei *et al.* (2001). Concerning the influence of the recovery medium, low pH reduces heat resistance parameter value  $\delta$  of *Salmonella typhimurium*. A similar effect was observed for the thermal inactivation of bacterial spore (Cook and Brown, 1965; Lopez, Gonzalez, Mazas, Gonzalez, Martin & Bernardo, 1997) but never for vegetative bacteria cells.

186 Equation 2 was used to describe the effect of the heating medium pH and the recovery 187 medium pH on  $\delta$  values. This model does not take interactions between these two 188 environmental factors into account. To estimate the weight of the possible interaction, a 189 variance analysis was performed. For the studied pH range, the weight of heating and 190 recovery medium pH represent 62.9% and 34.7% respectively, while the unexplained 191 variance comprising interactions represented only 2.4% of the total variability. This 192 observation concerning *Salmonella typhimurium* is in agreement with Couvert et al. (1999) 193 results related to *Bacillus cereus* spores. Such observations led us to neglect interactions and 194 to retain the simple Equation 2 without crossed term. Model parameters (Eq 2) determined for 195 *Salmonella typhimurium* (Table 5) were estimated from the whole set of  $\delta$  values, according 196 to factorial design.

The determination coefficient between experimental and calculated values ( $R^2$ :0.958) and 197 Figures 3a and 3b illustrate the goodness of fit of the model. The high  $z_{pH}$  parameter value 198 8.25, indicates a poor influence of the heating medium pH on the heat resistance for this 199 Salmonella strain. For other salmonella strains few data allow to evaluate this  $z_{pH}$  parameter. 200 From Blackburn et al. (1997), Casadei et al. (2001) and Mañas, Pagán, Raso and Condón 201 (2003) data, an optimal pH value appears to be variable and often lower than 7. This 202 observation leads to replace a fixed reference  $pH^*$ , equal to 7, by a variable optimal pH 203 parameter  $(pH_{opt})$  to be estimated Eq 3. 204

$$\log \delta = \log \delta_{opt} - \left(\frac{T - T^*}{z_T}\right) - \left|\frac{pH - pH_{opt}}{z_{pH}}\right| \text{ Eq 3}$$

Parameters  $z_{pH}$ ,  $pH_{opt}$  and  $z_T$  estimated from published values are shown Table 4. Regarding the studied bacterial strain and different bacterial species,  $z_{pH}$  values estimated from data of literature and our own data are lower than  $z_{pH}$  obtained in this work, even for the same *Salmonella typhimurium* strain ATCC 11331. This difference could be explained by errors on D value estimates due to the fitting of concave curves by linear regressions or growth condition and physiology state of bacteria which can have produced stress protein.

Concerning the effect of the pH of the recovery medium, the  $z'_{pH}$  value (3.6) indicates a higher influence of this factor than that of the heating medium pH. It is generally accepted that the pH of the recovery medium exerts a large influence on the apparent heat resistance of spores: *D*-values decrease as pH is reduced (Cook and Brown, 1965; Yokoya and York, 1965; Cook and Gilbert, 1968; Mallidis and Scholefield, 1986; Santos and Zarzo, 1996; Lopez et al., 1997). Observed  $z'_{pH}$  values, which characterized the influence of the recovery medium pH, for *Salmonella typhimurium* cannot be compared with values regarding other Salmonella strains or species. As far as we know, no data which could be compared to our results, are available from literature.

The primary model derived from the Weibull distribution and describing non linear survival kinetics associated with Bigelow type secondary models, can be easily used to optimise heat treatment process calculations taking the heating and recovery pH influences into account. For example, compared to a heat treatment in food at pH 7, a heat treatment for *Salmonella typhimurium* in food at pH 5, could reduce the heating time to a 3.5 ratio, or, with the same heating time, could reduce the heating temperature of 2.25°C with the same lethal efficiency.

In practice, to ensure safety of acid foods, heating pH is the pH of food before heat treatment. Because foods represent both the heating and the recovery medium, the input recovery pH is likely to keep the value of the heating pH. However, it frequently occurs that a decrease of food pH is observed during the heat treatment. In this case, for safety reasons, it is recommended to retain the value of the pH which is measured immediately after the heat treatment.

This work confirms the impact of low recovery medium pH on the apparent heat resistance. This influence allows reducing heat treatments with the same safety objective to keep better nutritional and sensory quality of foods. This approach could be extended to other vegetative strains and species which would require further data: p and z parameter values, related to heating temperatures and heating and recovery medium pH. From these data, standard values of  $z_{pH}$  and  $z'_{pH}$  could be defined according to a similar approach to the one that was

- implemented for the standard  $z_T$  value equal to 7°C for most resistant vegetative cells and
- 240 input in pasteurisation process calculations.

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- 338

339	Table legends
340	Table1
341	log $N_0$ and $\delta$ estimates and their confidence interval coefficients as functions of the heating
342	temperature.
343	
344	Table 2
345	log $N_0$ and $\delta$ estimates and their confidence interval coefficients as functions of the heating
346	and the recovery medium pH.
347	
348	Table 3
349	Parameters $z_T$ estimated (from log $\delta$ values determined) from our own (data) and published
350	data for different Salmonella strains
351	
352	Table 4
353	$z_{T, z_{pH}}$ and $pH_{opt}$ estimates from published classical D values for different Salmonella species
354	
355	Table 5
356	$z_{pH}$ , $z'_{pH}$ , $pH'_{opt}$ estimates for Salmonella Typhimurium ATCC 13311 (log $\delta$ values)

358	Figure legends
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359 Figure 1

- 360 Weibull shape parameter values and their confidence interval 95% evaluated for each kinetic
- 361 of heating temperature, and heating and recovery medium pH
- 362
- 363 Figure 2
- 364 Survival kinetics experimental data and fitted curves, with p-value equal1.67, at different
- heating medium pH for different recovery pH (pH 7  $\bullet$ , pH 6.5  $\Box$ , pH6 $\blacktriangle$ , pH5.5  $\bigtriangledown$ , pH 5  $\blacklozenge$ )
- and at different temperatures (59°C7  $\bullet$ , 57°C  $\Box$ , 55°C  $\blacktriangle$ , 53°C  $\bigtriangledown$ ) heating and recovery pH
- 367 fixed at 7
- 368
- 369 Figure 3a 3b
- 370 Observed and calculated log  $\delta$  values for different conditions of heating and recovery medium
- 371 pH
- 372
- 373
- 374











383 figure 3a 



385
386
387 Figure 3 b
388

Heating				
temperature °C	$\log N_0$	CI 95%	$\delta$ minutes	CI 95%
59	7.730	± 0.385	0.619	$\pm 0.047$
57	7.522	± 0.352	2.495	± 0.182
55	7.837	± 0.297	5.846	± 0.690
53	8.077	± 0.315	12.741	±1.547

392 Table 1 

		Global p va	lue	1.677	±0.065
Heating	Recovery	logNo	CL 05%	$\delta$ minutes	CL 05%
pН	pН	105110	01 95%	0 minutes	01 95%
7.02	7	8.155	±0.244	6.734	±0.475
7.02	6.5	7.925	±0.244	6.922	±0.499
7.02	6	7.812	±0.257	6.010	±0.447
7.02	5.45	7.872	±0.281	4.949	±0.380
7.02	4.92	7.808	±0.302	3.696	±0.258
6.5	7	8.295	±0.257	5.956	±0.440
6.5	6.35	8.197	±0.257	6.059	±0.454
6.5	6	8.067	±0.257	6.178	±0.472
6.5	5.45	7.998	±0.281	5.063	±0.397
6.5	4.95	8.191	±0.321	3.276	±0.236
6	7	8.356	±0.270	5.657	±0.370
6	6.45	8.626	±0.281	5.041	±0.394
6	6	8.489	±0.284	4.683	±0.367
6	5.45	8.261	±0.323	3.760	±0.277
6	5	8.570	±0.352	2.483	±0.181
5.3	7	8.599	±0.283	4.458	±0.314
5.3	6.35	8.559	±0.282	4.506	±0.319
5.3	6	8.514	±0.299	4.273	±0.335
5.3	5.45	8.425	±0.320	3.513	±0.268
5.3	4.92	8.066	±0.336	2.731	±0.191
5	7	8.659	±0.300	4.107	±0.310
5	6.35	8.568	±0.302	3.728	±0.261
5	6	8.613	±0.319	3.857	±0.324
5	5.45	8.684	±0.322	3.047	±0.209
5	4.95	7.990	±0.356	2.628	±0.201
4.4	7	8.414	±0.321	3.171	±0.223
4.4	6.45	8.392	±0.321	3.338	±0.244
4.4	6	8.400	±0.322	3.114	±0.217
4.4	5.35	8.488	±0.352	2.458	±0.178
4.4	4.95	8.548	±0.394	1.836	±0.141
3.8	7	8.117	±0.322	3.170	±0.226
3.8	6.5	8.152	±0.351	2.722	±0.214
3.8	6	8.045	±0.398	2.281	±0.182
3.8	5.5	8.402	±0.394	1.935	±0.155
3.8	5	8.264	±0.455	1.611	±0.183

401 Table2

	Bacterial strain		Heating medium	n*	T°C range	$z_T^{\circ}C$	CI95 %	R²
	Salmonella typhimurium ATCC13311		Tryptone salt broth	4	51-59	4.64	±0.877	0.989
	Salmonella typhimurium ATCC13311	Jäckle <i>et al</i> . 1987	Tryptone salt broth	3	58.5-61.5	8.55	±1.796	1.000
	Salmonella typhimurium NCBI 10248	Mackey & Derrick 1986	Tryptone salt broth	5	50-59	3.44	±0.510	0.994
403 404 405	* n: data number Table3							

Bacterial strain		Heating medium	n	T°C range	$z_T^{\circ}C$	CI 95%	pH range	Z <sub>p</sub> H	CI 95%	$pH_{opt}$	R²
Salmonella typhimurium ATCC13311	Manas <i>et al.</i> 2003	Citrate buffer	8				4-7.7	3.27	0.969	5.75	0.939
Salmonella enteritidis P167807	Blackburn <i>et al</i> . 1997	Tryptone salt broth + HCl /NaOH	8				4.3-9.5	3.92	2.261	6.00	0.724
Salmonella typhimurium NCTC 74	Casadei <i>et al.</i> 2001	Nutrient broth + Citric acid	8	48-54	4.61	1.204	7 & 3	2.17	0.203	ND Fixed at 7	0.994
* n: data number	2001										

417 Table 4

		Parameter	CI 95%
		values	
	$\log \delta^{*}$ 55°C, pH7, pH'opt	0.851	±0.053
	$Z_{pH}$	8.254	±1.572
	$z'_{pH}$	3.655	$\pm 1.349$
	$pH'_{opt}$	6.805	$\pm 0.656$
	R <sup>2</sup>	0.958	
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426	Table 5		
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