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1. Introduction

With increased applications of electromagnetic waves in many aspects of our daily life, determining the effects of microwave irradiation on biological systems is not only essential, but far from fully elucidated. Notions of 'thermal' and 'non-thermal' effects (the latter is also referred to as 'microwave-specific effects') of microwaves have been under repeated scrutiny in the past few decades.^{1–10} Some recent literature argued that due to the unique manner of energy transfer by microwaves, many observations that are unique to microwave exposure can be attributed to 'selective heating' of substances in a system of interest.^{11,12} As a result, isolation of thermal from microwavespecific effects is challenging,^{13–15} further complicating the interpretation of experimental results.

In this respect, we previously demonstrated that a CEM (Matthews, NC) microwave reactor operating at 2.45 GHz coupled with a Coolmate system allows for systems under

Exposure to microwave irradiation at constant culture temperature slows the growth of *Escherichia coli* DE3 cells, leading to modified proteomic profiles[†]

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Despite a few decades of research, interest continues in understanding the potential influences of low energy microwave irradiation on biological systems. In the present study, growth of *E. coli* DE3 in LB media slowed in the presence of microwave irradiation (max. 10 W) while the temperature of cultures was maintained at 37 °C. Viable cell counts in microwave-irradiated cultures were also significantly lower. When microwave irradiation was ceased, *E. coli* growth was restored. A top-down proteomic analysis of total proteins isolated from control and microwave-irradiated *E. coli* cultures revealed differential abundance of 10 resolved protein spots, with multiple proteins identified in each following mass spectrometric analysis. Among these proteins, a number are involved in metabolism, suggesting alterations to metabolic activities following microwave irradiation. Furthermore, four amino acid-tRNA ligases were also identified, pointing to the possibility of stress responses in *E. coli* under microwave irradiation.

investigation to remain typically within ±1 °C throughout experiments (with occasional brief fluctuations of ± 2 °C), as measured in situ by a fibre optic temperature probe and an alcohol-based thermometer, when the system is subjected to low-power microwave irradiation while cooled simultaneously. We were particularly interested in studies in which relatively low microwave energy (up to 10 W) is used, as higher microwave output is, presumably, more likely to exacerbate the issue of local heating and generation of hotspots. With these considerations, we demonstrated that while the enzymatic activity of trypsin during digestion of substrates, such as casein, is increased by exposure to microwave irradiation (10 W) at constant bulk temperature,16 a similar impact was not found for α-amylase and phosphatase.¹⁷ We further demonstrated that exposure of PC-3 human prostate cancer cells to non-lethal microwave irradiation (10 W) at constant culture temperature (37 °C) likely led to perturbation of cell membrane properties, while no necrosis nor apoptosis was induced.18 Most recently, we showed that uptake of the anticancer compound doxorubicin by PC-3 and human breast cancer MCF-7 cells was significantly enhanced when the cells were exposed to nonlethal microwave irradiation.19

In the present study, we extended our investigation to probe the influence of microwave exposure on bacterial growth while the culture temperature was maintained constant. While many reports show no differences in microorganisms exposed to microwave irradiation as compared with controls,²⁰⁻²⁶ there is

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also literature showing that microorganisms respond differently to microwave irradiation of various frequencies, leading to altered growth rate or metabolic activities.^{21,27-36} Comparisons of results from the literature are compounded by differences in at least the following factors: frequency and power output of the microwave used, duration and pattern of irradiation, choice of biological system, method used in temperature measurement, and the extent of temperature fluctuation.

This study focused on the exposure of *E. coli* DE3 cells to microwaves at 2.45 GHz, with relatively low power output (up to 10 W), while the culture temperatures were monitored *in situ* with a fibre optic temperature probe, maintaining the culture temperatures at 37 °C through simultaneous cooling. We report herein that *E. coli* growth is slowed under these conditions. Furthermore, a top-down proteomic analysis of total proteins suggest that cellular metabolism is modified in bacteria exposed to microwave irradiation.

2. Results and discussions

2.1. Temperature control

In the present study, great care was taken to ensure that the temperature of the bacterial cultures was controlled through simultaneous cooling and constant stirring; cultures were kept sterile with the use of a foam plug (Fig. 1). Pre-cooled fluid (Galden HT 110) was circulated through the jacket surrounding the culture tube in order to maintain culture temperatures. As we have previously established,¹⁷ the temperatures measured by a fibre optic temperature probe and an alcohol-based thermometer are very consistent. For simplicity, a fibre optic temperature probe was used here, as it allowed us to readily keep track of culture temperatures in real-time using the Synergy program. As is shown in Fig. S1,† temperatures of the culture were maintained at 37 °C (typically ± 1 °C with occasional brief fluctuations of ± 2 °C) throughout the 27.5 h growth period.

It is worth noting that as the fibre optic temperature probe provides feedback to the microwave, the microwave power output from the magnetron was not constant. As shown in Fig. S2,† while the maximal power output was set at 10 W, actual power output was below 10 W (>6 W for most of the irradiation period), but did not exceed this pre-set maximal value.



Fig. 1 E. coli culture setup in a CEM microwave reactor.



Fig. 2 Growth curves of *E. coli.* ■: cultured at 37 °C in a shaking incubator at 260 rpm; ●: cultured with stirring at 37 °C in a tube in the presence of microwave irradiation; ▲: cultured with stirring in a tube at 37 °C in an oil-bath in the absence of microwave exposure. All experiments were an average of biological triplicates. **P* < 0.05; ***P* < 0.01 calculated for OD₆₀₀ of cultures treated with microwave and control in oil bath.

2.2. E. coli growth curve

As shown in Fig. 2, the growth rates of *E. coli*, both in the presence and absence of microwave exposure, were slower than that in a shaking incubator, which is the result of less efficient oxygen transfer in the culture tubes under stirring compared with cultures in a shaking incubator. The growth of *E. coli* in the presence of maximal 10 W microwave irradiation at 37 °C was found to be significantly slower than control cultures not exposed to microwave irradiation.

2.3. Dissolved oxygen (DO) level in tube cultures

Considering the possibility that the slower growth of *E. coli* in the presence of microwave irradiation might be due to lower DO concentration when cultures are exposed to microwave irradiation, experiments were carried out to compare the DO levels in tube cultures isolated from the environment with foam plugs, in the presence and absence (control) of microwave irradiation.

Table S1[†] shows that the DO levels are slightly lower in the LB media treated with microwave irradiation relative to the control. This observation is consistent with a recent study that demonstrated efficient degassing by microwave exposure, however, at much higher temperatures.³⁷

2.4. Recovery of growth in the *E. coli* culture exposed to microwave irradiation

We were also interested in exploring whether the slower growth of *E. coli* subjected to microwave irradiation is reversible when microwave exposure stops. To this end, after *E. coli* cultures were exposed to maximal 10 W microwave irradiation at 37 °C for 5 h, the cultures were then incubated at 37 °C in an oil-bath, and growth rate (measured by OD_{600}) was assessed. As depicted in Fig. 3, the difference in cell density between the microwave treated and control cultures decreased when microwave irradiation was stopped. Viability tests revealed that 21 h after microwave exposure ceased, the difference in viable cell counts in both cultures was not significant (Fig. 4).



Fig. 3 Recovery of *E. coli* growth after ceasing microwave exposure at 5 h as indicated by the decreased difference in OD₆₀₀. Both control and microwave-treated cultures were carried out in biological triplicates.



Fig. 4 Viable cell counts 21 h after microwave exposure as compared with control. Graph shows mean \pm std. dev.

2.5. Two-dimensional gel electrophoresis (2DE) of total proteins isolated from *E. coli* DE3 cultures

Studies have shown that physical phenomenon such as electromagnetic waves can act as stimuli that may trigger responses in biological systems.^{38,39} Based on the current observations that *E. coli* cells grew more slowly in the presence of microwave irradiation, we became interested in studying the protein profiles, which could provide insight into potentially altered biological processes in *E. coli* under microwave exposure.

In order to investigate this scenario, total proteins were isolated from both control and microwave-treated *E. coli* cultures for 2DE-based proteomic analysis. Proteins from three biological replicates from both control and microwave-treated *E. coli* cultures were extracted and analyzed by 2DE in technical triplicates (all gel images are available in Fig. S3[†]).

Changes in spot densities between the resolved proteomes of microwave-treated *vs.* control *E. coli* were identified by quantitative image analysis. Here, 10 spots were identified as changing significantly, with three decreasing and seven increasing in abundance (Fig. 5).

2.6. Mass spectrometric (MS) analysis

Using the criteria described in the Experimental section, within these 10 resolved spots a total of 42 proteins were identified by comparing the MS data against known *E. coli* K12 proteins (Table 1). These proteins were grouped into two clusters according to the spot intensity seen in the 2D gels, one containing 16 proteins with overall decreased abundance and another containing 26 proteins with overall increased abundance in the microwave-treated cultures, respectively. Fig. S4† shows the LC-MS chromatograms of the 10 resolved protein spots.

Analysis of the proteins with decreased levels revealed that many proteins are involved in metabolic pathways, including the citric acid cycle, such as aconitate hydratase B, pyruvate dehydrogenase E1 component, aldehyde–alcohol dehydrogenase, 2-oxoglutarate dehydrogenase E1 component, NADH-





Table 1	Summary of the 42 proteins identified by mass spectrometry. The molecular weight (MW) and isoelectric point (pl) values determined by
2DE <i>vs.</i>	theoretical are presented for comparison

Protein spot	Abundance in irradiated <i>E. coli</i> [fold-change (<i>p</i> -value ^{<i>a</i>})]	Protein ID	Accession	Coverage (%)	No. of unique peptides	Theoretical MW [kDa]/pI	Calculated ^b MW [kDa]/pI	Sequest score
1	↓1.3 (0.004)	Aconitate hydratase B	P36683	44	38	93.4/5.4	83.4/4.9	846.66
		Pyruvate dehydrogenase E1 component	P0AFG8	25	22	99.6/5.7		85.02
		Aldehyde–alcohol dehydrogenase	P0A9Q7	17	12	96.1/6.8		71.4
		2-Oxoglutarate dehydrogenase E1 component	P0AFG3	12	9	105/6.5		23.09
		NADH-quinone oxidoreductase subunit G	P33602	9	9	100.2/6.3		22.31
		Leucine-tRNA ligase	P07813	9	7	97.2/5.3		17.1
		Elongation factor G	P0A6M8	8	4	77.5/5.4		9.61
		DNA mismatch repair protein MutS	P23909	3	2	95.2/5.55		4.3
		DNA gyrase subunit A	P0AES4	2	2	96.9/5.2		4.22
2	↑3.0 (<0.001)	GTP-binding protein TypA/BipA	P32132	13	6	67.3/5.3	73.4/4.8	5.33
		Catalase-peroxidase	P13029	8	5	80/5.3		5.31
		Formate acetyltransferase 1	P09373	4	3	85.3/6.0		6.01
		Polyribonucleotide nucleotidyltransferase	P05055	3	2	77.1/5.2		5.21
3	$\uparrow 2.0 \ (<0.001)$	Formate acetyltransferase 1	P09373	16	11	85.3/6.0	72.7/5.3	6.01
		Phosphoenolpyruvate-dependent phosphotransferase system	P37177	3	2	83.7/5.8		5.78
4	1.6 (<0.001)	Exoribonuclease 2	P30850	14	7	72.4/5.6	70.6/5.2	5.62
	,	Chaperone protein ClpB	P63284	7	5	95.5/5.5		5.52
		Dihydrolipoyllysine-residue acetyltransferase	P06959	5	3	66.1/5.2		5.17
		Glycine-tRNA ligase beta subunit	D00061	2	2	76 8/5 4		5.44
		Rifunctional polymyrin resistance protein ArnA	P00901	2	2	70.8/3.4		5.44 6.97
5	↑1 4 (0 001)	Chaperone protein HtpG	P01673	3 40	23	74.2/0.9	67 7/4 7	5.21
5	114 (0.001)	Lycine-tPNA licese heat inducible	DOAGNE	4J 26	10	57 9/5 2	07.7/4.7	5.21
		Dibydrolinovllysine-residue acetyltransferase	P06050	20	12	57.8/5.2		5.17
		component of pyruvate dehydrogenase complex	F00939	14	1	00.1/3.2		5.17
		Proline-tRNA ligase	P16659	8	4	63.7/5.2		5.24
		30S ribosomal protein S1	P0AG67	6	4	61.1/5.0		4.98
		Polyribonucleotide nucleotidyltransferase	P05055	3	2	77.1/5.2		5.21
		Methionine-tRNA ligase	P00959	3	2	76.2/5.9		5.86
6	$\downarrow 1.5 (< 0.001)$	D-Tagatose-1,6-bisphosphate aldolase subunit GatZ	P0C8J8	18	7	47.1/5.8	53.7/5.2	5.77
		Periplasmic pH-dependent serine endoprotease DegQ	P39099	8	3	47.2/6.0		5.95
		Dihydrofolate synthase/folylpolyglutamate synthase	P08192	5	2	45.4/5.8		5.8
		Xaa-Pro aminopeptidase	P15034	6	2	49.8/5.4		5.39
		Bifunctional protein HldE	P76658	5	2	51.0/5.4		5.41
7	\uparrow 1.5 (0.006)	NADH-quinone oxidoreductase subunit F	P31979	12	5	49.3/6.9	49.9/6.6	6.86
8	\uparrow 1.9 (<0.001)	Aspartate aminotransferase	P00509	17	7	43.5/5.8	43.3/5.3	5.77
		NADPH-dependent curcumin reductase	P76113	7	2	37.6/5.8		5.82
		Aminomethyltransferase	P27248	5	2	40.1/5.6		5.57
		Maltose-binding periplasmic protein	P0AEX9	6	2	43.4/5.7		5.71
		Fructose-bisphosphate aldolase class 2	P0AB71	6	2	39.1/5.9		5.86
9	$\downarrow 1.5 (0.006)$	DNA protection during starvation protein	P0ABT2	50	12	18.7/6.1	16.9/5.6	6.11
		50S ribosomal protein L13	P0AA10	10	2	16.0/9.9		9.91
10	\uparrow 1.4 (0.011)	10 kDa chaperonin	P0A6F9	65	6	10.4/5.2	13.4/4.9	5.24
		30S ribosomal protein S6	P02358	13	2	15.7/5.0		5

^{*a*} Significantly different from control, as displayed in the Delta2D quantitation table (Student's *t*-test). ^{*b*} Calculated in Delta2D, calibrated with Bio-Rad 2-D PAGE standards.

quinone oxidoreductase subunit G, D-tagatose-1,6-bisphosphate aldolase subunit, periplasmic pH-dependent serine endoprotease, dihydrofolate synthase/folylpolyglutamate synthase and bifunctional protein HldE.

For the 26 proteins with increased levels, four enzymes required for the aminoacyl-tRNA biosynthesis pathway were identified, glycine-tRNA ligase (beta subunit), lysine-tRNA ligase

(heat inducible), proline-tRNA ligase and methionine-tRNA ligase.

These results suggest that exposure of *E. coli* to microwave irradiation under the experimental conditions led to slower growth, likely due to stress imposed by microwave exposure. In response to the stimulation, metabolism of *E. coli* cells is down-regulated. The finding that four enzymes involved in tRNA

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synthesis were up-regulated also corroborates this hypothesis, as the literature points to the regulation of cellular tRNA levels as a response to stresses, such as starvation and heat.⁴⁰⁻⁴²

Taken together, the proteomic results from this study support the possibility that exposure of *E. coli* to non-lethal microwave irradiation at constant culture temperatures can be perceived by the microorganism as a stimulus. In adaptation, the bacteria modulate their metabolism and tRNA biosynthesis.

3. Materials and methods

Chemicals were purchased from Sigma-Aldrich unless otherwise stated, and were used without further purification. *E. coli* DE3 was a gift from Professor Charles Déspres' lab (Department of Biological Sciences, Brock University).

3.1. Microwave irradiation

A CEM Discovery Coolmate microwave system that operates at 2.45 GHz was used for microwave irradiation, and controlled with Synergy software from CEM (Matthews, NC). The standard Reaction Vessel (CEM part number 168302) was used as culture tubes, inserted into the standard Attenuator Assembly (CEM part number 542476), which provides cooling to the culture through circulating coolant. Solvay Solexis H Galden Coolant Zt 130 was pre-cooled in the coolant reservoir of Coolmate by liquid nitrogen. Temperature of cultures was preset as desired in Synergy and maintained through adjustment of the temperature of coolant and flow rates. The instrument and culture set-up are described in Fig. 1. Speed of the stirring bar was determined with an Omega HHT13 tachometer.

3.2. Generation of *E. coli* growth curves from cultures in a shaking incubator

Single bacterial colonies from LB agar plates were grown in LB broth in a shaking incubator at 37 $^{\circ}$ C and 260 rpm until an OD₆₀₀ in a range of 0.8–0.9 was reached. Aliquots (200 µl) were taken at appropriate intervals, and absorbance was read at 600 nm using a Biotek 405U microtitre plate reader.

3.3. Generation of *E. coli* growth curves from tube cultures in the presence of microwave and in hotplate oil-bath

100 μ l of cultures grown in the shaking incubator was added to fresh LB media (4.9 ml, to make a 1 : 50 dilution) in the reaction vessel in the CEM microwave reactor and exposed to a maximal 10 W microwave irradiation while the culture temperature was maintained at 37 °C through simultaneous cooling with constant stirring (stirring speed set as "high" at 800 \pm 20 rpm). The speed of the stirring bar was measured in rpm by an Omega HHT13-Kit tachometer. The culture tube was kept sterile by a foam plug. Aliquots (200 μ l) were taken at appropriate intervals, and absorbance was read at 600 nm using a Biotek 405U microtitre plate reader.

For growth recovery experiments, microwave irradiation was terminated after 5 h and the culture tube was transferred to an oil-bath and incubated at 37 °C. Growth was monitored for 26 h after initial inoculation by taking aliquots (200 μ l) for OD₆₀₀

measurement. A total viable cell count test was carried out at the end of each experiment.

As a control, *E. coli* cultures were incubated in an oil-bath at 37 °C and stirred. The speed of the stirring bar was measured by a tachometer to be 800 ± 20 rpm. Culture growth was monitored for 27.5 h after inoculation. A total viable cell count test as described below was carried out at the end of each experiment.

3.4. Total viable cell counts

To determine the total viable cell count (colony-forming units, CFU, per ml) at the end of each growth curve experiment, bacterial cultures were diluted up to 10^8 -fold by serial dilution. Aliquots (100 µl) of each dilution were streaked on an LB agar plate. Single colonies were counted after 18 h of incubation at 37 °C in a static incubator.

3.5. DO measurements

The apparatus used for measuring DO levels was made in-house (Fig. 6). Instead of an external cooling jacket, coolant was circulated through an internal cooling tube inserted into the culture tube that contained uninoculated LB media used in the growth experiments, isolated from the environment by a foam plug. The temperature inside the tube was monitored by a fibre optic temperature probe.

LB media (35 ml) was irradiated with microwave as described above, with stirring. At 1 and 2 h, the foam plug and cooling coil were removed, the vessel was quickly sealed, and DO was measured by inserting a Vernier Micro SD Optical Dissolved Oxygen probe into the vessel. DO levels were recorded after values stabilized (usually after two minutes). DO levels in nonirradiated controls were measured accordingly.

3.6. Bacterial culture and total protein extraction

LB media was inoculated with single colonies of *E. coli* DE3 and incubated in a shaking incubator at 37 °C and 260 rpm until an OD₆₀₀ in a range of 1.32–1.36 was reached. A portion of this preculture (4.5 ml) was then added to fresh LB media (40.5 ml, to make a 1 : 10 dilution) in the reaction vessel in CEM microwave reactor and exposed to a maximal 10 W microwave irradiation while the culture temperature was maintained at 37 °C through



Fig. 6 Setup for DO measurement. LB media isolated from the environment with the use of a foam plug were cooled through circulation in an internal cooling tube.

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simultaneous cooling and stirring was set at high. The speed of the stirring bar was measured by a tachometer (Omega HHT13-Kit) to be 800 \pm 20 rpm. The control was prepared similarly, except that the reaction vessel was placed in a water bath on a hotplate. The reaction vessels were isolated from the environment by foam plugs. After 5 h, cultures were transferred to 50 ml-Falcon tubes and centrifuged at 4255 RCF for 20 minutes at 4 °C. The supernatants were discarded and the cell pellets were washed three times by resuspension in 1× PBS (30 ml), and centrifugated as above. Supernatant-free cell pellets were flash frozen with liquid nitrogen and was stored at -80 °C. Three biological replicates per condition were prepared for analysis.

Pellets were homogenized *via* automated frozen disruption and solubilized in protein/kinase/phosphatase inhibitorsupplemented 2DE lysis buffer as described previously.⁴³ Sample aliquots were snap-frozen and stored at -80 °C. Total protein concentrations were determined using a solid-phase dot-blot assay as described previously,⁴⁴ with BSA as the calibration standard.

3.7. 2DE and in-gel protein detection

All methods for immobilized pH gradient strip (IPG) rehydration, isoelectric focusing (IEF), and subsequent sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) were carried out as described in detail previously.45 Briefly, 100 µg total protein from each sample was reduced and alkylated prior to passive rehydration into a 7 cm pH 3-10 non-linear (Ready-Strip[™] IPG Strip (Bio-Rad, Hercules, CA)) for 16 h. IEF in a Protean i12 IEF Cell (Bio-Rad, Hercules, CA) was carried out at 4000 V for 37 500 VH with several electrode wick changes during linear voltage ramping to facilitate desalting. Focused IPGs were equilibrated with reduction and alkylation prior to further protein resolution in large-format ($18 \times 17 \times 0.1$ cm) 12.5% T gels. Two IPGs were adjacently resolved per large gel, yielding 'mini-tall' gels (i.e. 7 cm first-dimension and 18 cm long seconddimension). Unstained protein marker (10-200 kDa; New England Biolabs, Ipswich, MA) were resolved in parallel to approximate resolved proteoform molecular weights. Electrophoresis was carried out in a cold room at 4 °C, for 20 min at 300 V then 120 V to completion (~18 h). Gels were fixed and stained using highly sensitive colloidal Coomassie Brilliant Blue (cCBB) as described previously.⁴⁶ Three technical replicates per biological replicate were assessed.

3.8. Gel imaging and analysis

Destained gels were imaged *via* transmissive densitometry using the GS-900TM Calibrated Densitometer (Bio-Rad, Hercules) at maximal scanning resolution. 16-Bit greyscale TIFF images were exported for analysis. Quantitative image analysis was performed using Delta2D (Decodon GmbH, Greifswald, Germany) following the manufacturerrecommended workflow (image alignment through to statistical analysis), with automated processes (*i.e.* warping, consensus spot pattern) manually refined to ensure accuracy.⁴⁷ A Welch's *t*-test was carried out for all protein spots across the control and treatment groups using the following parameters: *p*-value based on all permutations, critical *p*-value of 0.01, and false discovery rate from permutations to exceed no more than 5 proteins. From the resulting list of significantly altered spots (18), any spot with a relative standard deviation (RSD) of >40% of the normalized volume in either experimental group was excluded. The remaining spots, the majority of which exhibited \leq 30% normalised volume RSD across all replicates, were considered genuine after again assessing quality by manual comparison to ensure the highest possible reliability.

For isoelectric point (pI) and molecular weight (kDa) calibration, one mini-tall gel in which 2-D PAGE standards (Bio-Rad, Hercules, CA) were resolved, and one mini-tall gel in which 10 μ g *E. coli* sample spiked with 2-D PAGE standards was resolved (to ensure correct alignment) were prepared as described above. Calibration gel images were imported, warped, and annotated in Delta2D according to manufacturer recommendations for use of the in-built calibration tool.

3.9. Spot picking and in-gel digestion

Protein spots of interest were manually excised from several gels from both experimental groups, pooled, and transferred to microcentrifuge tubes. In-gel tryptic digestion was carried out as described previously⁴⁷ using 3 ng μ l⁻¹ trypsin with digestion performed at 4 °C for 30 min followed by 12 h at ambient temperature. Peptide solutions were recovered into microcentrifuge tubes and dried in a speed vacuum. Samples were shipped at ambient temperature for LC-MS/MS analysis.

3.10. Mass spectrometry analysis and protein identification

Mass spectrometry analysis was carried out using a Q-Exactive mass spectrometer. Data-dependant acquisition was preformed on the top 10 peptides per duty cycle with automatic switching between MS and MS/MS. Full-scan MS mode (375-1600 m/z) was operated at a resolution of 70 000 with automatic gain control (AGC) and a target of $1 imes 10^6$ ions. Ions selected for MS/MS (fixed first mass 100 m/z) were subjected to the following parameters: resolution 17 500, target of 1×10^5 ions, 1.5 m/z isolation window, normalized collision energy 27.0 V and dynamic exclusion 20.0 s. Source ionization parameters were as follows: spray voltage, 1.9 kV; capillary temperature, 280 °C; and s-lens RF level, 50.0. Chromatographic separation of peptides was accomplished using a Zorbax 300SB-C18 column (3.5 µm i.d. \times 150 mm, particle size 5 μ m, pore size 100 Å, Agilent Technologies, Wilmington, DE). Here, peptides were loaded onto a Zorbax 300SB-C18 trap cartridge at a flow rate of 2 µl per minute for 10 min. After washing with 0.1% formic acid the peptides were eluted using a 5-40% B gradient for 30 min at a flow rate of 250 nl min⁻¹ acid (mobile phase A = 0.1% formic acid; mobile phase B = 0.1% formic acid in acetonitrile) on an Agilent 1260 capillary/nano system.

LC-MS/MS results were searched using Proteome Discoverer (version 2.2, Thermo Scientific against the Uniprot *E. coli* (strain K12) (4340 entries)) and the Proteome Discoverer contaminants database in which raw files were searched using the Sequest HT

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algorithm. Peptides produced by digestion with trypsin, with a maximum of two missed cleavages, were matched using precursor and fragment mass tolerances of 10 ppm and 0.04 Da, respectively. Carbamidomethylation of cysteine residues (C) was selected as a static modification. Oxidation (M), deamidation (NQ) and acetyl (protein N-term) were chosen as dynamic modifications. Peptide spectrum matches (PSMs) were verified based on *q*-values set to 1% false discovery rate (FDR).

4. Conclusions

When E. coli cultures were exposed to up to 10 W microwave irradiation, while temperature was maintained at 37 \pm 1 $^{\circ}\mathrm{C}$ with occasional brief fluctuations of ± 2 °C, the growth slowed relative to non-irradiated cultures. When microwave irradiation was terminated, however, growth was restored. The identifications of the altered proteins suggest that the microwave exposure likely leads to the alteration of the metabolic profiles of the bacteria, however the long-term effects of such exposure on bacterial physiology remain unclear at this time. In connection with our previous findings, it appears that exposure of biological systems to non-lethal microwave irradiation can lead to alterations of enzymatic activities, membrane properties, and cellular metabolism. Future work will focus on the validation of microwave exposure related biological processes in E. coli and transcriptomic analysis of the proteins involved in these processes.

Conflicts of interest

The authors declare no conflict of interest.

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