Background: *C. glutamicum* has traditionally been grown in neutral-pH media for amino acid production, but in a previous article we reported that this microorganism is a moderate alkaliphile since it grows optimally at pH 7.0–9.0, as shown in fermentor studies under tightly controlled pH conditions. We determined the best pH values to study differential expression of several genes after acidic or basic pH conditions (pH 6.0 for acidic expression and pH 9.0 for alkaline expression). Thus, it was interesting to perform a detailed analysis of the pH-adaptation response of the proteome of *Corynebacterium glutamicum* ATCC 13032 to clarify the circuits involved in stress responses in this bacterium. In this paper we used the above indicated pH conditions, based on transcriptional studies, to confirm that pH adaptation results in significant changes in cytoplasmatic and membrane proteins.

Results: The cytoplasmatic and membrane proteome of *Corynebacterium glutamicum* ATCC 13032 at different pH conditions (6.0, 7.0 and 9.0) was analyzed by classical 2D-electrophoresis, and by anion exchange chromatography followed by SDS-PAGE (AIEC/SDS-PAGE). A few cytoplasmatic proteins showed differential expression at the three pH values with the classical 2D-technique including a hypothetical protein cg2797, L-2.3-butanediol dehydrogenase (ButA), and catalase (KatA). The AIEC/SDS-PAGE technique revealed several membrane proteins that respond to pH changes, including the succinate dehydrogenase complex (SdhABCD), F0F1-ATP synthase complex subunits β, α and δ (AtpF, AtpH and AtpA), the nitrate reductase II α subunit (NarG), and a hypothetical secreted/membrane protein cg0752. Induction of the F0F1-ATP synthase complex β subunit (AtpD) at pH 9.0 was evidenced by Western analysis. By contrast, L-2.3-butanediol dehydrogenase (ButA), an ATPase with chaperone activity, the ATP-binding subunit (ClpC) of an ATP-dependent protease complex, a 7 TMHs hypothetical protein cg0896, a conserved hypothetical protein cg1556, and the dihydrolipoamide acyltransferase SucB, were clearly up-regulated at pH 6.0.

Conclusion: The observed protein changes explain the effect of the extracellular pH on the growth and physiology of *C. glutamicum*. Some of the proteins up-regulated at alkaline pH respond also to other stress factors suggesting that they serve to integrate the cell response to different stressing conditions.
Background

Since its discovery [1], **Corynebacterium glutamicum** ATCC 13032 is one of the most important organisms used industrially to produce not only various amino acids, but also vitamins, organic acids, proteins and diverse raw chemicals [2-4]. This strain is a facultative anaerobic Gram-positive soil bacterium with a high G+C content. Recently, the entire genome sequence of **C. glutamicum** ATCC 13032 has been determined in parallel in Japan [5] and in Germany [6]. Availability of the genome sequence allowed considerable progress in gene organization studies, transcriptional analyses [7-11], proteome [12,13] and evolutionary studies of this strain [14].

Among the most influential environmental factors in the growth and physiology of bacteria are temperature and extracellular pH. Due to the fluctuations of the pH in the nature microorganisms have developed throughout the evolution diverse adaptation strategies to minimize the damage induced by the stress of an acid or basic environment [15]. Most of the studies in this field were made in *E. coli* [16] or *Bacillus* species [17-19]. For example, in enteric bacteria changes in pH contribute to disease. Low pH enhances expression of numerous virulence factors such as the *ToxR-ToxT* virulence regulon in *Vibrio cholerae* [20], the *phoP-phoQ* regulon of *Salmonella enterica* [21], and the pH 6 antigen of *Yersinia pestis* [22]. In *E. coli* low pH accelerates acid consuption and proton export, while coinducing oxidative stress, possibly through increased production of oxygen radicals. On the other hand, high pH accelerates proton import while repressing the energy-expensive systems of flagellar biosynthesis and chemotaxis [16]. In Gram-positive soil bacterium *Bacillus subtilis* the σ^W^ regulon is induced as a response towards an alkaline shock [19].

In *C. glutamicum*, proteomic methods have been successfully used in several studies to analyze physiological responses to different stimuli such as ammonium or nitrogen starvation [23,24], growth on glucose or acetate [25], heat shock [10], exposure to herbicides [26], or more recently the metabolic response in an H^+^-ATPase-defective mutant [27]. While numerous responses to different stresses are well studied, the pH-stress response in *C. glutamicum* remains poorly understood, and relatively few proteomic studies have been done in this field, most of them in enteric bacteria [28,29].

The technique most frequently used in *C. glutamicum* for proteome analysis has been the 2D-electrophoresis in combination with mass spectrometry [13,30,31]. In most cases this technique works satisfactorily for cytoplasmatic proteins but not for membrane proteins, due to extremes in pI and in hydrophobicity observed for many membrane proteins. Therefore, intense efforts have been made to develop diverse alternative methods for the analysis of integral membrane proteins [32-34]. One technology particularly useful for the separation and relative quantification of intact membrane proteins is the consecutive separation with anion-exchange chromatography and SDS-PAGE [35,36].

Using a combination of these techniques we observed significant patterns of membrane protein changes in response to pH variations and showed an overlapping of pH stress with other stress responses.

Methods

**Bacterial strains and growth conditions**

*Corynebacterium glutamicum* ATCC 13032 was grown in trypticase soy broth (TSB) at 30°C and at different pH conditions (pH 6.0 as acidic, pH 7.0 as control, and pH 9.0 as alkaline condition), respectively, in three identical BIOSTAT 5-liter fermentors equipped with automatic pH control. The pH values were maintained at ± 0.1 units of the initial pH conditions. The pH was controlled automatically by dropwise addition of 1 M HCl or 1 M KOH. The cultures were grown aerobically in stirred fermentors. The *C. glutamicum* growth kinetics at pH values were as described previously [9].

**Preparation of cytoplasmic protein extracts**

Cells of *C. glutamicum* ATCC 13032 from a 50 ml culture at three different pH conditions, were harvested in the mid-exponential growth phase (OD600, 3.5–4.0), by centrifugation for 10 min at 4500 × g. Preparation of cytoplasmic protein extracts and 2-DE analysis were conducted as described in [10]. Briefly, the cells were harvested by centrifugation, washed, and disrupted in a Fastprep machine (BIO 101). The disrupted cells were centrifuged at 16000 × g to remove cell debris and particulate matter. The cell-free supernatant was treated with Benzonase (Merck) and the proteins were concentrated and precipitated by acetone. Finally the protein pellet was dried and resuspended in rehydration buffer. Protein concentrations of the crude extracts were determined by the Bradford method [37]. Precast IPG strips with linear pH gradients of 4.5 to 5.5, and 4.0 to 7.0, were used for the isoelectric focusing (IEF) step. After the SDS-PAGE second dimension, gel spots were quantified relative to each other by densitometry using the ImageMaster™ 2D Platinum Software (GE Healthcare Life Science). Proteins were regarded as regulated if the corresponding ratios referring to the relative volume of the spots changed more than two-fold and if this regulation pattern was found in all biological replicates.
Preparation of membrane protein extracts. AIEC/SDS-PAGE (2D-IEC)

*C. glutamicum* membrane protein extracts were prepared by following the method described by Schluesener and co-workers [35] from 1 liter of cells grown in TSB at different pH conditions to the mid-exponential growth phase (OD600, 3.5–4.0). Briefly, cells were disrupted in a French Press (Thermo Spectronic) with 4 passages at 20,000 psi. Unbroken cells and cell debris were removed by centrifugation twice at 5,000 × g and 4°C. Membranes were enriched by ultracentrifugation at 100,000 × g and 4°C for 30 min. The resulting pellet was resuspended gently with ice-cold PBS buffer and ultracentrifugation was repeated, and membranes were washed with 2.5 M NaBr to remove membrane-associated proteins. The resulting membrane fraction was solubilized in buffer containing 2% (w/v) ASB-14 and applied onto an anion exchange column (Poros20 HQ material, Applied Biosystems). Fractions eluted from the anion exchange column were precipitated with trichloroacetic acid [38], and SDS-PAGE was performed according to Laemmli [39] using gels with concentrations of 12% acrylamide + bisacrylamide and 1% crosslinking. Gels had a separation length of 20 cm. Proteins were stained with colloidal Coomassie and gels were scanned on an image scanner with the LabScan software (Amersham Biosciences). The scanner was calibrated with a grayscale marker (Kodak), and the same settings were applied for all gels. Scanning was carried out at 300 dpi and 8-bit grayscale. Gel bands were quantified relative to each other by densitometry using the software Scion Image (version 4.0.2; Scion Corporation, http://www.scioncorp.com). Proteins were regarded as regulated if they passed at least one of these two criteria: (i) the corresponding ratios referring to the relative volumes of the spots changed more than two-fold and if this regulation pattern was found in all biological replicates; (ii) p-value < 0.1, from Student’s t-test (paired, two tailed). In many cases, all or most members of an operon showed similar regulation factors; the succinate dehydrogenase complex and the F0F1-ATPase operon was particularly consistent.

In-gel tryptic digestion and MALDI-TOF PMF protein identification

Cytoplasmatic protein spots were excised from gels and digested with modified trypsin (Promega) as described in [11]. Membrane protein bands were excised from the Maxi-size gels, destained following the protocol of Hellman *et al* [40], and were also digested with modified trypsin (Promega). The peptide mass fingerprints were determined with Voyager DE-Pro Instrument in reflector mode (Applied Biosystems). Identification was performed using the MASCOT software [41]. Parameters of the search were set as in [28].

Western-Blotting of AtpD

Immunological detection of AtpD was performed using 25 μg of cytoplasmic or membrane protein extracts. Proteins were diluted in loading buffer (62 mM Tris-HCl, pH 6.8, 3% SDS, 5% β-mercaptoethanol, 10% glycerol and 0.01% bromophenol blue), and incubated for 5 min at 95°C. Membrane protein samples were not boiled but incubated for 30 min at 60°C to avoid aggregation. After brief cooling, the mixture was centrifuged (5 min, 12,000 × g) and the supernatant was loaded onto the gel. Proteins were analyzed by SDS-PAGE (1 hour, at 30 mA per gel) and electrotransferred onto PVDF membranes applying a current of 1 mA/cm² of gel at maximum for 1 hour. "Prestained Protein Molecular Weight Marker" (MBI Ferments), ranging in size between 120 kDa and 20 kDa were used as molecular weight markers. Membranes were blocked by incubating in PBST [137 mM NaCl, 1.5 mM KH2PO4, 7.9 mM Na2HPO4, 2.7 mM KCl, 0.5% Tween 20 (v/v)] with 3% nonfat milk powder for 1 h at room temperature. Primary antibody was added to this buffer (1:10,000), and the blot was incubated for 2 h or overnight. Membranes were washed 4 times thoroughly in PBST, and then incubated with the anti-rabbit secondary antibody conjugated to horseradish peroxidase (Sigma-Aldrich). Proteins were detected using p-coumaric acid, 3-aminoptalhydrazide (Luminol, Sigma-Aldrich), and 30% H2O2 as substrates.

Results

Cytoplasmatic proteome changes due to pH adaptation

Using three different pH conditions (pH 6.0, pH 9.0 and control pH 7.0) defined in a previous study [9], protein changes in 2D-gels were analyzed from four independent experiments (biological replicates). Initially, precast IPG strips with linear pH gradients of 4.0 to 7.0 [10] were used for the isoelectric focusing step, in which most of the *C. glutamicum* proteins were found within the pl range 4.0–7.0. With this first approach three proteins were detected that showed different abundance in the three pH conditions (Fig. 1). Another protein spot seemed to have different molecular mass at the three pH conditions (indicated as spots 4, 5 and 6 in Fig. 1). To achieve a more detailed resolution from all of these proteins, an ampholite pH range of 4.5 to 5.5 was used (Fig. 2). This 'enlarged 2D-gel' conditions confirmed the same protein spots observed initially. All of these proteins were unequivocally identified by MALDI-TOF PMF analysis (Table 1), and in all cases no more than one protein was found in each gel spot. The two proteins that showed clearly a large increase in response to acidic pH (spots 1 and 2) were identified as: i) a hypothetical protein cg2797 and, ii) ButA which corresponds to L-2,3-butanediol dehydrogenase/acetoine reductase. The only protein (spot 3) that increased clearly in response to alkaline pH was identified as a catalase (KatA). Moreover, modifications by proteolytic process-
Table 1: Cytoplasmic proteins whose levels increase or decrease significantly in response to pH shock*a

| Spot | Name           | Accession no. | Function                                                                 | Cal. MW kDa | Calc. pl | Factor Mowseb | pH 6.0/7.0 ratio | S.D.c | pH 9.0/7.0 ratio | S.D.c | pH 9.0/6.0 ratio | S.D.c |
|------|----------------|---------------|--------------------------------------------------------------------------|-------------|----------|---------------|-----------------|-------|-----------------|-------|-----------------|-------|
| 1    | Hypot. protein | cg2797        | (2R)-phospho-3-sulfolactate synthase (comA)                              | 30.3        | 4.77     | 96            | ↑3.21           | 0.06  | 1.32            | 0.22  | ↓0.42           | 0.25  |
| 2    | ButA           | cg2958        | L-2,3-butanediol dehydrogenase/acetoin reductase                         | 27.0        | 4.51     | 89            | ↑2.18           | 0.14  | 0.85            | 0.17  | ↓0.43           | 0.10  |
| 3    | KatA           | cg0310        | Catalase                                                                | 58.6        | 5.13     | 255           | 0.73            | 0.03  | 1.45            | 0.12  | ↑2.03           | 0.08  |
| 4    | Sod            | cg3237        | Iron/manganese superoxide dismutase                                      | 22.0        | 5.15     | 83            |                 |       |                 |       |                 |       |
| 5    | Sod            | cg3237        | Iron/manganese superoxide dismutase                                      | < 22.0      | 5.15     | 92            |                 |       |                 |       |                 |       |
| 6    | Sod            | cg3237        | Iron/manganese superoxide dismutase                                      | < 22.0      | 5.15     | 97            |                 |       |                 |       |                 |       |

*aOnly one protein was proposed by MASCOT software for each spot. Arrows indicate up- or down-regulation. In boldface are shown the proteins whose ratios change more than two fold.
bMowse factor, significance level provided by MASCOT software.
cStandard deviation; n = 4
Comparison of 2D-gels of cytoplasmic proteins of *Corynebacterium glutamicum* ATCC 13032 grown at pH 7.0, control condition (upper panel), versus pH 6.0 and pH 9.0, acid and basic conditions (left and right lower panels, respectively). The linear pH range used in first dimension was 4.0 to 7.0. In the space between upper and lower panels the spots 4, 5 and 6 identified by MALDI-TOF PMF are shown in more detail. Molecular masses, in kDa, are indicated on the right side of the upper panel.

**Figure 1**

Comparison of 2D-gels of cytoplasmic proteins of *Corynebacterium glutamicum* ATCC 13032 grown at pH 7.0, control condition (upper panel), versus pH 6.0 and pH 9.0, acid and basic conditions (left and right lower panels, respectively). The linear pH range used in first dimension was 4.0 to 7.0. In the space between upper and lower panels the spots 4, 5 and 6 identified by MALDI-TOF PMF are shown in more detail. Molecular masses, in kDa, are indicated on the right side of the upper panel.
Comparative expanded (ampholite range 4.5 to 5.5) 2D-gel analysis of *Corynebacterium glutamicum* ATCC 13032 cytoplasmic proteins at pH 7.0, control condition (upper panel), versus pH 6.0 and pH 9.0, acid and basic conditions (left and right lower panels, respectively). To increase the resolution of the spots, a linear pH range of 4.5 to 5.5 was applied in the first dimension. Protein spots 4, 5 and 6 identified by MALDI-TOF PMF are shown in more detail in the space between upper and lower panels. Molecular masses in kDa are indicated on the right side of the upper panel.
ing due to basic pH could be responsible for the occurrence of three protein spots (4, 5 and 6) with identical pI but apparently different molecular mass. These three spots correspond to superoxide dismutase protein (Sod).

**Membrane proteome changes in response to pH stress**

A comparative analysis of the membrane proteome was performed with three independent experiments (biological replicates), and using the same acidic and basic pH culture conditions as for cytoplasmatic proteins. Membrane proteins were resolved using a combination of anion exchange chromatography (AIEC) as first dimension, and SDS-PAGE as second one [35]. Fig. 3 shows an AIEC separation of the *C. glutamicum* membrane fractions at different pH values after the washing and solubilization steps. Fig. 4 shows the SDS-PAGE separation in Maxi-size gels of *C. glutamicum* membrane fractions obtained from cultures grown at different pH values. For most of the proteins no differences in expression were visible. At the present time there is no satisfactory software for the comparison of bands in this type of gels, and therefore, the first screening of the gels to choose the protein spots that seemed to increase or decrease their intensity, was performed visually [36]. Several differences were detected by comparing the protein patterns obtained at the different pH values. Seventeen proteins were observed that showed significant changes in response to different pH values (Fig. 4). Six of them (spots 2, 10, 11, 12, 16 and 17) were drastically up-regulated at pH 6.0, while eight proteins (spots 1, 3, 4, 5, 6, 7, 13 and 15) were up-regulated at pH 9.0. Additionally, we observed other three proteins (spots 8, 9 and 14) that change in response to pH variations although the data were not significant enough to define an up-regulation at pH 9.0 (spots 8 and 9), or a regulation at pH 6.0 or
Separation of the solubilized membrane fractions from *C. glutamicum* ATCC 13032 in control condition (upper panel), versus acidic and basic conditions (left and right lower panels, respectively). Before separation in the first dimension by AIEC, the membranes were washed twice with 2.5 M NaBr and solubilized in buffer containing 2% (w/v) ASB-14. After the TCA precipitation of the AIEC fractions, SDS-PAGE was used as second dimension in Maxi size gels. Fraction numbers of the AIEC are given at the top of the gels. Numbers indicate the protein spots with differences among the 3 pH values chosen for MALDI-TOF analysis and are listed in Table 1. Molecular masses (in kDa) are shown on the left side of the panels.
| Spot | Name          | Accesion no. | Function                          | L<sup>a</sup> | Calculated MW (kDa) | Calculated pI | pH induction<sup>b</sup> | Regulation factors<sup>c</sup> |
|------|---------------|--------------|-----------------------------------|---------------|---------------------|---------------|---------------------|--------------------------------|
|      |               |              |                                   |               |                     |               |                     | 6.0/7.0          | 9.0/7.0          | 9.0/6.0          | 9.0/6.0          | S.D.          | S.D.          | S.D.          | S.D.          |
| 1    | Sdh CD        | cg0445       | Succinate dehydrogenase CD        | 5             | 28.2                | 9.82          | pH 9.0             | 0.68±           | 0.15             | 1.30±           | 0.06            | 2.04±         | 0.52          |
| 2    | Hypothetical  | cg0896       | protein (7 TMHs)                  | 7             | 10.9                | 4.35          | pH 6.0             | 0.15            | 0.76             | 1.18            | 0.16            | 0.6±          | 0.14          |
|      | Sdh A         | cg0446       | Succinate dehydrogenase A         | a             | 7.40                | 5.40          | pH 9.0             | 0.61            | 0.18             | 1.26            | 0.20            | 2.85±         | 1.07          |
| 4    | Sdh B         | cg0447       | Succinate dehydrogenase B         | a             | 2.65                | 5.43          | pH 9.0             | 0.61            | 0.18             | 1.41±           | 0.22            | 2.52±         | 0.77          |
| 5    | AtpF          | cg1364       | F0F1-type ATP synthase b subunit  | a             | 20.9                | 4.97          | pH 9.0             | 0.48±           | 0.12             | 1.26            | 0.20            | 2.24±         | 0.24          |
| 6    | AtpH          | cg1365       | F0F1-type ATP synthase delta subunit | a            | 28.7                | 5.36          | pH 9.0             | 0.55            | 0.25             | 0.85            | 0.21            | 1.75±         | 0.46          |
| 7    | AtpA          | cg1366       | F0F1-type ATP synthase alpha chain | a             | 58.6                | 4.77          | pH 9.0             | 0.60±           | 0.08             | 0.88            | 0.11            | 1.47±         | 0.16          |
| 8    | AtpD          | cg1368       | F0F1-type ATP synthase beta chain<sup>h</sup> | a            | 52.4                | 4.63          | ---<sup>c</sup>    | 0.61±           | 0.10             | 0.68±           | 0.06            | 1.12±         | 0.08          |
| 9    | GluB          | cg2137       | Glutamate secreted binding protein | a             | 31.5                | 3.89          | ---<sup>c</sup>    | 0.84            | 0.36             | 0.98            | 0.22            | 1.30          | 0.38          |
| 10   | ClpC          | cg2963       | ATPase with chaperone activity (heat shock protein) | a             | 101.5               | 4.93          | pH 6.0             | 2.91            | 2.38             | 1.00            | 0.20            | 0.54          | 0.24          |
|   | Protein Name                  | Gene ID | Localization      | pH 6.0 | pH 9.0 | Abbreviation | Regulation Factor | p-value | Ratio 2-fold | p-value |
|---|--------------------------------|---------|--------------------|--------|--------|--------------|-------------------|---------|--------------|---------|
|11 | CysK                           | cg2833  | c                  | 32.6   | 4.68   | O-Acetylserine (Thiol)-Lyase | ↑1.81*  | 0.88         | ↑1.56*  | 0.22       | 0.91    | 0.42    |
|12 | ButA                           | cg2958  | c                  | 27.0   | 4.51   | L-2.3-butanediol dehydrogenase/acetoin reductase | ↑1.58*  | 0.37         | 0.78    | 0.16       | 0.59    | 0.25    |
|13 | Putative secreted/membrane protein | cg0752  | s                  | 49.6   | 5.06   | Uncharacterized BCR | ↓0.30*  | 0.03         | 0.83    | 0.09       | ↑2.79*  | 0.59    |
|14 | Pks                            | cg3178  | a                  | 172.2  | 4.40   | Polyketide synthase | ---*   | 0.90         | 0.29    | 0.63*      | 0.11    | 0.77    | 0.25    |
|15 | NarG                           | cg1344  | a                  | 139.2  | 5.74   | Nitrate reductase 2 alpha subunit | ↑1.62*  | 0.85         | 1.05    | 0.14       | 0.95    | 0.48    |
|16 | Conserved hypothetical protein | cg1556  | s                  | 33.2   | 9.70   | ---                     | 3.21*  | 1.75         | 1.38*  | 0.09       | 0.58    | 0.31    |

|   | SucB                            | cg2421  | c                  | 70.7   | 4.14   | Dihydrolipoamide acyltransferase | ---*   | 1.75         | 1.38*  | 0.09       | 0.58    | 0.31    |

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a) Localization: c: cytoplasmic, a: membrane-associated, s: secreted, digits indicate numbers of transmembrane helices.
b) See discussion.
c) The statistical results of three biological culture replicates were not significant enough to define an up-regulation at pH 9.0.
d) The statistical results of three biological culture replicates were not significant enough to define a regulation at pH 6.0 or 9.0.
e) Regulation factors were obtained from three biological culture replicates for each pH condition. Arrows indicate up- or down-regulation. In boldface are shown the proteins whose ratios change more than two-fold.
f) Standard deviation; n = 3.
g) Significant difference (t-test, p < 0.1).
h) Despite the p-value > 0.1 (0.17), and the 9.0/6.0 ratio is below 2, this protein was regarded as pH regulated by Western-blotting (see Results).
9.0 (spot 14). All of these proteins were unequivocally identified by MALDI-TOF PMF analysis, and in all cases only one protein was found in each spot.

Table 2 summarizes the pH-regulated proteins of the C. glutamicum membrane fractions. The six proteins that showed clearly a drastic increase in response to acidic pH (spots 2, 10, 11, 12, 16 and 17) were respectively identified as: i) a hypothetical protein (cg0896) with 7 transmembrane helices (TMHs), ii) the ATP-binding subunit of an ATPase with chaperone activity (ClpC), iii) CysK which corresponds with O-acetylserine thiol-lyase, iv) L-2,3-butanediol dehydrogenase/acetoine reductase (ButA), v) a conserved hypothetical protein cg1556, and vi) a dihydrolipoamide acyltransferase (SucB).

The proteins clearly induced in response to alkaline pH (spots 1, 3, 4, 5, 6, 7, 13 and 15) were respectively identified as: i) the three subunits of the succinate dehydrogenase complex (SdhCD, SdhA and SdhB), ii) subunits b, δ and α of the F_{0}F_{1}-ATP synthase complex (AtpF, AtpH and AtpA), iii) a hypothetical secreted/membrane protein cg0752, and iv) the α subunit of the nitrate reductase II (NarG). In this group we have included again CysK, since it is also induced at pH 9.0 but less than under acidic stress (pH 6.0) (see Table 2 and Discussion). The three proteins with poorly significant abundance changes (spots 8, 9 and 14) were identified as: i) β subunit of the F_{0}F_{1}-ATP synthase complex (AtpD), ii) a secreted glutamate binding protein (GluB), and iii) Pks, which corresponds with a polyketide synthase, a type of enzymes that are frequently involved in secondary metabolism. All of these protein bands were quantified relative to each other using densitometry and the expression factor (up- or down-regulation) found in three independent separation experiments is shown in Table 2.

**Immunological detection of AtpD protein**

The data on the AtpD protein in the membrane fractions were not significant, unlike the rest of the subunits of the F_{0}F_{1}-ATP synthase complex identified in this study. To analyze the expression of this protein in more detail western-blot analyses were performed with an antibody against a peptide fragment of Spinach AtpD, using the same samples from the three pH conditions (three biological replicates) described above. The immunological detection of AtpD protein is shown in Fig. 5; a hybridization band of about 53 kDa was found in the membrane fractions of the three cultures, but not in the cytoplasmic proteins. These bands correspond perfectly with the expected size of the AtpD protein. The intensity of the AtpD band in cells grown at pH 9.0 was 63.2% higher than at pH 6.0 (Fig. 5), and 26.7% higher at pH 9.0 than at pH 7.0. Therefore, we concluded that AtpD is induced at alkaline pH (9.0). These results at the translational level for the F_{0}F_{1}-ATP synthase beta chain correlate well with those reported at the transcriptional level [9], and with those observed for the other proteins of F_{0}F_{1}-ATP synthase complex in this work. In summary, all the studied components of the F_{0}F_{1}-ATP synthase complex are up-regulated at alkaline pH.

**Discussion**

Up to date, little information is available concerning the physiology of C. glutamicum when grown at different extracellular pH conditions (alkaline or acid), although respiration and membrane transport processes may be affected by these changes in external pH. We used two different strategies to study the analysis of the C. glutamicum cytoplasmatic and membrane proteome at different pH conditions.

The first approach was to analyze the proteome by the classical two-dimensional polyacrylamide gel electrophoresis (2-D PAGE). Two proteins were identified as acidic pH induced (Spots 1 and 2, Table 1). A hypothetical protein cg2797 was found to be up-regulated 3.2-fold and the L-2,3-butanediol dehydrogenase/acetoine reductase showed a minor up-regulation factor of 2.18 (Table 1). No functional domains were detected in the cg2797 protein and homology searches revealed only one protein, ComA [a (2R)-phospho-3-sulfolactate synthase] in the actinobacteria *Rubrobacter xylanophilus* (Rxyl_2563) that show 57% identity. This protein is involved in the Coenzyme M (CoM; 2-mercaptoethanesulfonic acid) biosynthesis, which was thought to be exclusive of methanogenic bacteria until Krum and Ensign [42] also found it in *Xantobacter Py2* and in *Rhodococcus rhodochrous* B276. In the hyperthermophile marine methanogen *Methanococcus jannaschii*, ComA has been described as the first enzyme required for CoM biosynthesis [43]. This enzyme catalyzes the addition of sulfite to phosphoenolpyruvate forming L-2-phospho-3-sulfolactate, which through several reactions (conducted by the other three enzymes of the pathway, ComB, ComC and ComDE) yields coenzyme M. Since *C. glutamicum* is a Gram-positive bacteria (actinobacteria) related to *Rhodococcus*, it seems to be possible that ComA of *C. glutamicum* is...
involved in the synthesis of coenzyme M. Moreover, this gene seems to form an operon with another hypothetical protein cg2796. There is experimental evidence [44] that this operon is repressed by a transcriptional factor DtxR [45], homologous to the diphtheria toxin repressor DtxR of C. diphtheriae, which is involved in regulation of iron metabolism in C. glutamicum.

The second pH-regulated protein is the L-2,3-butanediol dehydrogenase/acetoin reductase (ButA). This protein catalyzes the formation of 2,3-butanediol from pyruvate with acetalactate and acetoin as intermediates. Proteomic studies in C. glutamicum showed a sharp increase in expression of ButA after heat shock [46], and the same was observed using microarrays (C. Barreiro and J. F. Martín, unpublished results). Interestingly, after heat-shock a significant drop in intracellular pH and an increase of the acetate level was observed in E. coli [47]; all three enzymes involved in the 2,3-butanediol synthesis were found to be strongly induced by acetate in several bacteria [48,49]. Besides, in our laboratory another C. glutamicum protein identified as Pqo (formerly PoxB) was found moderately up-regulated in response to heat shock [10]. This protein is annotated as pyruvate:quinone oxidoreductase (pyruvate oxidase) that catalyzes the reaction from pyruvate to acetate [50]. These data show that both stressful situations (pH and temperature) up-regulate this route, and the ButA response may serve to integrate different environmental stresses. This is not surprising, since in several bacteria it has been observed that the pH interacts with other environmental factors such as oxygen, temperature or salt concentration.

The catalase (KatA), was found to be up-regulated 2.0-fold at basic pH (pH 9.0/pH 6.0 ratio) (Table 1). This protein has also been observed to be up-regulated in transcriptomic studies after heat shock (C. Barreiro and J. F. Martín, unpublished results). In addition, KatA is repressed by the AraC-type regulator RipA under iron limitation conditions [51].

In all these cases, the observed modification of the proteins is the result of an adaptation phenomenon to prolonged incubations at acidic or basic pH. This includes transcriptional up- or down-regulation and post-transcriptional regulation. The latter may include post-translational protein modifications. The superoxide dismutase (Sod) seems to have modifications by proteolytic processing due to basic pH. This situation has also been observed in S. coelicolor proteomic studies by Hesketh and co-workers [52], where they reported several other protein spots with lower than expected molecular weights and often different pl values, suggesting modifications by proteolytic processing. Both proteins (KatA and Sod) have been found in most organisms, aerobic and anaerobic, and play a key role in cellular protection against the highly toxic reactive oxygen species (ROS) such as superoxide radical anions (O$_2^-$), hydrogen peroxide (H$_2$O$_2$) and hydroxyl radicals (OH$^-$). In C. glutamicum Sod belongs to the Mn-Sod type, which uses manganese as cofactor. This protein catalyzes the conversion of superoxide radicals, generated indirectly in aerobic respiration, to molecular oxygen and hydrogen peroxide; the latter is broken to water by catalase or peroxidase. In most microorganisms, the Sod protein has a constitutive expression, although in E. coli it was observed that the Mn-Sod is an inducible enzyme after exposure to oxygen or to superoxide radicals [53]. Recently, both catalase and superoxide dismutase genes of Shewanella oneidensis have been shown to be induced after oxidative stress generated by the extracellular alkaline pH [54]. The data obtained in this study suggest that C. glutamicum at an alkaline extracellular pH also develops this protective response against oxidative stress.

The number of proteins identified by the 2D procedure, that show significant changes in expression level as a function of pH, was very low. This observation is not surprising since work in Bacillus indicates that only a small number of cytoplasmic proteins change in response to pH stress [55]. This phenomenon reflects the fact that the intracellular pH (cytoplasmatic) does not show drastic changes in response to variations in the extracellular pH [56].

On the other hand, the proteins in the cell surface or those that are partially or entirely exposed to the external environment are likely to be more susceptible to changes in the extracellular pH than the cytoplasmic ones. For this reason the C. glutamicum proteome was analyzed by a second approach, the newly reported two-dimensional separation AIEC/SDS-PAGE [35,36,57]. This technique is much more effective than the previous one for resolution of integral membrane proteins (IMPs) and membrane-associated proteins (MAPs). Our results indicate that the proteome obtained by this method at pH 6.0 is very different from that observed at pH 7.0 and 9.0. This difference can be even observed in the elution profiles after the ion exchange chromatography (first step of the AIEC/SDS-PAGE) (Fig. 3). Using this procedure seventeen proteins could be identified, of which two are integral transmembrane proteins (5–7 TMHs), while ten are membrane-associated, two are secreted and three are cytoplasmic (Fig. 4, Table 2).

Two hypothetical proteins (cg0896, cg1556) were up-regulated 2.0 and 1.6-fold respectively at acidic pH, and other hypothetical secreted/membrane protein cg0752 was found up-regulated at alkaline pH about 2.8-fold (Table 2). The first of these three proteins (cg0896) is a membrane integral protein, which contains 7 transmem-
brane spanning domains according to the TMpred program [58]. Very recently, this protein was also found to be induced in C. glutamicum during growth on citrate [59], an anion that acidifies the culture medium. This protein may play an important role in acid stress responses.

Protein cg1556 that shows homology to members of the DoxD-family was found to increase 1.6-fold at acidic pH and a similar regulation factor (1.7) has been obtained for this protein after heat shock in C. glutamicum [46]. DoxD is a subunit of the terminal quinol oxidase present in the plasma membrane of Acidianus ambivalens [60]. Under normal growth conditions, the cytochrome bc1-a3 supercomplex is used as terminal electron acceptor (terminal oxidase) and cannot be easily substituted by alternative menaquinol oxidation pathways [61]. Interestingly, Kusumoto et al. [62] suggested that a third terminal oxidase exists, besides cytochrome a3 and cytochrome bd [see also [63]]. This DoxD-like protein may be an alternative oxidase.

The protein spot 10 was found to be induced 2.9-fold at pH 6.0 and was identified as a ATP-dependent protease (ClpC). This heat shock protein is one of the components of the Clp holoenzyme (ClpC, ClpP1 and ClpP2). The activation of ClpC might play a role in the degradation of misfolded proteins due to acidification; it may work also as a chaperone to assist the refolding of proteins. The clpC and clpP1P2 genes of C. glutamicum respond also to heat stress, they are positively regulated by the transcriptional factor σH (σHI) under a severe heat shock [64]. Moreover, in C. glutamicum subsp. flavum the transcriptional factor σHI controls the expression of σB under acidic stress, cold and heat shock [65]. These results suggest a double positive regulation of the expression of clpC mediated by the same sigma factor under two stress situations (pH and temperature) that appear intertwined in different microorganisms. The Bacillus subtilis heat-shock proteases are induced by heat stress but also by general stress conditions such as exposure to salt or ethanol, or starvation for glucose, phosphate or oxygen [66].

CysK, an O-acetylserylserine (thiol)-lyase was up-regulated 1.8-fold at basic pH but also showed a modest increase at pH 6.0 with a regulation factor of 1.56. In C. glutamicum proteome studies this protein has been identified as a phosphoprotein involved in the biosynthetic pathway of cysteine; it catalyzes the step from O-acetylserylserine to cysteine. Rey et al. [67] identified a repressor protein named McbR which is found to regulate several genes involved in the biosynthesis of methionine and cysteine, such as cysK. E. coli proteome studies [68] have shown that basic pH induces several amino acid metabolic enzymes, particularly CysK. The up-regulation of the cysK gene after an oxidative stress generated by alkaline extracellular pH was also observed in Shewanella oneidensis transcriptomic studies [54]. These authors suggest that high pH appears to enhance the cellular demand for cysteine, which may be needed to repair oxidative damages. This role of CysK agrees with the results obtained in this work with KatA and Sod that are up-regulated for protection against oxidative stress.

The L-2,3-butanediol dehydrogenase/acetoain reductase protein (ButA) was also found 1.6-fold induced at acidic pH in the membrane protein gels, supporting the results previously obtained in the cytoplasmic gels.

A dihydrolipoamide acyltransferase protein (SucB) was found 3.2-fold up-regulated at pH 6.0. Little is known about the encoding gene sucB; Kalinowski et al. [6] annotated it by similarity to the 2-oxoglutarate dehydrogenase complex subunit E2o of the citric acid cycle from other organisms, including odhB in Gram-positives or sucB in E. coli.

All three subunits of the succinate-DH complex were up-regulated at alkaline pH with a regulation factor ranging from 2.0 to 2.8 (Table 2). Succinate dehydrogenase is a tightly membrane-bound enzyme catalyzing the oxidation of succinate to fumarate. This enzyme is part both of the TCA cycle and of the aerobic respiratory chain. This enzyme complex consists of three subunits SdhA, SdhB and SdhC; the first two form the membrane-associated cytoplasmatic domains and the third constitutes the integral membrane domain. These three genes are likely to form an operon with two ORF encoding hypothetical proteins (cg0448 and cg4001) since they are oriented in the same direction and are coordinately up-regulated when C. glutamicum is grown on acetate [25] or in citrate [59]. Furthermore, this operon is regulated negatively by the RipA regulator [50] and positively by DtxR transcriptional regulator [44].

Interestingly, the subunits b, δ, α and β of the F0F1-ATP synthase complex (AtpF, AtpH, AtpA and AtpD) were also found up-regulated at alkaline pH. The induction of AtpD was confirmed by western blot studies. This is in accordance with a previous work in which we established that the abundance of the mRNA of the F0F1-ATP synthase operon is increased at alkaline extracellular pH [9]. The F0F1-ATP synthase operon of E. coli has been shown also to be induced by alkaline extracellular pH [16]. Furthermore, another recent study showed that this complex is also highly induced (at the mRNA and protein levels) when C. glutamicum is grown on citrate [59]. Our hypothesis is that the cells increased the expression of the ATP-ase (a reversible enzyme system) to compensate the alteration of the proton gradient produced by the alkaline pH.
Finally, the alpha subunit of nitrate reductase II named NarG was up-regulated 2.6-fold at alkaline pH. This enzyme is responsible for the nitrate reduction to nitrite and consists of two membrane-associated cytoplasmic domains NarG and NarH bound to a heme-containing membrane anchor (NarI). The nitrate reductase responsible for this reduction is encoded by the narKGHII gene cluster. The reduction of nitrate in C. glutamicum by the NarGHI complex under anaerobic conditions generates a electrochemical proton gradient, as happens in E. coli, because the quinol oxidation occurs at the outside of the cytoplasmatic membrane and the reduction of nitrate takes place in the cytoplasm [63]. Although C. glutamicum has been always regarded as an aerobic microorganism, a work has recently published showing that C. glutamicum grows in the presence of nitrate as a terminal electron acceptor [69].

Interestingly, we observed in our data that succinate dehydrogenase and proteins of the respiratory chain, i.e. the alpha subunit of nitrate reductase II and subunits of the F$_{1}$F$_{0}$-ATP synthase complex, showed between 1.4 and 2.8-fold higher abundance at alkaline pH than at acidic pH. The transcriptional regulation of the corresponding ATP-synthase genes [9] correlates well with this increase. In preliminary experiments, we found a membrane-bound NADH dehydrogenase also induced at alkaline pH (data not shown). Some of the observed changes may be indirectly caused by the change in pH and are not directly involved in the maintenance of pH homeostasis as occurs in other bacteria, e.g. it has been reported that in B. subtilis alkaline stress results in phosphate starvation [70,71].

Conclusion

Taking all these results together and the fact that the succinate dehydrogenase is the only enzyme that is part of both the citric acid cycle and the respiratory chain, a possible mechanism of coordinate regulation by alkaline pH of the respiratory chain and the tricarboxylic acid cycle is likely to exist.

In summary, the findings reported in this article suggest a complex interaction between pH, temperature and oxidative stresses. All these stress factors trigger expression of sets of identical or similar genes that appear to serve to integrate diverse stress signals that in turn activate the cell defences against the stressing conditions.

Authors’ contributions

MBI analyzed the cytoplasmic proteome, performed the western-Blotting of AtpD and wrote the manuscript draft. DS analyzed the membrane proteome and was supervised by AP. CB participated in the design of the 2-D PAGE experiments and helped to draft the manuscript. JFM prepared the research project, supervised the experiment work and was responsible for the preparation of the final version of the manuscript.

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