Acid Tolerance of *Streptococcus macedonicus* as Assessed by Flow Cytometry and Single-Cell Sorting

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An in situ flow cytometric viability assay employing carboxyfluorescein diacetate and propidium iodide was used to identify *Streptococcus macedonicus* acid tolerance phenotypes. The logarithmic-phase acid tolerance response (L-ATR) was evident when cells were (i) left to autoacidify unbuffered medium, (ii) transiently exposed to nonlethal acidic pH, or (iii) systematically grown under suboptimal acidic conditions (acid habituation). Stationary-phase ATR was also detected; this phenotype was gradually degenerated while cells resided at this phase. Single-cell analysis of *S. macedonicus* during induction of L-ATR revealed heterogeneity in both the ability and the rate of tolerance acquisition within clonal populations. L-ATR was found to be partially dependent on de novo protein synthesis and compositional changes of the cell envelope. Interestingly, acid-habituated cells were interlaced in longer chains and exhibited an irregular pattern of active peptidoglycan biosynthesis sites when probed with BODIPY FL vancomycin. L-ATR caused cells to retain their membrane potential after lethal challenge, as judged by ratiometric analysis with oxonol [DiBAC4(3)]. Furthermore, F-ATPase was important during the induction of L-ATR, but in the case of a fully launched response, inhibition of F-ATPase affected acid resistance only partially. Activities of both F-ATPase and the glucose-specific phosphoenolpyruvate-dependent phosphotransferase system were increased after L-ATR induction, distinguishing *S. macedonicus* from oral streptococci. Finally, the in situ viability assessment was compared to medium-based recovery after single-cell sorting, revealing that the culturability of subpopulations with identical fluorescence characteristics is dependent on the treatments imposed to the cells prior to acid challenge.

The existence of common strategies employed by gram-positive bacteria, particularly lactic acid bacteria, in response to acid challenge has emerged only recently (14, 53). However, generalizations about specific mechanisms involved in such responses could prove premature, since it has been shown that these mechanisms can be species or even subspecies specific (14). A typical example relevant to this report is the current knowledge about streptococcal responses to acid stress. Linkages of acid production (acidogenicity) or tolerance (acid-uricity) with pathogenicity have fueled recent interest in the basic physiology of oral streptococci (44). Today, *Streptococcus mutans* is probably the best-studied *Streptococcus* species in terms of acid stress physiology, but it is uncertain whether the behavior of this species is truly representative of the genus. For instance, a recent report on *S. sobrinus* revealed key differences from *S. mutans* concerning the involvement of F-ATPase and glucose-specific phosphoenolpyruvate-phosphotransferase system (PEP-PTS) activities in the manifestation of acid adaptation (35). Additionally, comparison of the recently completed genome sequence of *S. thermophilus* with those of other streptococci showed the inactivation or absence of genes involved in virulence, both of which were attributed to its adaptation to the milk environment (4). Therefore, more detailed studies of the acid stress physiology of food-related lactic acid streptococci are needed.

*S. macedonicus* was first isolated from traditional Greek Kasseri cheese (51), and since then it has been shown to participate in the fermenting floras of different traditional dairy products (10, 18, 43). *S. macedonicus* ACA-DC 198 exhibits antimicrobial activity against a number of important food spoilage and pathogenic bacteria, due to the production of a bacteriocin peptide named macedocin (23). In the quest for new biopreservatives, *S. macedonicus* is a promising candidate to be used as a protective culture (5). In parallel to practical application research, we wish to unravel the basic physiology of the bacterium in order to rationally incorporate it into processes.

We previously adapted an in situ viability assay for *S. macedonicus*, combining carboxyfluorescein diacetate (cFDA) and propidium iodide (PI) as viability markers (41). The cFDA-PI two-color flow cytometric assay has been applied for the study of a number of lactic acid bacterial species under thermal (52), bile salt (2), and ethanol (25) stress. The advantages of technologies that assess populations at the single-cell level over traditional culture-based techniques have been described clearly (9, 16, 48). Even though such approaches can reveal the heterogenic behavior of clonal populations—an important piece of information, especially for stress physiology (6)—their application is still far less than common. One of the major drawbacks in applying flow cytometry to bacterial populations is the natural clumping of cells (36). To overcome this problem, *S. macedonicus* cell chains were disaggregated by a mild...
sonication procedure prior to flow cytometric analysis, ensuring the assessment of the physiological status on a cell-by-cell basis, as we have described before (41).

The aim of this study was to assess the acid stress physiology of *S. macedonicus* by using dynamic cell staining techniques with different fluorochromes and flow cytometry. The main objective for implementing this approach was to determine the contribution of *S. macedonicus* intrapopulation heterogeneity to the overall behavior of the bacterial population. In a first step, we evaluated the ability of *S. macedonicus* to acquire acid tolerance (AT) after various treatments. In a second step, we attempted to reveal central cellular processes involved in the mechanism of acid tolerance. Finally, we wanted to correlate cells’ fluorescence labeling status with culturability by examining medium-based recovery after single-cell sorting. Notably, this is one of very few studies concerning a food-related lactic *Streptococcus* species under acid stress conditions.

**MATERIALS AND METHODS**

**Microorganism and growth conditions.** The gram-positive bacterium *S. macedonicus* strain ACA-DC 198 was isolated from naturally fermented Greek Kasseri cheese. The strain was grown at 37°C in MRS broth (Oxoid Ltd., Basingstoke, Hampshire, United Kingdom) supplemented with 200 mM morpholinepropanesulfonic acid (MOPS) and adjusted to pH 7.0 with 5 N NaOH (MRS-MOPS). MOPS has been employed previously to maintain neutral pH in batch-type cultures for the study of acid tolerance (26, 45). In this medium, *S. macedonicus* exhibited better growth with the pH of the culture remaining above 6.8 until late logarithmic phase (data not shown) than in unbuffered MRS adjusted to pH 7.0 or in M17 medium. Stock cultures were maintained on Preserver beads (Technical Service Consultants, Heywood, Lancashire, United Kingdom) at 80°C.

**AT phenotypes and acidic lethal challenge.** Different treatments of *S. macedonicus* were performed in order to reveal possible AT phenotypes (Fig. 1). For induction of the logarithmic-phase acid tolerance response (L-ATR), cells which were subcultured in MRS-MOPS were inoculated into fresh MRS-MOPS, grown until mid-log phase (approximately 10⁹ cells; optical density [OD], ~0.6), recovered by centrifugation, and transiently exposed in MRS, pH 6.0 or 5.5, for 1 h at 37°C. Alternatively, cells were inoculated in unbuffered MRS, pH 7.0, and left to progressively acidify the medium through glycolysis until mid-log phase (approximately 10⁸ cells; OD, ~0.6). For acid habituation, both subculture and final growth until mid-log phase were carried out in MRS, pH 6.0 (approximately 5 × 10⁸ cells; OD, ~1.2). Furthermore, acid-habituated cells were subcultured twice in MRS-MOPS, and mid-log-phase cells of the second subculture were assessed for possible retention of L-ATR. To investigate stationary-phase ATR, cells were obtained after an overnight subculture and final growth for 12, 24, and 48 h in MRS-MOPS (approximately 10¹⁰ cells; OD, ~1.2).

In all cases, untreated and treated cells were harvested by centrifugation and resuspended in MRS, pH 3.5 (adjusted with HCl). Samples of 100 μl were removed aseptically after 15 or 30 min (lethal challenge), instantly diluted in 900 μl of buffered peptone water, pH 7.2 (Merck, Darmstadt, Germany), to sharply stop the effect of the acidic environment, centrifuged, and resuspended in 100 μl phosphate-buffered saline (PBS), pH 7.0, for subsequent labeling with fluorochromes.

**Treatment with inhibitors during induction of L-ATR.** Antibiotics (purchased from Sigma, St. Louis, MO) were used at the following final concentrations: 100 μg/ml chloramphenicol, 40 μg/ml actinomycin D, 40 μg/ml cefulisin, 10 μg/ml vancomycin, 10 μg/ml penicillin, and 0.2 or 0.4 mM N,N′-dicyclohexylcarbodi-imide (DCCD) (see below). Typically, antibiotics were added to the culture 20 min prior to and for the 30 min of acid adaptation at pH 5.5. DCCD was added at a low concentration (0.2 mM) 20 min prior to and for the 60 min of acid adaptation at pH 5.5 or at a high concentration (0.4 mM) during only the last 40 min of acid adaptation performed under the same conditions. All samples were challenged at pH 3.5 for 15 min. Final concentrations of antibiotics were higher than the MICs since cells were exposed to the inhibitors for relatively short periods of time. In all cases, control unchallenged cultures were examined and it was ensured that none of the antibiotics exhibited any bactericidal effect on its own during the treatment.

**F-ATPase and glucose-specific PEP-PTS assays.** F-ATPase and glucose-specific PEP-PTS activities were assayed in permeabilized cells following protocols previously described for other streptococcal species (21, 42). In detail, 40 ml of culture to be assayed was washed once and resuspended in 1.8 ml membrane buffer (75 mM Tris, pH 7.0, 10 mM MgSO₄). Toluene was added to a concentration of 10% (vol/vol), and the suspension was vortexed for 30 s and subjected to two rounds of freeze-thawing. Cells were collected, resuspended in 1 ml

**FIG. 1.** Experimental design for the study of the *S. macedonicus* acid tolerance response. The sequence of subcultures and incubation conditions that led to nonadapted cells, autoacidified cells, acid-adapted cells (by transient exposure to nonlethal acidic pH), acid-habituated cells, cells reversed from acid habituation, and stationary-phase cells is presented schematically. o/n, overnight.
membrane buffer, and aliquoted in 100-μl samples. F-ATPase activity was assayed in terms of the release of inorganic phosphate in 50 mM Tris-maleate buffer, pH 7.0, containing 10 mM MgSO4, and 50 μM ATP. Sonication was assayed by the malachite green method, using a commercially available kit (R&D Systems, Inc., Minneapolis, MN). Glucose-specific PEP-PTS activity of permeabilized cells was assayed by the reaction of NADH in the presence of lactate dehydrogenase and pyruvate that was produced from PEP in response to the addition of glucose. Permeabilized cells were incubated in 100 mM Tris-maleate buffer, pH 7.0, 20 mM MgCl2, 1 mM NaF, 40 mM glucose, and 5 mM PEP for 30 min. Two hundred microliters of cleared supernatant at 0 and 30 min were removed and mixed with 300 μl of double-distilled water. Five hundred microliters of NADH solution (0.21 mM/ml NaOH, 1.5 M Tris, pH 7.0, 0.021% NaN3) and 3 μl of lactate dehydrogenase were added to the mixture. The utilization of NADH was recorded at 340 nm. The difference in the amounts of NADH between time zero and 30 min reflects the amount of glucose phosphorylated. In both F-ATPase and glucose-specific PEP-PTS assays, total protein was assayed by the Bradford method (Bio-Rad Laboratories, Inc., Munich, Germany).

**RESULTS**

Table phenotypes of *S. macedonicus* as assessed by flow cytometry. The ability of *S. macedonicus* to develop ATR was investigated. Control mid-log-phase cells grown in MRS-MOPS as well as mid-log-phase cells which were (i) transiently exposed to MRS, pH 6.0 or 5.5, for 1 h prior to lethal challenge, (ii) autoacidified in unbuffered MRS, pH 7.0 (pH 5.8 ± 0.2 at the time of sampling), or (iii) acid habituated (AH) in MRS, pH 6.0 (pH 5.3 ± 0.1 at the time of sampling), were exposed to MRS, pH 3.5, for 15 min. Unstressed and heat-killed mid-log-phase cells were always analyzed in parallel as additional controls. The differences in the patterns of fluorescent subpopulations after lethal challenge were compared to elucidate possible induction of L-ATR (Fig. 2A). ATR would be expected to increase the percentage of the cFDA^-PI- (viable) and/or cFDA^-PI+ (injured) subpopulation, accompanied by a decrease of the percentage of the cFDA^-PI+ (dead) subpopulation after treatment, in comparison to the control (25). Indeed, all treatments resulted in protection of *S. macedonicus* against lethal challenge (Tukey’s test; α = 0.05) (Fig. 2B).

L-ATR phenotypes acquired after transient exposure to non-lethal acidic pH (pH 6.0 or pH 5.5) and acid habituation resulted in the highest level of (but equal among them) protection, while autoacidification protected *S. macedonicus* cells to a lesser extent (Tukey’s test; α = 0.05). Increasing the time of lethal challenge from 15 to 30 min revealed that all treatments protected *S. macedonicus* to the same extent (Tukey’s test; α = 0.05) (Fig. 2C). Further exposure of the cells at pH 5.5 for 45 min resulted in all samples being almost 100% dead by cytometry (data not shown), indicating that none of the treatments could lead to any rescue of *S. macedonicus*, at least for the detection limits of the analysis. In order to investigate whether acid habituation of *S. macedonicus* in MRS, pH 6.0, led to the selection of acid-resistant mutants, cells of this cul-
FIG. 2. (A) Flow cytometric analysis of S. macedonicus acid tolerance at mid-log phase. Nonadapted (a) or acid-adapted (b to f) cells, after exposure to pH 3.5 for 15 min, were labeled with cFDA and PI and analyzed by flow cytometry. Prior to acid challenge, bacteria were adapted either by autoacidification (b), transient exposure to pH 6.0 and 5.5 (c and d), or acid habituation at pH 6.0 (e). Acid habituation was also reversed after regrowth of habituated cells in MRS-MOPS (f). As additional controls, unstressed (g) and heat-killed (h) mid-log-phase cells were analyzed in parallel. (B) Percentages (means ± standard deviations) of fluorescent subpopulations of the samples presented in panel A. (C) Percentages (means plus standard deviations) of fluorescent subpopulations of samples challenged at pH 3.5 for 30 min in the same order as that presented in panel A.
turance decreased along with the time cells resided in stationary-phase cultures at 12 and 24 h were more resistant to challenge, acid tolerance was evident within the first 10 min of acid challenge, as determined by flow cytometry of cultures prior to (unchallenged) or after (challenged) exposure to MRS, pH 3.5, for 15 min. The number of cFDA-PI staining (data not shown). The presence of the inhibitors prior to and during the process of acid adaptation radically influenced the ability of S. macedonicus mid-log-phase cells to launch a full L-ATR, as judged by their aptitude for overcoming lethal challenge (Fig. 4). In detail, all antibiotics except for vancomycin resulted in partial inhibition of the L-ATR, as the increase of viable cells and decrease of dead cells in comparison to those of the nonadapted control did not match the magnitude of the corresponding changes observed for the adapted control (Tukey’s test; α = 0.05). Most interestingly, partial inhibition of L-ATR was accompanied by a significant increase in injured cells in comparison to both nonadapted and adapted controls (Tukey’s test; α = 0.05). Vancomycin was the only inhibitor that completely abolished L-ATR and rendered cells more susceptible to lethal challenge in comparison to the nonadapted control, since the numbers of both viable and injured cells decreased, while the number of dead cells marginally increased (Tukey’s test; α = 0.05). Vancomycin has been shown to inhibit cell wall biosynthesis by being incorporated into nascent peptidoglycan chains (15). The commercial availability of the fluorescent derivative BODIPY FL vancomycin allowed us to assess possible changes in the cell wall formation profile due to acid adaptation. Flow cytometric analysis and CLSM did not reveal any obvious differences between nonadapted and acid-adapted (at pH 5.5) populations (Fig. 5A and B). Interestingly, though, acid-habituated (at pH 6.0) cells exhibited an almost 1.5-fold increase of the mean side scatter intensity and a >2.5-fold increase of the mean fluorescence intensity in comparison to nonadapted cells (Fig. 5A). Microscopic examination demonstrated that acid habituation resulted in the formation of long cell chains coin-
ciding with the increase of the side scatter intensity, while peptidoglycan synthesis occurred at nearly every division septum and cell equator (Fig. 5B). Nongrowing stationary-phase cells could not be labeled with BODIPY FL vancomycin, verifying its specificity for active sites of cell wall biosynthesis for *S. macedonicus* (Fig. 5A).

**L-ATR leads to retention of MP of *S. macedonicus* cells after lethal challenge.** DiBAC<sub>4</sub>(3) ratiometric analysis was em-

![Graph A](image1)

**FIG. 3.** Kinetics of acid adaptation of *S. macedonicus*. Mid-log-phase cells were adapted to pH 5.5 for 0 to 60 min and exposed to lethal challenge for 15 or 30 min (A and B, respectively). Percentages (means ± standard deviations) of fluorescent subpopulations, as determined by flow cytometry after cFDA-PI labeling, are presented.

![Graph B](image2)

**FIG. 4.** Effects of inhibitors on the induction of acid tolerance in *S. macedonicus*. Prior to acidic lethal challenge at pH 3.5 for 15 min, mid-log-phase cells were either left unadapted (control) or acid adapted at pH 5.5 for 30 min in the presence of chloramphenicol (Chl), actinomycin D (Act D), cerulenin (Cer), vancomycin (Van), or penicillin (Pen) or in the absence of any inhibitor (adapted). Percentages (means ± standard deviations) of fluorescent subpopulations, as determined by flow cytometry after cFDA-PI labeling, are presented.
ployed to compare the MP of nonadapted and acid-adapted cells after acidic lethal challenge. The log ratio of green fluorescence to side scatter enhanced the resolution of live, depolarized, and dead cells (Fig. 6) compared to that we previously reported (41), since this type of analysis corrects for DiBAC₄(3) fluorescence variations due to cell size heterogeneity (2) or cell clumping (38). The membrane potential of nonadapted cells was completely dissipated after lethal challenge, giving a single peak of the ratio parameter with a mean intensity higher than that of depolarized cells and similar to that of heat-killed cells (Fig. 6, peak d). In contrast, acid-adapted cells were separated into three subpopulations after lethal challenge (Fig. 6, peak e). The majority of the cells retained normal membrane potential (55.8%), while the other two groups coincided with either the depolarized or the heat-killed population (19.0% and 25.2%, respectively). These results indicate that induction of L-ATR enables cells to resist the abolishment of membrane potential caused by lethal challenge.

**Involvement of F-ATPase and glucose-specific PEP-PTS activities in L-ATR of *S. macedonicus***. DCCD was employed as an inhibitor of F-ATPase. Mid-log-phase *S. macedonicus* cells were exposed for 1 h to pH 5.5, and DCCD was added either at 0.2 mM 20 min prior to and during the 1 h of acid adaptation or at 0.4 mM for only the last 40 min of treatment (Fig. 7). Inhibition of F-ATPase activity throughout acid adaptation resulted in abolishment of L-ATR, as lethal challenge resulted in >99% dead cells. In contrast, adapting *S. macedonicus* cells for 20 min before inhibiting F-ATPase activity (for the last 40 min at pH 5.5) allowed a partial L-ATR. DCCD had no effect on the viability of control unchallenged cultures (data not shown). Additionally, since DCCD causes covalent modification of F-ATPase (28), it should be presumed that inhibition persisted throughout lethal challenge. All of these data indicate an important role for F-ATPase during the induction of L-ATR, but its activity appears to be dispensable to a degree once L-ATR has been launched fully (i.e., after exposure for 20 min at pH 5.5).

The activities of F-ATPase and the glucose-specific PEP-PTS were further assessed in permeabilized *S. macedonicus* cells (Fig. 8A and B, respectively). F-ATPase activity of acid-adapted cells (by transient exposure to pH 5.5) and AH mid-log-phase cells increased 2.6-fold compared to that of nonadapted cells, while for autoacidified cells there was a lesser,
2-fold increase (Tukey’s test; \( \alpha = 0.05 \)). Glucose-specific PEP-PTS activity of \emph{S. macedonicus} increased to the same extent for all three AT phenotypes (>3.6-fold). Acid-adapted cells in the presence of chloramphenicol had the same F-ATPase and PEP-PTS activities as control cells. While it has been reported that cell membrane fatty acid composition influences both F-ATPase and PEP-PTS activities (21), in our case a marginal, not statistically significant decrease of both enzymatic activities was observed when cells were treated with cerulenin during acid adaptation.

**Single-cell sorting of \emph{S. macedonicus} subpopulations according to their cFDA-PI fluorescence status.** In our previous work on acid-stressed cells of \emph{S. macedonicus} (41), we already found that the cFDA-positive cells detected after acid challenge showed recovery when sorted onto conventional agar plates. In order to investigate the effect of acid adaptation on the culturability of \emph{S. macedonicus} cFDA-PI fluorescent subpopulations, nonadapted and transiently acid-exposed (to pH 5.5 for 30 min) mid-log-phase cells were challenged at pH 3.5 for 15 min, labeled with the fluorochromes, and sorted onto both optimal and suboptimal media. Unchallenged control mid-log-phase cells showed recovery above 96% in all different medium types. This indicated that neither sonication treatment nor sorting had any serious impact on the cells’ culturability and that the suboptimal media used only delayed growth (data not shown). Additionally, sorted cFDA\(^+\) PI\(^+\) cells were unable to recover in all cases (data not shown), as previously reported (41). Systematically, cFDA\(^+\) PI\(^-\) cells exhibited a higher capacity to form colonies than did the corresponding cFDA\(^+\) PI\(^+\) cells (binomial comparison of proportions; \( P < 0.05 \)) (Fig. 9). Most importantly, shifting the pH of the sorting medium to 6.0 or 5.5 caused a drastic decrease in the recovery of both cFDA\(^+\) PI\(^-\) and cFDA\(^+\) PI\(^+\) cells derived from nonadapted samples. In contrast, the recovery of the two subpopulations for acid-adapted cells was not considerably influenced under such conditions. Furthermore, sorting cFDA\(^+\) PI\(^-\) and cFDA\(^+\) PI\(^+\) cells onto MRS-MOPS agar plates supplemented with 1% NaCl revealed that the observed differences in the recovery of

![FIG. 6. Ratiometric analysis of membrane potential changes in \emph{S. macedonicus} cells after acidic lethal challenge with DiBAC\(_4\)(3). Untreated (a), depolarized (b), and heat-killed (c) mid-log-phase \emph{S. macedonicus} cells served as controls. Nonadapted (d) or acid-adapted (at pH 5.5 for 30 min) (e) mid-log-phase cells were exposed to pH 3.5 for 15 min. All samples were labeled with DiBAC\(_4\)(3) and analyzed by flow cytometry. Results are presented as single-parameter histograms of the log ratios of green fluorescence to side scatter. M1, M2, and M3 were defined by the untreated, depolarized, and heat-killed cells’ distributions, respectively.](http://aem.asm.org/)

![FIG. 7. Effect of DCCD on the induction of acid tolerance in \emph{S. macedonicus}. Prior to lethal challenge, cells were nonadapted (a), acid adapted at pH 5.5 for 60 min, with DCCD being added either 20 min prior to and during the 60 min of acid adaptation (b) or only during the last 40 min of the treatment (c), and acid adapted under the same conditions in the absence of DCCD (d). Percentages (means ± standard deviations) of fluorescent subpopulations, as determined by flow cytometry after cFDA-PI labeling, are presented.](http://aem.asm.org/)
these two subpopulations between nonadapted and acid-adapted samples were not solely dependent on the pH of the sorting medium. In summary, these results show that subpopulations identical in fluorescence characteristics may differ significantly in their potential to recover.

**DISCUSSION**

In this study, we employed flow cytometry along with different fluorescence-based assays to characterize the acid stress responses of *S. macedonicus*. Our data clearly show that all three manipulations imposed on mid-log-phase *S. macedonicus* cells resulted in induction of L-ATR. In detail, transient exposure to nonlethal acidic pHs (pH 6.0 and pH 5.5) and continuous growth at suboptimal pH 6.0, i.e., acid habituation (first described for *Escherichia coli* [24]) of mid-log-phase cells, prior to lethal challenge, resulted in equal protection, making these phenotypes indistinguishable in terms of their survival. Interestingly, the acidifying capacity of *S. macedonicus* also stimulated a protective response. This type of L-ATR was weaker than those in response to the other treatments and could probably be ascribed to physiological changes similar to those described for *Lactococcus lactis* autoacidified cells (19, 46). Stationary-phase ATR was also verified, constituting a growth phase resistance pattern for *S. macedonicus*. The response decayed progressively along with the time that the cells resided at stationary phase. Apparently, prolonged exposure to pH 4.3 (i.e., the pH value at stationary phase) caused the degeneration of nongrowing *S. macedonicus* cells, making them more susceptible to low pH. This was also evident by the increasing numbers of both injured and dead cells exhibited by the control stationary-phase cultures. Similar findings have been reported for *S. mutans* (47) and *L. lactis* (37). Our batch-type culture system cannot rule out the compensation of low pH for the induction of the observed stationary-phase ATR, which has been shown to be essential in the case of *Lactobacillus acidophilus* (32). All the same, *S. mutans* and *L. lactis* grown at constant pH have also been reported to exhibit stationary-phase ATR (1, 49).

Subsequently, flow cytometric analysis of acid-stressed *S. macedonicus* cells following acid adaptation revealed the heterogenic behavior of the bacterium, similar to that of *Oenococcus oeni* adapted to ethanol stress (25). We demonstrated not only that *S. macedonicus* cells differ in the ability to induce L-ATR but that there are also cell-to-cell deviations in the rate
of this induction. Such an individualized response of a population to an external stimulus (in this case, exposure to sublethal pH) has been linked to asynchronous cell cycle progression, differences in cell physiological status, and stochastic variations (6, 9).

We then employed different inhibitors to elucidate the possible involvement of central cellular mechanisms in the manifestation of acid tolerance of mid-log-phase *S. macedonicus* cells. L-ATR was partially inhibited by both chloramphenicol and actinomycin D, suggesting that part of the acid resistance phenotype is dependent on de novo protein synthesis that seems to be the result of a genetically programmed response. Indeed, chloramphenicol has been shown to inhibit induction of L-ATR in *L. lactis* (46), *Listeria monocytogenes* (39), and *Lactobacillus sanfranciscensis* (17). In addition, acid-adapted oral streptococci are enriched in membrane long-chain monounsaturated fatty acids in order to decrease proton membrane permeability (20–22). Inhibition of fatty acid biosynthesis of *S. macedonicus* by cerulenin caused a decrease of L-ATR to some extent. Furthermore, cell wall formation proved to be important for the induction of L-ATR, as vancomycin and penicillin led to severe inhibition of resistance to lethal challenge. Previous work with mutant strains of *S. mutans*, *L. monocytogenes*, and *Enterococcus faecalis* have also indicated that proper cell wall biosynthesis is necessary for acid resistance (8, 12, 50). Further investigation with fluorescent vancomycin that labeled sites of nascent peptidoglycan biosynthesis revealed that acid-adapted cells exhibited an abnormal pattern of active cell wall biosynthesis sites, which correspond to recent or forthcoming division sites (15). This observation may indicate severe perturbations of cell cycle progression due to continuous growth at low pH and is in agreement with the fact that the FtsZ and FtsA proteins (which belong to the cell division machinery and are located at septum formation sites) were found to be up-regulated when *S. mutans* was grown at pH 5.2 (54). The increase in the length of cell chains of AH cells could be attributed to impairment of enzymatic activities that are necessary for cell segregation after division due to low pH, similar to those described for *S. mutans*, *S. pneumoniae*, and *S. thermophilus* (7, 11, 31). It should also be mentioned that the increase of the length of cell chains due to acid habituation may be an advertent adaptation response, as it has been shown that dechained *S. mutans* cells were less aciduric than chained ones (27) and that chains can give growth advantages to *S. thermophilus* under aerobic conditions (34).

Since acid-challenged mid-log-phase *S. macedonicus* cells suffer severe cell membrane damage and since acid adaptation causes changes to the cell membrane composition in an attempt to reduce the extensive proton influx (22), the perturbation of cell MP was examined as a possible consequence of acid stress. After acid challenge, nonadapted cells exhibited completely dissipated MP. Significantly, under the same conditions, the majority of acid-adapted cells retained normal MP. This difference clearly indicates that physiological mechanisms induced during acid adaptation actively increase the ability of cells to maintain ion gradients across the cell membrane during acid challenge. Since the MP of streptococcal cells is mainly attributed to the proton-translocating F-ATPase (29) and since...
it has been reported that several streptococci, as well as L. lactis, cope with high extracellular proton concentrations by increased F-ATPase activity (3, 40), we investigated the involvement of this enzymatic system in L-ATR of S. macedonicus. Our findings strongly suggest that F-ATPase activity is important during the induction of L-ATR, probably in order to maintain the proper pH gradient (ΔpH) necessary for the function of key cellular components needed for the induction of the response. However, once L-ATR is fully induced, F-ATPase activity is only partially required, since other physiological changes that already have occurred may aid the survival of cells during lethal challenge (i.e., newly synthesized proteins, changes in the cell envelope, etc.). In favor of this reasoning, nonadapted mid-log-phase L. monocytogenes cells were found to be more susceptible to acid killing after DCCD treatment than were acid-adapted cells (13). It was further determined that both F-ATPase and glucose-specific PEP-PTS activities increased in all three logarithmic-phase AT phenotypes in comparison to those in nonadapted cells. The increase of F-ATPase activity in autoclaved cells was lower than that for the other two phenotypes and probably reflects the decreased acid tolerance of this phenotype. The up-regulation of both enzymatic systems in S. macedonicus was due to de novo protein synthesis, similar to the case for other streptococci (30, 33, 35, 44). In addition, a fabM deletion mutant of S. mutans which is unable to produce unsaturated fatty acids was sensitive to low pH and exhibited significantly increased F-ATPase and glucose-specific PEP-PTS activities (21). In the case of S. macedonicus, inhibition of fatty acid biosynthesis by cerulenin did not influence the increased activities of F-ATPase and glucose-specific PEP-PTS observed during acid adaptation. Up-regulation of the glucose-specific PEP-PTS under acid adaptation conditions may be necessary for the increased glucose intake required to compensate for the depletion of ATP pools caused by enhanced F-ATPase activity (35). To the best of our knowledge, S. macedonicus is the first Streptococcus species reported to induce both enzymatic activities during acid adaptation. This feature demonstrates that S. macedonicus responds differently to acidic conditions than do the majority of oral streptococci, which typically up-regulate F-ATPase and down-regulate the glucose-specific PEP-PTS (44), or S. sobrinus, which up-regulates only the glucose-specific PEP-PTS and retains a stable F-ATPase level (35).

Previous sorting experiments attempted to establish a direct correlation between the fluorescence status of stressed bacteria after cFDA-PI labeling and culturability, i.e., cFDA-PI, cFDA-PI, and cFDA-PI events have been reported to correspond to viable, injured, and dead cells, respectively (2, 25). The reduced recovery of the sorted cFDA-PI cells compared to that of cFDA-PI. S. macedonicus cells on optimal medium is in accordance with these findings. Sorting cFDA-PI and cFDA-PI cells of nonadapted acid-stressed S. macedonicus samples onto suboptimal media demonstrated that cells referred to as injured by culture-based techniques (i.e., cells that are able to replicate under optimal conditions but unable to replicate under suboptimal, nonlethal conditions) exist in both populations. In contrast, recovery of these two fluorescent subpopulations derived from acid-adapted samples after acid challenge remained more or less unaffected by the different recovery conditions, thus precluding the existence of injured cells in culture-based terms in both subpopulations. This leads us to the conclusion that classical injury, as detected by the ability of a cell to grow under adverse conditions, may not be reflected fully by cFDA-PI labeling. Finally, while S. macedonicus cells can be separated into three discrete stages of activity and permeability by cFDA-PI labeling after lethal challenge, our findings strongly indicate that the recovery potential of any given fluorescent subpopulation may be influenced by the physiological status of the cells prior to lethal challenge. This was evident because the regrowth patterns of nonadapted and acid-adapted cells were substantially different.

Here we elucidated traits of the hitherto unknown acid stress physiology of S. macedonicus. Multiparameter flow cytometry combined with single-cell sorting revealed effectively diverse physiological changes that S. macedonicus underwent during acquisition of acid tolerance. Our findings should facilitate a better understanding of the physiology of food-related lactic streptococci that will allow their rational utilization in industrial processes. We will continue to refine this approach, along with molecular techniques, such as RNA arbitrarily primed PCR, to further establish possible differences among the different AT phenotypes we described and to determine their relevance to the food environment.

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