Identification of the Streptococcus mutans LytST two-component regulon reveals its contribution to oxidative stress tolerance

Sang-Joon Ahn¹, Ming-Da Qu², Elisha Roberts², Robert A Burne¹ and Kelly C Rice²*

Abstract

Background: The S. mutans LrgA/B holin-like proteins have been shown to affect biofilm formation and oxidative stress tolerance, and are regulated by oxygenation, glucose levels, and by the LytST two-component system. In this study, we sought to determine if LytST was involved in regulating lrgAB expression in response to glucose and oxygenation in S. mutans.

Results: Real-time PCR revealed that growth phase-dependent regulation of lrgAB expression in response to glucose metabolism is mediated by LytST under low-oxygen conditions. However, the effect of LytST on lrgAB expression was less pronounced when cells were grown with aeration. RNA expression profiles in the wild-type and lytS mutant strains were compared using microarrays in early exponential and late exponential phase cells. The expression of 40 and 136 genes in early-exponential and late exponential phase, respectively, was altered in the lytS mutant. Although expression of comYB, encoding a DNA binding-uptake protein, was substantially increased in the lytS mutant, this did not translate to an effect on competence. However, a lrgA mutant displayed a substantial decrease in transformation efficiency, suggestive of a previously-unknown link between LrgA and S. mutans competence development. Finally, increased expression of genes encoding antioxidant and DNA recombination/repair enzymes was observed in the lytS mutant, suggesting that the mutant may be subjected to increased oxidative stress during normal growth. Although the intracellular levels of reaction oxygen species (ROS) appeared similar between wild-type and lytS mutant strains after overnight growth, challenge of these strains with hydrogen peroxide (H₂O₂) resulted in increased intracellular ROS in the lytS mutant.

Conclusions: Overall, these results: (1) Reinforce the importance of LytST in governing lrgAB expression in response to glucose and oxygen, (2) Define a new role for LytST in global gene regulation and resistance to H₂O₂, and (3) Uncover a potential link between LrgAB and competence development in S. mutans.

Keywords: Stress, Oxygen, Competence, Cid/Lrg system, Streptococcus mutans

Background

Streptococcus mutans is considered the primary causative agent of dental caries, and when transiently introduced into the bloodstream following daily dental hygienic practices such as toothbrushing and flossing, this bacterium can also cause potentially lethal infective endocarditis (IE) [1-4]. In both infectious scenarios, the virulence of S. mutans depends upon its ability to form biofilms and to withstand extreme changes in environmental conditions, including fluctuations in oxygen, shear stress, as well as nutrient source and availability. For example, in the oral cavity, S. mutans must be able to rapidly alter its expression of transporters and metabolic enzymes to catabolize a variety of host-derived dietary carbohydrates. Internalized carbohydrates are metabolized through the glycolytic pathway, resulting in the accumulation of acidic end-products in the environment, which favors the growth of S. mutans and other acid-tolerant cariogenic species. Repeated cycles of acidification can lead to a net demineralization of tooth enamel and the development of caries. Sucrose, a common dietary sweetener, can also be utilized by...
S. mutans for the production of extracellular polysaccharides [5-8] that facilitate bacterial adhesion and biofilm formation. Aeration has also been found to have a profound effect on carbohydrate metabolism and biofilm formation by S. mutans [9-11]. It is therefore not surprising that there is overlap in the genetic regulatory circuits responsive to carbohydrate metabolism, aeration/oxidative stress resistance and control of biofilm formation in S. mutans, which include CcpA [12-14], Rex [15], and Frp [16].

More recently, an emerging trend in the study of bacterial biofilms has been a focus on the contribution of bacterial cell death and autolysis to biofilm adherence, maturation, and dispersal. It has been demonstrated in a wide variety of bacteria that death and lysis of a subpopulation of cells can facilitate biofilm formation due to the release of DNA into the extracellular environment (eDNA) [17-22]. Likewise, cell death and lysis have been implicated in dispersal of cells from a mature biofilm [23-25]. In Staphylococcus aureus, the Cid/Lrg system has been shown to be involved in the regulation of cell death, autolysis, and biofilm formation [17,21,26-28]. Characterization of S. aureus cid and lrg mutants has revealed that these operons have opposing effects on cell death and murein hydrolase activity [27,29]. These observations, combined with the fact that LrgA and CidA share structural features with the bacteriophage lambda family of holin proteins [29], have led to the hypothesis that CidA and LrgA control cell death and lysis in a manner analogous to effector and inhibitor holins, respectively [26,30]. Bacteriophage holins are small membrane proteins that oligomerize in the cell membrane, acting as “molecular clocks” that regulate the timing and lysis of the host cell during lytic infection [31]. For example, the lambda S holin regulates cell death and lysis by the formation of large lipid-excluding “rafts” that promote cytosolic leakage as well as access of the phage-encoded endolysin (murein hydrolase) to the cell wall [32-34]. S. aureus CidA and LrgA have recently been shown to oligomerize into high-molecular-mass complexes in a cysteine disulfide bond-dependent manner, a biochemical feature also shared with holin proteins [35]. Although the molecular details of how Cid and Lrg function to control cell death and lysis have not yet been completely elucidated, the fact that cid and lrg homologues have been identified in a wide variety of bacterial and archaean genomes supports a fundamental and conserved role for this system in cell physiology [30,36].

In previous work it was determined that expression of potential cidAB and lrgAB homologues in S. mutans is highly responsive to carbohydrate availability [12,37] and oxygenation [11]. Given the potential importance of these genes to biofilm development in S. mutans, we previously characterized a panel of S. mutans cid and lrg isogenic mutants and found that a subset of these genes did indeed influence biofilm formation, production of glucosyltransferases (enzymes that synthesize extracellular glucan polymers that contribute to biofilm adhesion), and oxidative stress tolerance [37]. In this study it was also found that, as demonstrated previously in S. aureus [38,39], the S. mutans LytST two-component system was required for activation of lrgAB expression, but not cidAB expression [37]. Genes homologous to lytST appear to be present in most Gram-positive organisms that contain lrgAB [30] and these genes are often linked to another, inferring an important role for this two-component system in fine-tuning lrgAB expression in response to external environmental signals. Therefore in this study, we sought to determine if LytST is involved in regulation of lrgAB expression in response to glucose and oxygenation in S. mutans, and to elaborate on the contribution of LytST to cellular homeostasis and global control of gene expression.

Results

Effects of oxygenation and glucose metabolism on S. mutans lrg and cid expression

The LytST two-component regulatory system has been shown to positively regulate lrgAB expression in a wide variety of bacteria, including various staphylococcal [38-40] and Bacillus species [41,42], as well as in S. mutans [37]. The conserved nature of this regulation in Gram-positive bacteria, combined with the known effects of LytST and LrgAB on cell death/lysis [29,38,39,43], biofilm development [21,37,38], and oxidative stress resistance [37], suggests that LytST and LrgAB are central regulators of physiologic homeostasis. However, little is known about the environmental and/or cellular cues to which LytS responds. In S. aureus and B. anthracis, it has been shown that lrgAB expression is responsive to disruption of cell membrane potential in a LytST-dependent manner [41,44]. However, we were unable to determine whether this regulation also occurs in S. mutans, as treatment with membrane-potential disrupting agents (gramicidin, carbonyl cyanide m-chlorophenylhydrazone) did not have a measurable effect on membrane potential, as assessed by staining with DIOC2 (3) [data not shown].

In previous studies, it was shown that oxygen and glucose metabolism have a pronounced effect on lrg and cid expression in S. mutans, but the specific role of LytS, if any, in this regulation was not addressed [11,37]. Therefore, S. mutans UA159 and its isogenic lytS mutant were grown under aerobic and low-oxygen conditions to exponential (EP) and stationary (SP) growth phases in media containing 11 mM or 45 mM glucose. Quantitative real-time reverse transcriptase PCR (qRT-PCR) was performed on RNA isolated from cultures at each time
point to assess changes in _lrg_ expression (Figure 1). In UA159, stationary phase _lrgA_ expression was upregulated 365-fold relative to exponential phase when grown under 11 mM glucose and low-oxygen conditions (Figure 1A). Although mutation of _lytS_ resulted in a severe loss of stationary phase _lrgA_ induction in cells grown in 11 mM glucose, _lrgA_ expression was not completely abolished. When grown under aerobic conditions and 11 mM glucose, stationary phase _lrgA_ expression was upregulated 2500-fold relative to exponential phase in the wild-type strain (Figure 1A), confirming previously-published observations that aerobic growth promotes _lrgA_ expression [11]. However, stationary-phase _lrgA_ expression was still induced 216-fold in the _lytS_ mutant during aerobic growth, suggesting that (1) other as-yet-unknown regulators also contribute to the positive control of _lrgA_ expression during aerated growth, and (2) LytST is a predominant regulator of _lrgA_ expression during low oxygen growth, compared to aerobic growth. Under low-oxygen and aerated cultures, stationary phase induction of _lrgA_ expression was dramatically reduced when grown in 45 mM glucose, and similar levels of expression were observed in the wild-type and _lytS_ mutant (Figure 1B), suggesting that growth in high levels of glucose abrogates oxygen-dependent regulation of _lrgA_ by LytST. Consistent with previously-published data [37], LytS did not appear to have a measurable effect on _cidA_ expression under any of the growth conditions tested here (data not shown). In summary, LytST-dependent regulation of _lrgA_ expression is much more pronounced during low-oxygen growth and at low glucose levels.

### Microarray analysis of the LytS regulon

Based on the transcriptional data presented above, the effects of LytST regulation on _lrgA_ expression are most evident while _S. mutans_ is growing under conditions of low-oxygen (5% CO₂) with a lower concentration of glucose. To begin to explore how LytST impacts critical phenotypes of _S. mutans_, RNA expression profiles in UA159 and the _lytS_ mutant were compared using an RNA microarray approach. RNA was isolated from early exponential and late exponential growth phases from static planktonic cultures grown in BHI (containing 11 mM total glucose) at 37°C in a 5% CO₂ atmosphere (Additional file 1: Table S1 and Additional file 2: Table S2). At early exponential growth phase, loss of LytS affected the expression of 40 genes (12 upregulated and 28 downregulated; _P_ < 0.005; Additional file 1: Table S1). Most of the upregulated genes in early exponential phase displayed only a modest increase in expression and included genes involved in DNA repair, purine/pyrimidine metabolism, competence, and a number of unassigned and hypothetical ORFs. RNA transcripts that were strongly down-regulated greater than 10-fold in cells lacking LytS during early exponential growth included those annotated as bacitracin/surfactin/gramicidin synthesis proteins, transport and binding proteins, and _LrgA_. In contrast, loss of LytS affected the expression of a much larger number of genes in late exponential phase (136 genes total), with 79 upregulated transcripts and 57 downregulated transcripts (_P_ < 0.001; Additional file 2: Table S2). Aside from dramatically decreased _lrgA_ expression, affected genes included those involved in amino acid and co-factor biosynthesis, carbohydrate and fatty acid metabolism, stress adaptation, toxin production, DNA repair/recombination,

### Figure 1

**Figure 1 LytS-dependent expression of lrgA in _S. mutans_.** Overnight cultures were diluted in THYE, containing either 11 mM (A) or 45 mM glucose (B) to an OD₆₀₀ₐₚ = 0.02 and grown at 37°C as static cultures at 5% CO₂ (“low-O₂”) or as aerobic shaking cultures at 250 RPM (“aerobic”). RNA was harvested at exponential (EP) and stationary phase (SP). Reverse-transcription, real-time PCR reactions, and determination of copy number were performed as described previously using _lrgA_ and 16S-specific primers [37,77]. Fold-change expression of _lrgA_ and 16S under each growth condition was calculated by dividing the gene copy number of each test sample by the average gene copy number of UA159 EP. Data was then normalized by dividing each _lrgA_ fold-change value by its corresponding 16S fold-change expression value. Data represent the average of 3 biological replicates. Dark grey bars represent UA159 and light grey bars represent _lytS_ mutant. Error Bars represent the standard error (SEM).
protein synthesis, transcriptional regulation, and competence, as well as multiple hypothetical and/ or unassigned ORFs (Additional file 2: Table S2 and Figure 2). A subset of genes was differentially expressed as a function of the loss of LytS in both early exponential and late exponential growth phases (Additional file 1: Table S1 and Additional file 2: Table S2). These included many genes encoded by the S. mutans genomic island TnSMu2 [45] (SMU.1335c, 1339-1342, 1344c-1346, 1354c, 1360c, 1363c, 1366c), ssbA, comYB, and lrgAB. Given that these genes were regulated by LytS in both growth phases examined, it is possible that they are under the direct control of LytST. To validate the microarray data, qRT-PCR was performed on late exponential phase wild-type and lytS mutant RNA to assess expression of 14 of the affected genes. As shown in Table 1, the expression ratios (lytS mutant/wild-type) for each gene obtained by real-time PCR were similar to the microarray results. Interestingly, expression ratios of these genes were all close to 1.0 when comparing expression between the wild-type strain and a lrgAB mutant (Table 1), indicating that the differential expression patterns observed in the lytS

| Table 1 Real-time PCR validation of RNA microarray results |
|------------------------------------------------------------|
| Gene | Microarray | Real-time PCR |
|------|------------|---------------|
| (SMU.1985) comYA (comYB) | 22.9927 | 6.8449 | 0.8163 |
| SMU.1967 ssbA | 5.5803 | 4.1076 | 0.8791 |
| (SMU.1515) vicR (vicX) | 2.6764 | 1.7647 | 1.0267 |
| SMU.924 tpx | 2.4148 | 3.6168 | 1.058 |
| SMU.1739 fabF | 2.2443 | 2.0333 | 1.084 |
| SMU.1666 livG | 2.1183 | 3.4331 | 1.099 |
| SMU.80 hrcA | 0.4953 | 0.6107 | 1.0204 |
| SMU.1424 pdhD | 0.4769 | 0.4031 | 1.2004 |
| SMU.580 xseA | 0.29849 | 0.5409 | 1.1398 |
| SMU.1600 celB | 0.2186 | 0.2825 | 1.2979 |
| SMU.113 pfk | 0.1597 | 0.176 | 1.3578 |
| SMU.82 dnaK | 0.1523 | 0.2652 | 0.9907 |
| SMU.1344 fabD | 0.0223 | 0.012 | 1.0637 |
| SMU.1341 grs | 0.0008 | 0.0121 | 1.1027 |

Results are expressed in fold-change (mutant/wild-type).

**Figure 2** Distribution of functions of genes affected by loss of LytS at late exponential phase. Statistical analysis was carried out with BRB array tools (http://linus.nci.nih.gov/BRB-ArrayTools.html/) with a cutoff P value of 0.001. The 136 genes differentially expressed at P ≤0.001 are grouped by functional classification according to the Los Alamos S. mutans genome database (http://www.oralgen.lanl.gov/).
mutant were not a consequence of down-regulated lrgAB expression.

Investigation of the effect of LytST and LrgAB on competence
In analyzing the microarray data in Additional file 1: Table S1 and Additional file 2: Table S2, it appeared that the gene most highly upregulated in response to loss of LytS in both phases of growth was comYB (SMU.1985), a homologue of the B. subtilis comGB gene that encodes part of an ABC transporter essential for DNA binding-uptake during competence in S. mutans [46]. Interestingly, a comYB mutant of S. mutans was shown to be unaffected in competence signaling, but showed reduced biofilm formation, which was thought to be a function of its inability to bind biofilm matrix eDNA [47]. Since the lytS mutant displayed an increase in comYB expression (Additional file 1: Table S1 and Additional file 2: Table S2), we hypothesized that this strain may display alterations in its ability to form biofilm and/or its transformability during genetic competence. However, the lytS mutant did not display any appreciable difference in its ability to form static biofilm in the presence of glucose or sucrose (data not shown), and likewise, did not display a difference in its ability to uptake plasmid DNA in a quantitative competence assay, relative to the wild-type strain (Figure 3). Since lrgAB expression is so strongly regulated by LytST, the ability of isogenic lrgA, lrgB, and lrgAB mutants to uptake plasmid DNA via competence was also assessed (Figure 3). Of all the mutants tested, the lrgA mutant was the most severely impaired in its ability to uptake plasmid DNA relative to the parental strain, displaying a 26- and 24-fold decrease in transformation efficiency in the presence and absence of competence-stimulating peptide (CSP), respectively (Figure 3), suggesting that LrgA is somehow involved in genetic transformation in a CSP-independent manner. This finding has particular significance considering that LrgAB has been linked to regulation of cell death and lysis in S. aureus [21,29] and S. mutans [37], and these physiological processes are also extremely important during natural competence. It is interesting to note that, similar to the competence results described here, the lrgA mutant was previously shown to display decreased glucose-dependent biofilm formation and decreased glucosyltransferase production, whereas the lrgB and lrgAB mutants behaved in a manner similar to the parental strain [37]. These phenotypic patterns suggest that the presence of LrgB alone, rather than the lack of LrgA, may be responsible for the biofilm and competence phenotypes observed in the lrgA mutant.

Effect of LytST on oxidative stress tolerance
Previously, our investigations disclosed a strong link between oxidative stress tolerance and the Cid/Lrg system [37], a role for these genes that had not been described in other organisms. Specifically, we found that lrgAB, lrgB, cidAB, and cidB mutants exhibited reduced growth in the presence of paraquat, and growth of lrgAB, cidAB, and cidB mutants on BHI agar plates in aerobic conditions was almost completely inhibited [37]. It is therefore interesting to note that in the lytS microarray results (Additional file 2: Table S2), genes encoding antioxidant and DNA repair/recombination enzymes were significantly upregulated in the lytS mutant in late exponential phase. These included yghU and tpx, encoding the putative anti-oxidant enzymes glutathione S-transferase and thiol peroxidase, respectively, as well as recA, which encodes a single-stranded DNA exonuclease protein that facilitates DNA repair in response to oxidative stress [48-51]. To further investigate the effect of lytS and lrgAB on oxidative stress tolerance, wild-type, lytS, and lrgAB mutants were grown as planktonic static BHI cultures in aerobic atmosphere and in the presence and absence of H2O2 (Figure 4). When challenged with H2O2, UA159 experienced an increased lag phase of growth, and the overall OD of the culture was 10-25% less than the untreated culture until 20 h growth. Under these assay conditions, the lrgAB mutant displayed a
dramatic growth defect in both the presence and absence of H$_2$O$_2$. It is interesting to note that this aerobic growth defect was also previously observed when the lrgAB mutant was grown in aerobic atmosphere on BHI agar plates [37]. The lrtS mutant displayed an increased lag in growth relative to UA159 when cultured in the presence of H$_2$O$_2$, but OD values were comparable to the wild-type strain by 16 h growth. These results suggest that the LytST regulon impacts the ability of cells to grow under conditions of oxidative stress.

The cell-permeable fluorescent dye CM-H$_2$DCFDA (Invitrogen Molecular Probes) was also used to assess intracellular ROS in UA159 and the lrtS mutant (Figure 5). This fluorescent compound is oxidized in the presence of H$_2$O$_2$ and other reactive oxygen species (ROS) and is considered a general indicator of intracellular oxidative stress [52,53]. This analysis revealed that stationary-phase cultures of the wild-type and lrtS mutant strains had similar “endogenous” intracellular levels of ROS (Figure 5, light grey bars). When stationary-phase cells from each strain were loaded with CM-H$_2$DCFDA and then challenged with 5 mM H$_2$O$_2$ (Figure 5, dark grey bars), a greater increase in fluorescence was observed in the lrtS mutant relative to UA159 ($P = 0.009$, Mann–Whitney Rank Sum Test), suggesting that loss of LytS has an impact on the ability of the cells to detoxify H$_2$O$_2$ and/or other intracellular ROS.

**Discussion**

The transcriptome analyses presented in this study have revealed that the LytST two-component system has a widespread effect on gene expression in *S. mutans*. A much higher number of transcripts were affected by the lrtS mutation in late exponential phase and the magnitude of changes in expression was greater ($n = 136$ genes, Additional file 2: Table S2) relative to early-exponential phase ($n = 40$ genes, Additional file 1: Table S1), where most genes exhibited only a modest (1-2 fold) change in expression. These differences in gene expression patterns are unlikely to be an indirect function of altered lrgAB expression in the lrtS mutant, as expression of lrtS-regulated genes was unaltered in an lrgAB mutant relative to the wild-type strain (Table 1). Taken together, these observations suggest that LytST exerts control over its transcriptome in a growth-phase dependent manner, and to our knowledge, this is the first study that has compared the scope of LytST regulation at different phases of growth. Interestingly, RNA microarray studies of lrt mutants have also been performed in *S. aureus* [38], *S. epidermidis* [40], and *B. subtilis* [42]. As we have observed here in *S. mutans*, a global effect of LytST on gene expression was also noted in *S. aureus* and *S. epidermidis* [38,40]. In *S. aureus*, LytST appeared to exert
primarily positive effects on gene expression in exponential phase when aerobic cultures were grown in media containing excess (35 mM) glucose, as only 7 genes were found to be upregulated in the lytS mutant [38]. In *S. epidermidis*, a large number of genes were up- or down-regulated as a function of the presence of LytST during exponential phase during aerobic growth in medium containing 12 mM glucose [40]. In contrast, mutation of lytS only appeared to affect the expression of lytST itself and genes encoding lrgAB and cipAB homologues in *B. subtilis* [42]. However, due to the differences in growth conditions used (glucose levels and/or culture aeration) and the differing metabolic pathways present in these organisms, it is difficult to establish direct correlations between these studies and the *S. mutans* microarray results presented here.

As demonstrated previously [37], expression of lrgAB was also shown to be tightly controlled by the LytST two-component system in *S. mutans* in this study. Specifically, we have found that LytST-dependent expression of lrgAB is regulated in part by glucose metabolism and oxygen in *S. mutans* (Figure 1). Furthermore, control of lrgAB expression by LytST appears to be highly growth-phase dependent: lrgAB expression in the lytS mutant exhibited only a modest decrease in expression in early exponential phase (0.49 relative to UA159, Additional file 1: Table S1), whereas lrgAB expression was down-regulated some 200-fold in the lytS mutant at late exponential phase (Additional file 2: Table S2). Alternatively, it is possible that control of lrgAB expression by LytST is related to higher glucose availability during early exponential phase. Although detailed mechanistic studies have not yet been performed, there is mounting evidence that these proteins are critical for oxidative stress resistance in *S. mutans*: (1) lrgAB expression is highly regulated by oxygen ([11] and this study); (2) a lrgAB mutant was defective in aerobic growth on BHI agar plates [37]; (3) a lrgAB mutant displayed a decreased growth rate in the presence of paraquat (a superoxide-generating agent) relative to the wild-type strain [37]; and (4) a lrgAB mutant displayed a strong growth defect during static planktonic aerobic growth in BHI in the presence and absence of H₂O₂ challenge (this study). Interestingly, a link between LrgAB and oxidative stress was also demonstrated in *S. aureus*, where lytSR and lrgAB expression were upregulated 2-5 fold in response to azurophilic granule proteins, H₂O₂, and hypochlorite [54].

In agreement with a role for LrgAB in oxidative stress resistance, several LytST-regulated genes identified in this study have also been implicated in bacterial oxidative stress responses. Upregulated potential oxidative stress genes include yghU, a putative anti-oxidant enzyme [50], tpx, a predicted thiol peroxidase [55], and recF, a single-stranded DNA exonuclease protein that facilitates DNA repair in response to oxidative stress [51]. Conversely, several genes belonging to the TnSMu2 gene cluster (SMU.1334c – SMU.1359) were downregulated in the lytS mutant. These genes are annotated as encoding a series of gene products involved in bacitracin and gramicidin synthesis [56], but more recently have been shown to be responsible for nonribosomal peptide and polyketide (NRP/PK) biosynthesis of a pigment that enhances aerobic growth and tolerance to H₂O₂ challenge in *S. mutans* UA159 [45]. The altered expression of one or more of these genes may explain, in part, the increased ROS accumulation that was observed in the lytS mutant when challenged with H₂O₂ (Figure 5). Furthermore, it was previously found that a two-component system responsible for positive regulation of the NRP/PK genes was located on the TnSMu2 genomic island of UA140 but not in UA159 [45]. This observation, combined with the microarray results performed here (Additional file 1: Table S1 and Additional file 2: Table S2) suggest that LytST may have taken over some of the regulatory functions of this non-core-genome two-component system that is missing in UA159.

Interestingly, H₂O₂ has also been shown to be a potent stimulator of competence and eDNA release in *S. sanguinis* [57], *S. gordonii* [57,58], and *S. pneumoniae* [59]. Although the effects of H₂O₂ on *S. mutans* competence, cell lysis, and eDNA release have not been directly measured, it has been shown that growth under aerobic conditions promotes competence in *S. mutans* [47], and that expression of competence-related genes is upregulated during aerobic growth [11]. The results presented here have demonstrated that expression of comYB, a gene encoding a component of the DNA-binding uptake system in *S. mutans* [47] was upregulated 2-fold in early exponential phase and 22-fold in late exponential phase in the lytS mutant (Additional file 1: Table S1 and Additional file 2: Table S2). The significance of high-level comYB expression in the lytS mutant at late exponential phase is unclear, given that maximal *S. mutans* competence develops in actively-growing populations [60,61]. Accordingly, upregulation of comYB expression did not correlate with increased transformability of the lytS mutant under the conditions tested in this study (Figure 3). However, it was found that the lrgA mutant displayed a significant reduction in competence. It has been recently reported that only a subpopulation of *S. mutans* culture lyses in response to CSP, and this lysis event is controlled in part by the CipB bacteriocin and the Cipl immunity protein [62]. Subsequent microarray analysis of a cipl (immunity protein) mutant showed that both lytST and lrgAB expression were highly upregulated in the cipl mutant [63]. These results, combined with the fact that LrgA/B has been shown to be involved in regulating cell lysis and eDNA release in *S. aureus* [21,29], lends strong support to the idea that LrgA plays
an important role during competence, possibly by altering membrane permeability or by modulating murine hydrolase activity.

The \textit{S. mutans} \textit{comY} operon consists of nine co-transcribed genes, of which the first eight genes are either essential to or significantly affect competence [46]. The ninth gene of this operon is predicted to encode acetate kinase (AckA), an enzyme that catalyzes the inter-conversion of acetyl-phosphate and acetate [46,64]. For micro-organisms with an inefficient or incomplete TCA cycle such as \textit{S. mutans}, AckA-mediated conversion of acetyl-phosphate to acetate is thought to be a critical mechanism of generating ATP [reviewed in [65]].

Since \textit{ackA} (\textit{comY}) was previously found to be upregulated in \textit{S. mutans} during aerated growth [11], it is possible that LytST is involved in the regulation of energy generation through the phosphate acetyltransferase (Pta)-AckA pathway during aerobic growth and/or during oxidative stress. In this respect, it has recently been reported that an \textit{S. mutans pta} mutant was more susceptible to both acid and oxidative stresses [66].

The ability of \textit{S. mutans} to combat \textit{H2O2} stress is critical for its survival in the oral cavity, yet \textit{H2O2} detoxifying mechanisms and their regulation have not been extensively-characterized in this organism, limited primarily to the ScnRK and VicRK two-component systems [67,68], \textit{ropA} [69], \textit{brpA} [70], \textit{lucS} [71] and genomic island \textit{TnSMu2} [45]. \textit{H2O2} has been shown to have potent antibacterial effects on \textit{S. mutans} [72], and it is thought that \textit{H2O2} produced by other oral streptococcal species serves as an antagonist against \textit{S. mutans}. For example, \textit{S. sanguinis} and \textit{S. gordonii} have been shown to produce \textit{H2O2} via pyruvate oxidase under aerobic growth conditions, and this \textit{H2O2} production allows them to compete effectively against \textit{S. mutans} when co-cultured under aerobic growth conditions [57]. It is therefore possible that the \textit{S. mutans} LytST regulon mediates a pleiotropic protective response against these \textit{H2O2}-producing niche competitors. On-going and future studies by our group will focus on experimental testing of this hypothesis.

\section*{Conclusions}

In summary, the LytST two-component system has been shown to have a pleiotropic effect on gene expression in \textit{S. mutans}. This is congruent with microarray analyses of \textit{lytS} mutants in \textit{S. aureus} [38] and \textit{S. epidermidis} [40]. However, unlike in other organisms, we have been able to identify a pattern of LytS-mediated gene expression that suggests a role for this regulon in responding to oxidative/\textit{H2O2} stress. Although we have not yet been able to identify the external signal to which LytS responds, it is likely linked to an oxidative stress-sensing mechanism, such as \textit{H2O2}-mediated membrane damage (ie. lipid peroxidation) via its large number of transmembrane domains, or oxygen/ROS interactions with its predicted cytoplasmic GAF domain, a ubiquitous signaling domain that has been shown to detect changes in the redox state of bound iron or oxygen in \textit{Mycobacterium tuberculosis} [73-75]. Establishing mechanistic links between the LytST regulon, \textit{H2O2} resistance, and competence regulation will provide valuable new insights into \textit{S. mutans} survival and virulence in the oral cavity.

\section*{Methods}

\subsection*{Bacterial strains, media, and growth conditions}

For all experiments, frozen glycerol stocks of \textit{S. mutans} UA159 and its isogenic \textit{lytS} (SAB111; \textit{ΔlytS::NPKmr}), \textit{lrgA} (SAB113; \textit{ΔlrgA::NPSpβ}), \textit{lrgB} (SAB119; \textit{ΔlrgB::NPEEm}), and \textit{lrgAB} (SAB115; \textit{ΔlrgAB::ΩKm}) mutants [created previously in [37] were freshly streaked for isolation on either Todd Hewitt Yeast Extract (THYE) or Brain Heart Infusion (BHI), containing selective antibiotic as appropriate: kanamycin (Km) – 1000 μg/ml, erythromycin (Em) – 10 μg/ml, spectinomycin (Sp) - 1000 μg/ml]. With the exception of SAB115 (\textit{lrgAB} mutant), all mutants were created using non-polar (NP) antibiotic-resistance markers [37]. Unless otherwise indicated, all \textit{S. mutans} cultures were grown as static cultures in BHI or THYE broth at 37°C and 5% \textit{CO2}.

\subsection*{Analysis of \textit{lrgAB} expression}

To measure the effects of oxygen and glucose on \textit{lrg} expression, overnight THYE cultures of UA159 and the \textit{lytS} mutant (\textit{n} = 3 biological replicates each, grown at 0 RPM, 37°C and 5% \textit{CO2}) were each inoculated to an OD600 = 0.02 into THYE containing either 11 mM or 45 mM aerobic) was calculated by dividing the gene copy number of each test sample by the average gene copy number of UA159 EP. Data was then normalized by dividing each \textit{lrgA} fold-change expression value by its corresponding 16S fold-change expression value.

\subsection*{RNA microarray analysis of UA159 and \textit{lytS} mutant}

To assess the effect of LytS on global gene expression, overnight BHI cultures of UA159 and \textit{lytS} mutant (\textit{n} = 3...
biological replicates per strain) were diluted to an OD600 = 0.02 in BHI, and grown as static cultures at 37°C and 5% CO2. Total RNA was isolated from each culture at early-exponential (OD600 = 0.15) and late exponential phase (OD600 = 0.9), using previously-published methods [77]. RNA microarray analysis was performed using S. mutans UA159 microarrays provided by The Institute for Genomic Research, and previously-described methods and data analysis [11,70,78]. In brief, 2 μg total bacterial RNA was used in each reverse-transcription and cDNA labeling reaction (performed as described in [70,78]), and a single preparation from each culture was hybridized per microarray slide in a Maui hybridization chamber (BioMicro Systems, Salt Lake City, UT). The resulting microarray slides were scanned, analyzed, and normalized using TIGR Spotfinder software (http://www.tigr.org/software/), and in-slide replicate analysis was performed with the TIGR microarray data analysis system (MIDAS; http://www.tigr.org/software/). Statistical analysis was carried out with BRB array tools (http://linus.nci.nih.gov/BRB-ArrayTools.html/) with a cutoff P value < 0.005 for the early exponential-phase data and P < 0.001 for the late exponential phase data. To validate the microarray results, real-time quantitative RT-PCR was performed on a subset of the differentially-expressed genes, as described previously [77,79]. All real-time PCR primers were designed with Beacon Designer 4.0 software (Premier Biosoft International, Palo Alto, CA), and standard curves for each gene were published elsewhere [80]. The microarray data obtained from these studies have been deposited to NCBI's gene expression omnibus (GEO) [81] (GEO Accession #GSE39470) and comply with MIAME guidelines [82].

Quantitative competence assays
To compare the ability of UA159 and its isogenic lytS, lrgA, lrgB, and lrgAB mutants to take up exogenously-added plasmid DNA, a quantitative competence assay was performed on n = 4-6 biological replicates of each strain using a previously-published protocol [83] with the following modifications: Overnight cultures of each strain were diluted to an OD600 = 0.02 in BHI, and grown in a 96-well plate to an OD600 = 0.15 prior to addition of 500 ng plasmid DNA with and without 100 ng CSP. Plasmid pAT28 (encoding spectinomycin resistance; [84]) was used to assess transformation efficiency in UA159, lytS, lrgB, and lrgAB mutants. Because the lrgA mutant was generated with a spectinomycin-resistance cassette [37], plasmid pORi23 [encoding erythromycin resistance; [85]] was used to assess transformation efficiency in UA159 and lrgA mutant. After 2.5 h incubation in the presence of plasmid DNA +/- CSP, cultures were serially diluted and plated on BHI agar with and without selective antibiotic. CFU/ml of each culture were enumerated after 48 h growth at 37°C and 5% CO2, and transformation efficiencies were calculated as the percentage of transformants (CFU/ml on BHI + selective antibiotic) among total viable cells (CFU/ml on BHI).

H2O2 assays
To assess the ability of UA159, lytS and lrgAB mutants to grow in the presence of H2O2, overnight cultures of each strain (n = 6 biological replicates) were each diluted 40-fold into BHI. 1 ml aliquots of each diluted culture were either untreated or challenged with 1 mM H2O2. Aliquots of each (500 μl per well, 2 wells total) were then immediately transferred to an optically-clear 48-well tissue culture plate (Costar 3548), which was incubated for 20 h at 37°C (aerobic atmosphere) in a Biotek Synergy microplate reader. OD600 measurements of each well were recorded at 2 h intervals.

Oxidative stress measurements
To assess intracellular oxidative stress in UA159 and lytS mutant, single isolated colonies of each strain (n = 3-6 biological replicates per strain) were inoculated into culture tubes containing 4 ml BHI, and grown in “low-O2” conditions (37°C, 0 RPM, 5% CO2). After 20 h growth, 2 × 1 ml aliquots of each culture were harvested by centrifugation in a microcentrifuge (3 min at 13,000 RPM). The culture supernatants were discarded, and cell pellets were each resuspended in 1 ml Hanks Buffer (HBSS) containing 5 μM chloromethyl 2,7'-dichlorofluorescein diacetate (CM-H2DCFDA; Invitrogen Molecular Probes), a cell-permeable fluorescent compound that is oxidized in the presence of H2O2 and other reactive oxygen species (ROS) and is considered a general indicator of cellular oxidative stress [52,53]. Cell suspensions were incubated at 37°C for 60 min to “load” the cells with CM-H2DCFDA, followed by centrifugation (3 min at 13,000 RPM). Supernatants were discarded, and cell pellets were washed once with HBSS prior to resuspension in 1 ml HBSS or in 1 ml HBSS containing 5 mM H2O2. Each cell suspension was transferred into triplicate wells (200 μl per well) of an optically-clear 96 well plate (Costar 3614), and the plate was transferred to a Biotek Synergy microplate reader. Fluorescence in relative fluorescence units (RFU; using 492-495 nm excitation and 517-527 nm emission) and OD600 readings of each well were recorded after 30 min incubation at 37°C.

Statistical analysis
All statistical analyses, unless otherwise indicated, were performed using SigmaPlot for Windows 11.0 software (Build 11.0.0.75, Systat Software, Inc.).
Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
SJA carried out the RNA microarray experiments and associated data analysis, performed all real-time PCR studies, participated in the conception and design of the study, and helped draft the manuscript. MDQ carried out all of the RNA isolations for comparing the effects of glucose and oxygenation on *Streptococcus mutans* gtf adherence of *Streptococcus mutans* genes in caries induction in the specific-pathogen-free rat model. Infect Immun 1993, 61(9):3811–3817.

References
1. Deonarine B, Lazar J, Gill MV, Cunha BA: Quadri-valvular endocarditis caused by *Streptococcus mutans*. Clin Microbiol Infect 1997, 3(1):139–141.

2. Biswas S, Bowler IC, Bunch C, Prendergast B, Webster DP: *Streptococcus mutans* gtf infected endocarditis complicated by vertebral discitis following dental treatment without antibiotic prophylaxis. J Med Microbiol 2010, 59(Pt 10):1253–1259.

3. Ullman RF, Miller SJ, Stampfer MJ, Cunha BA: *Streptococcus mutans* endocarditis: report of three cases and review of the literature. Heart Lung 1988, 17(2):209–212.

4. Vose JM, Smith PW, Henny M, Colan D: Recurrent *Streptococcus mutans* endocarditis. Am J Med 1987, 82(3) Spec No:363–362.

5. Yamashita Y, Ueno H, Kato K, Kusumi S, Kato Y, Sekiguchi T: Molecular characterization of a *Streptococcus mutans* mutant altered in environmental stress responses. J Bacteriol 1993, 175(19):6220–6228.

6. Yamashita Y, Takehara T, Kuramitsu HK: Role of the *Streptococcus mutans* gtf genes in caries induction in the specific-pathogen-free rat model. Infect Immun 1993, 61(9):3811–3817.

7. Yamashita Y, Takehara T, Kuramitsu HK: Role of the *Streptococcus mutans* gtf genes in caries induction in the specific-pathogen-free rat model. Infect Immun 1993, 61(9):3811–3817.

8. Munro CL, Michalek SM, Macrina FL: Sucrose-derived exopolymers have site-dependent roles in *Streptococcus mutans*-promoted dental decay. FEMS Microbiol Lett 1995, 128(3):327–332.

9. Ooshima T, Matsumura M, Hashimoto T, Kawai H, Yamao S, Fujiiya T: Contributions of three glycosyltransferases to sucrose-dependent adherence of *Streptococcus mutans*. J Dent Res 2001, 80(7):1672–1677.

10. Munro CL, Michalek SM, Macrina FL: Sucrose-derived exopolymers have site-dependent roles in *Streptococcus mutans*-promoted dental decay. FEMS Microbiol Lett 1995, 128(3):327–332.

11. Ahrn SJ, Cunha BA: Changes in biochemical and phenotypic properties of *Streptococcus mutans* during growth with aeration. Appl Environ Microbiol 2003, 69(8):2517–2527.

12. Ahrn SJ, Cunha BA: Effects of oxygen on biofilm formation and the AtIA autolysis of *Streptococcus mutans*. J Bacteriol 2007, 189(17):6293–6302.

13. Ahrn SJ, Cunha BA: Effects of oxygen on virulence traits of *Streptococcus mutans*. J Bacteriol 2007, 189(25):8519–8527.
83. Seaton K, Ahn SJ, Sagstetter AM, Burne RA: A transcriptional regulator and ABC transporters link stress tolerance, (p)ppGpp, and genetic competence in Streptococcus mutans. J Bacteriol 2011, 193(4):862–874.

84. Trieu-Cuot P, Carlier C, Poyant-Salmeron C, Courvalin P: A pair of mobilizable shuttle vectors conferring resistance to spectinomycin for molecular cloning in Escherichia coli and in gram-positive bacteria. Nucleic Acids Res 1990, 18(14):4296.

85. Que YA, Haefliger JA, Francioli P, Moreillon P: Expression of Staphylococcus aureus clumping factor A in Lactococcus lactis subsp. cremoris using a new shuttle vector. Infect Immun 2000, 68(6):3516–3522.

doi:10.1186/1471-2180-12-187

Cite this article as: Ahn et al: Identification of the Streptococcus mutans LyST two-component regulon reveals its contribution to oxidative stress tolerance. BMC Microbiology 2012 12:187.