Genome-Wide Investigation of *Pasteurella multocida* Identifies the Stringent Response as a Negative Regulator of Hyaluronic Acid Capsule Production

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**ABSTRACT** *Pasteurella multocida* is a Gram-negative capsulated bacterium responsible for a range of diseases that cause severe morbidity and mortality in livestock animals. The hyaluronic acid (HA) capsule produced by *P. multocida* serogroup A strains is a critical virulence factor. In this study, we utilized transposon-directed insertion site sequencing (TraDIS) to identify genes essential for *in vitro* growth of *P. multocida* and combined TraDIS with discontinuous density gradients (TraDISort) to identify genes required for HA capsule production and regulation in this pathogen. Analysis of mutants with a high cell density phenotype, indicative of the loss of extracellular capsule, led to the identification of 69 genes important for capsule production. These genes included all previously characterized genes in the capsule biosynthesis locus and *fs* and *hfq*, which encode known positive regulators of *P. multocida* capsule. Many of the other capsule-associated genes identified in this study were involved in regulation or activation of the stringent response, including *spoT* and *relA*, which encode proteins that regulate the concentration of guanosine alarmones. Disruption of the autoregulatory domains in the C-terminal half of SpoT using insertional mutagenesis resulted in reduced expression of capsule biosynthesis genes and an acapsular phenotype. Overall, these findings have greatly increased the understanding of hyaluronic acid capsule production and regulation in *P. multocida*.

**IMPORTANCE** The bacterial pathogen *P. multocida* can cause serious disease in production animals, including fowl cholera in poultry, hemorrhagic septicemia in cattle and buffalo, atrophic rhinitis in pigs, and respiratory diseases in a range of livestock. *P. multocida* produces a capsule that is essential for systemic disease, but the complete mechanisms underlying synthesis and regulation of capsule production are not fully elucidated. A whole-genome analysis using TraDIS was undertaken to identify genes essential for growth in rich media and to obtain a comprehensive characterization of capsule production. Many of the capsule-associated genes identified in this study were involved in the stringent response to stress, a novel finding for this important animal pathogen.

**KEYWORDS** *Pasteurella multocida*, capsule, hyaluronic acid, stringent response, *P. multocida*
P. multocida is classified into five capsular serogroups, A, B, D, E, and F, with strains from each serogroup producing structurally distinct glycosaminoglycan capsule polymers (4, 5). The capsule biosynthesis locus representing each P. multocida capsule serogroup has been identified (6–8). Genes within each capsule locus encode proteins required for monomer synthesis and activation, glycosyltransferases required to synthesize the specific capsule polymer, and proteins involved in the ABC-transport system and in anchoring of the capsule to the outer membrane (4, 9–12). Directed mutation of capsule export genes in P. multocida strain VP161 (serogroup A) and strain M1404 (serogroup B) generated acapsular mutants that were unable to cause systemic disease in chickens and/or mice (13, 14), proving that capsule is an essential P. multocida virulence factor. In the serogroup A strain VP161, genes in the hyaluronic acid (HA) capsule biosynthesis locus are positively regulated by the nucleoid-associated regulatory protein Fis and the RNA chaperone Hfq (15, 16).

The importance of capsule for P. multocida virulence makes it an ideal target for novel therapeutics and vaccines. However, a complete understanding of the cellular signals that control capsule production in P. multocida and the regulatory mechanisms involved is lacking. In this study, we first utilized transposon-directed insertion site sequencing (TraDIS) to identify genes essential for P. multocida strain VP161 growth in rich media. We then phenotypically screened mutants using TraDISort to identify genes important for HA capsule production in strain VP161. The TraDISort method using density gradient centrifugation has been used previously to separate transposon mutants with altered cell density, which for capsulated bacteria is largely dependent on the amount of extracellular capsule present (17). Mutants with a higher or lower cell density relative to that of the majority of the mutant population are then analyzed by TraDIS to identify the inactivated gene in each mutant (17). In this study, we used TraDISort to isolate P. multocida VP161 transposon mutants with a higher cell density, indicative of reduced extracellular capsule, and identified a total of 69 genes important for capsule production in strain VP161. Individual mutants representing several genes identified by TraDISort analysis were confirmed to have reduced HA capsule production compared to that of the parent strain. Several of the genes associated with the acapsular phenotype were involved in the regulation or activation of the stringent response, which was previously not known to affect capsule production in this bacterium.

RESULTS

Generation of a saturated P. multocida strain VP161-Tn7 Himar1 mutant library. A kanamycin-resistant P. multocida strain, VP161-Tn7, was constructed (Table S1) to allow for use as a recipient in conjugation experiments with donor Escherichia coli strain AL2972 (Table S1), harboring the Himar1 delivery vector pAL6114 (18). Each conjugation mating generated more than $1 \times 10^6$ P. multocida CFU when recovered on heart infusion (HI) media supplemented with kanamycin and spectinomycin. Approximately $6.45 \times 10^6$ P. multocida Himar1 transconjugants were pooled and stored in aliquots at −80°C for subsequent TraDIS experiments.

To identify genes that were essential for in vitro growth in HI and determine the quality and depth of the VP161-Tn7 Himar1 library, two aliquots (A and B) of the mutant library were grown separately in HI medium (37°C, 200 rpm) to mid-exponential growth phase (optical density at 600 nm [OD$_{600}$] of ~0.5). DNA was isolated from cells harvested from each culture and used to generate TraDIS libraries; two different concentrations of adapter ligated material were used to ensure there was sufficient template for library amplification. Four TraDIS libraries were successfully amplified and sequenced on an Illumina MiSeq, namely, A1 and B1 (100 ng template) and A2 and B2 (250 ng template). The data were analyzed using the Bio-TraDIS pipeline to identify the total number of transposon-specific reads and the location of Himar1 insertion sites in the VP161 genome (Fig. S1, Table S2). Comparison of unique Himar1 insertion sites per gene gave high correlation coefficients between the rich media TraDIS library replicates regardless of the amount of DNA template used (Fig. S2), allowing the data to be combined and thus increasing the number of unique Himar1 insertion sites across the genome and, therefore, the resolution. In total, 81,292 unique Himar1 insertion sites were identified (Table S2), representing a unique insertion every 28 bp of the VP161 genome and an average of 38 unique insertions per gene. An insertion index was calculated for each gene by dividing
the number of unique insertions by gene length, and insertion indexes for all genes were then used to generate two normally distributed data sets representing essential and nonessential genes (Fig. S3). Essential genes were determined to be those with an insertion index of less than 0.0134, defined as the intersection between the two curves. A total of 509 genes were identified as essential for *P. multocida* growth in HI broth (Table S3), encoding 473 predicted proteins, 35 tRNAs, and one 55 rRNA. Assignment of the encoded proteins to cluster of orthologous groups (COGs) revealed that a large proportion were predicted to be involved in translation, ribosomal structure and biogenesis. The 473 protein-encoding genes were compared to the database of essential genes (DEG) that contains genes identified as essential in other bacterial species (19). There were homologs in the DEG for 442 (93%) of the protein-encoding genes, indicating that TraDIS analysis of *P. multocida* grown in vitro in HI broth was a highly effective method to identify essential genes in this species (Table S3). Of the remaining 31 *P. multocida* essential genes encoding proteins, 17 encoded hypothetical proteins and 14 encoded proteins predicted to be involved in a range of processes, including the DNA repair and stress response, gene and protein regulation, nutrient synthesis, and contact-dependent growth inhibition (Table S3).

Isolation of acapsular *P. multocida* cells via Percoll discontinuous density gradient centrifugation. To separate mutants within the library based on cell density, TraDISort using consecutive Percoll discontinuous density gradients (10%, 40%, and 80% Percoll) was employed (Fig. S1 and Fig. S4A). Following the first centrifugation, the saturated *P. multocida* VP161-Tn7 Himar1 mutant library separated into an upper, low-density cell layer containing capsulated cells and a lower, high-density cell layer (Fig. S4A). Mutants within the upper cell layer following this first centrifugation were not examined further, as they were predicted to produce wild-type levels of capsule. The lower layer of cells containing high-density cells was recovered, as this was predicted to contain cells with reduced extracellular capsule. A sample of this population was plated onto rich media. Following overnight growth, approximately 50% of the colonies generated were mucoid, indicating that a single gradient separation was insufficient to fully separate the acapsular cells from the cells capable of producing capsule. It is known that *P. multocida* cells in the exponential growth phase produce significantly more extracellular capsule than cells in other phases of growth (15). To allow transient acapsular mutants the opportunity to produce extracellular capsule, the lower cell layer population was grown for 4 h in HI broth. The cells were recovered and then subjected to a second Percoll discontinuous density gradient. Following centrifugation, the cells separated into a top, low-density (termed LD2) cell layer and a bottom, high-density (termed HD2) cell layer. Each population was recovered and plated separately onto HI agar plates. All colonies generated from cells recovered from the LD2 cell layer displayed a capsulated phenotype (large, mucoid), and approximately 99% of the colonies generated from cells recovered from the HD2 layer displayed an acapsular phenotype (small, nonmucoid), confirming that consecutive density gradient centrifugations were required to successfully separate high-density mutants with an acapsular phenotype.

To confirm that mutants producing small and nonmucoid colonies represented acapsular mutants, genomic DNA was isolated from 35 nonmucoid colonies recovered from the HD2 cell layer and used separately as the template in a Sanger sequencing reaction with an outward-firing, Himar1-specific primer (BAP3453, Table S4) to identify the transposon insertion site in the genome of each mutant. The majority of the 35 Himar1 mutants examined had insertions within genes known to be required for capsule biosynthesis. Interestingly, several genes not previously known to be involved in capsule production were also identified, including pgm, ppx, and cpxA. Absorbance assays measuring the amount of extracellular HA capsular material produced by Himar1 mutants representing eight genes showed that all produced significantly reduced amounts of HA compared to the parent strain, VP161-Tn7 (Fig. S4B). Together, these data show that consecutive discontinuous Percoll density gradients were successful in isolating acapsular mutants from the large VP161-Tn7 Himar1 mutant library.

Identification of capsule-associated genes in VP161. Genomic DNA was extracted from the population of VP161-Tn7 Himar1 mutants recovered from the top, LD2 cell layer (predicted to be capsulated) and separately from the population of mutants in the bottom, HD2 cell layer (predicted to be acapsular). Separate TraDIS libraries were generated using
genomic DNA from both cell layers, sequenced on an Illumina MiSeq, and analyzed by Bio-TraDIS and modified scripts (Fig. S1). The number of unique Himar1 insertions was highly correlated between the LD2 and HD2 library replicates (Fig. S5A), and principal-component analysis of unique Himar1 insertions per gene in the TraDIS libraries generated using the HD2 cell layer compared to TraDIS libraries generated using the LD2 cell layer following consecutive discontinuous Percoll gradient centrifugations, with the \(-\log_{10} q\) value provided for each comparison. Genes in the HD2 cell layer with a log2 fold change of \(>2\) and a \(q\) value of \(<0.001\) increase in Himar1 insertions compared to those in the LD2 cell layer were identified as important for capsule production (both cutoffs shown as gray dotted lines). Known capsule biosynthesis genes are shown in blue, and known positive regulators of HA capsule production are shown in red. (C) TraDIS library analysis showing number of Himar1 insertions mapping to the VP161 capsule locus in the input Himar1 library (black) and after consecutive discontinuous Percoll gradients; top, LD2 cell layer (capsulated, blue), and bottom, HD2 cell layer (acapsular, green). The majority of mutants with Himar1 insertions in the capsule biosynthesis locus separated into the HD2 cell layer of the Percoll gradient, as shown by data showing a large increase in Himar1 insertions in these genes in mutants recovered from the bottom, HD2 cell layer compared to the insertions in the same genes from mutants recovered from the top, LD2 cell layer.

FIG 1 Overview of TraDISort library data. (A) The number of Himar1 insertions per 500 bp across the VP161 genome in the TraDIS libraries generated using either the top, LD2 cell layer (predominantly capsulated cells, insertions sites in blue) or the bottom, HD2 cell layer (predominantly acapsular, insertion sites in green). Cell layers LD2 and HD2 were recovered from the second discontinuous Percoll gradient centrifugation. Genes identified as important for \(P.\ multocida\) hyaluronic acid (HA) capsule production are indicated using black bars. (B) Volcano plot showing the log2 fold change in the number of Himar1 insertions per gene in the TraDIS libraries generated using the HD2 cell layer compared to TraDIS libraries generated using the LD2 cell layer following consecutive discontinuous Percoll gradient centrifugations, with the \(-\log_{10} q\) value provided for each comparison. Genes in the HD2 cell layer with a log2 fold change of \(>2\) and a \(q\) value of \(<0.001\) increase in Himar1 insertions compared to those in the LD2 cell layer were identified as important for capsule production (both cutoffs shown as gray dotted lines). Known capsule biosynthesis genes are shown in blue, and known positive regulators of HA capsule production are shown in red. (C) TraDIS library analysis showing number of Himar1 insertions mapping to the VP161 capsule locus in the input Himar1 library (black) and after consecutive discontinuous Percoll gradients; top, LD2 cell layer (capsulated, blue), and bottom, HD2 cell layer (acapsular, green). The majority of mutants with Himar1 insertions in the capsule biosynthesis locus separated into the HD2 cell layer of the Percoll gradient, as shown by data showing a large increase in Himar1 insertions in these genes in mutants recovered from the bottom, HD2 cell layer compared to the insertions in the same genes from mutants recovered from the top, LD2 cell layer.
index of &gt;0.8) was identified as important for HA capsule production in strain VP161 (Fig. 1B). This analysis identified 69 genes important for HA capsule production, including all genes in the capsule biosynthesis locus (Table 1, Fig. 1C), fis and hfq, which encode known positive regulators of capsule biosynthesis genes, and the novel capsule-associated genes identified by Sanger sequencing of individual mutants, namely, pgm, ppx, and cpxA. These data confirm that the use of two consecutive discontinuous density gradient centrifugations had successfully isolated acapsular mutants from the saturated VP161-Tn7 Himar1 library. Other genes that had a significant increase in the number of transposon insertions in this population of mutants were involved in a wide range of functions, including the stringent response, central metabolism and carbohydrate biosynthesis, and translation, transport, and ATP synthesis (Table 1). Interestingly, except for fis, no other genes encoding transcription factors were identified, suggesting that there are no other transcription factors that act as direct positive regulators of HA capsule biosynthesis in strain VP161.

Validation of novel capsule production genes. To confirm that the genes newly identified using TraDiSort were important for HA capsule production, TargeTron-directed mutagenesis of a number of genes was attempted. In total, three TargeTron mutants were successfully constructed; these mutants had insertions in ppx and ptsH and in the 3′ regulatory region of spoT (Table S1). The 3′ region of spoT was chosen for TargeTron insertion, as TraDiSort analysis indicated that the majority of HD2 spoT mutants had insertions in this region (Fig. 2). To add to these validation experiments, pgm and galU Himar1 mutants isolated from the HD2 cell layer were also included. Each mutant was confirmed to have a single transposon insertion by Sanger sequencing as described above.

To prove that the inactivated gene in each mutant was responsible for the acapsular phenotype, complementation experiments were conducted. The appropriate complementation plasmid and corresponding empty vector were used separately to transform each mutant to generate complemented mutant and empty vector control strains (Table S1). The amount of HA produced by each strain the corresponding complemented mutant was determined by HA absorbance assays using cells grown to mid-exponential growth phase. The amount of HA produced by each strain was compared to HA production in the corresponding parent strain harboring empty vector (VP161 for TargeTron mutants and VP171-Tn7 for Himar1 mutants). A VP161 hyaD TargeTron mutant (Table S1) was included as a known acapsular control strain. As expected, all mutants harboring empty vector produced significantly less HA than the parent strains VP161 and VP161-Tn7 (Fig. 3). All mutants provided with an intact copy of the appropriate gene in trans produced significantly more HA than the corresponding mutant harboring empty vector, although the amount of HA was not restored to the same level as the parent strain for all mutants (Fig. 3). Scanning electron microscopy (SEM) of the spoT TargeTron mutant was performed to show that loss of HA capsular material correlated with less capsule on the surface of the bacterium. For this experiment, the parent strain, VP161, and the VP161 hyaD TargeTron mutant were used as known capsular and acapsular controls, respectively. Imaging of cells in the mid-exponential phase of growth revealed that the wild-type VP161 cells had a ruffled outer surface indicative of a capsule layer that was absent in both the hyaD and spoT mutants harboring empty vector but was present in the complemented mutant strains (Fig. 4). Corresponding to less capsule, both the hyaD and spoT mutants had significantly reduced cell widths compared to wild-type VP161, and mutant cells provided with the appropriate complementation plasmid had a width similar to that of wild-type VP161 cells (Fig. 4). Together, these data show that novel capsule-associated genes identified by TraDiSort were required for HA capsule production in VP161.

Disruption of the 3′ regulatory region of spoT results in reduced expression of the capsule biosynthesis genes independent of fis. The proteins encoded by spoT and relA are homologs of proteins within the RelA/SpoT homolog (RSH) superfamily. In our study, TraDiSort spoT and relA mutants enriched in the HD2 cell population had insertions predominantly in the 3′ region of the gene (Fig. 2). In E. coli, RSH proteins play central roles in the stringent response to various stresses by controlling the levels of the guanosine alarmone molecules 5′-diphosphate-3′-diphosphate and GTP-3′-diphosphate [collectively named (p)(pp)Gpp]. RelA is a (p)(pp)Gpp synthase that in other bacteria has been shown to respond to
### TABLE 1 Genes identified using TraDiSort as important for hyaluronic acid capsule production in *P. multocida* strain VP161

| Gene name | VP161 locus tag | Predicted function | Log. fold change | q value | Insertion index ratio |
|-----------|-----------------|---------------------|-----------------|---------|----------------------|
| znuA      | PmVP161_0255    | High-affinity zinc uptake system protein ZnuA | 7.2            | 1.02E-43 | 3.1                  |
| gmk       | PmVP161_0260    | Guanylate kinase    | 7.1            | 0.000618 | 3.0                  |
| rpoZ      | PmVP161_0261    | DNA-directed RNA polymerase subunit omega | 6.8            | 3.22E-60 | 1.5                  |
| spoT      | PmVP161_0262    | Bifunctional (pppGpp synthase/hydrolase SpoT) | 6.0            | 1.99E-35 | 1.7                  |
| seqA      | PmVP161_0336    | Negative modulator of initiation of replication | 5.6            | 2.22E-49 | 0.9                  |
| glnD      | PmVP161_0442    | Bifunctional uridylyltransferase/uridylyl-removing enzyme | 7.6            | 7.88E-115 | 0.9                  |
| pmcC      | PmVP161_0549    | Release factor glutamine methyltransferase | 12.5           | 21.3E-52 | 26.0                 |
| truX      | PmVP161_0625    | tRNA pseudouridine synthase A | 4.1            | 3.03E-34 | 0.8                  |
| ubIX      | PmVP161_0693    | Flavin prenyltransferase UbIX | 3.9            | 3.79E-25 | 0.9                  |
| pta       | PmVP161_0698    | Phosphate acetyltransferase | 6.3            | 4.42E-06 | 1.6                  |
| dnaJ      | PmVP161_0735    | Chaperone protein DnaJ | 9.1            | 1.50E-52 | 3.5                  |
| rmpP      | PmVP161_0757    | Ribosome maturation factor RmpP | 9.9            | 2.84E-83 | 2.7                  |
| phyB      | PmVP161_0773    | Capsule phospholipid substitution protein | 12.7           | 5.07E-159 | 6.1                  |
| phyA      | PmVP161_0774    | Capsule phospholipid substitution protein | 12.5           | 1.16E-152 | 6.2                  |
| hyeE      | PmVP161_0775    | Hyaluronic acid capsule biosynthesis protein | 13.7           | 1.42E-141 | 6.0                  |
| hydA      | PmVP161_0776    | Hyaluronic acid synthase | 14.0           | 1.03E-131 | 8.4                  |
| hycA      | PmVP161_0777    | UDP-glucose 6-dehydrogenase | 13.5           | 8.45E-125 | 12.2                 |
| hyB       | PmVP161_0778    | Hyaluronic acid capsule biosynthesis protein | 12.7           | 5.64E-184 | 5.8                  |
| hexD      | PmVP161_0779    | Hyaluronic acid capsule export | 8.1            | 1.97E-22 | 3.7                  |
| hexC      | PmVP161_0780    | Hyaluronic acid capsule export | 7.8            | 2.08E-28 | 3.2                  |
| hexB      | PmVP161_0781    | Hyaluronic acid capsule transport protein HexB | 6.5           | 1.34E-10 | 2.2                  |
| hexA      | PmVP161_0782    | Hyaluronic acid capsule transport ATP-binding protein HexA | 5.3           | 7.72E-08 | 2.0                  |
| pmVP161_0825 | Hypothetical protein | 6.3            | 4.00E-76 | 0.9                  |
| holC      | PmVP161_0828    | DNA polymerase III subunit chi | 2.9            | 7.33E-09 | 1.5                  |
| gyrA      | PmVP161_0847    | DNA gyrase subunit A | 9.6            | 1.19E-23 | 3.0                  |
| pmVP161_0878 | Hypothetical protein | 11.9           | 1.36E-107 | 18.4                 |
| ptsH      | PmVP161_0905    | Phosphocarrier protein HPt | 9.8            | 6.26E-102 | 2.9                  |
| hqG       | PmVP161_0914    | RNA-binding protein HqG | 6.8            | 1.61E-63 | 2.5                  |
| znuB      | PmVP161_0986    | High-affinity zinc uptake system membrane protein ZnuB | 6.0           | 5.09E-09 | 1.1                  |
| secG      | PmVP161_1023    | Protein export membrane protein SecG | 10.8           | 7.65E-36 | 7.0                  |
| eepA      | PmVP161_1029    | Elongation factor P-(P)-beta-lysin ligase | 5.7           | 2.78E-61 | 0.9                  |
| quiD      | PmVP161_1046    | 6-Carboxy-5,6,7,8-tetrahydropyrimidin synthase | 7.1           | 1.63E-73 | 1.0                  |
| rnb       | PmVP161_1050    | Exoribonuclease 2 | 5.5            | 2.24E-44 | 1.2                  |
| efp       | PmVP161_1135    | Elongation factor P | 7.9           | 1.95E-16 | 7.0                  |
| eepB      | PmVP161_1136    | L-Lysine 2,3-aminomutase | 7.6           | 2.12E-32 | 1.7                  |
| a1sT_2    | PmVP161_1144    | Amino acid carrier protein last | 4.6           | 4.65E-31 | 1.1                  |
| reP       | PmVP161_1260    | Regulator of sigma-E protease ReP | 5.7           | 0.00018 | 1.6                  |
| cpxA      | PmVP161_1346    | Sensor histidine kinase CpxA | 2.5           | 1.94E-14 | 0.9                  |
| relA      | PmVP161_1370    | GTP pyrophosphokinase | 6.3           | 6.54E-74 | 1.1                  |
| purB      | PmVP161_1383    | Adenylosuccinate lyase | 4.0           | 4.34E-48 | 0.9                  |
| ppx       | PmVP161_1426    | Exopolysphosphatase | 8.6           | 9.75E-81 | 2.2                  |
| trmA      | PmVP161_1430    | tRNA/tmRNA [uracil-C(S)]-methyltransferase | 7.6           | 8.83E-92 | 1.2                  |
| pmVP161_1431 | Hypothetical protein | 9.2           | 1.14E-60 | 2.5                  |
| tufA_1     | PmVP161_1480    | Elongation factor Tu 1 | 6.0          | 1.33E-61 | 1.5                  |
| mioC      | PmVP161_1625    | Protein MioC | 3.0           | 0.000615 | 0.9                  |
| pmVP161_1628 | Hypothetical protein | 7.1           | 7.11E-94 | 2.2                  |
| atpB_1     | PmVP161_1629    | ATP synthase subunit a | 4.8           | 3.22E-56 | 4.5                  |
| atpB_2     | PmVP161_1630    | ATP synthase subunit c | 7.8           | 3.42E-63 | 2.1                  |
| atpF       | PmVP161_1631    | ATP synthase subunit b | 8.1           | 1.32E-34 | 3.5                  |
| atpG_2     | PmVP161_1632    | ATP synthase subunit delta | 4.3          | 1.10E-50 | 2.2                  |
| atpA       | PmVP161_1633    | ATP synthase subunit alpha | 8.0          | 1.42E-80 | 2.5                  |
| atpD       | PmVP161_1634    | ATP synthase gamma chain | 7.0          | 5.52E-68 | 2.9                  |
| pgl        | PmVP161_1693    | 6-Phosphogluconolactonase | 5.6           | 1.24E-30 | 1.0                  |
amino acid starvation. SpoT is a bifunctional (p)ppGpp synthase/hydrolase that in *E. coli* has been shown to respond to iron, phosphate, and fatty acid starvation (20–23). The C-terminal regulatory region of both proteins contains four domains, namely, the ThrRS, GTPase, and SpoT (TGS) domain, the helical domain, the conserved cysteine or zinc finger domain (CC/ZFD), and the aspartokinase, chorismate mutase, and TyrA (ACT) domain (Fig. 2) (24). There were transposon insertions across the entire *relA* gene following *in vitro* growth in HI, including the N-terminal domain encoding the synthase domain, indicating that RelA (p)ppGpp synthase activity is not essential for normal growth *in vitro* in rich media (Fig. 2). However, the TraDISort read data revealed that there was only a very small number of transposon sites

TABLE 1 (Continued)

| Gene name | VP161 locus tag | Predicted function                                    | Log2 fold change | q value | Insertion index ratio |
|-----------|-----------------|-------------------------------------------------------|------------------|---------|-----------------------|
| rpe       | PmVP161_1767    | Ribulose-phosphate 3-epimerase                        | 3.8              | 4.88E-45| 1.2                   |
| thiE      | PmVP161_1798    | Thiamine-phosphate synthase                           | 4.0              | 0.000551| 0.9                   |
| thiM      | PmVP161_1800    | Hydroxyethylthiazole kinase                           | 6.6              | 6.31E-05| 2.2                   |
| ssaB      | PmVP161_1804    | Aliphatic sulfonates import ATP-binding protein SsaB  | 6.6              | 5.96E-26| 2.5                   |
| galU      | PmVP161_1828    | UTP-glucose–1-phosphate uridylyltransferase           | 13.3             | 2.19E-175| 5.4                   |
| PmVP161_1879 | PmVP161_1879   | Hypothetical protein                                  | 7.1              | 7.15E-05| 1.5                   |
| rtaA_2    | PmVP161_1902    | Elongation factor Tu 1                               | 6.0              | 9.15E-63| 1.4                   |
| glmS      | PmVP161_1917    | Glutamine-fructose-6-phosphate aminotransferase (isomerizing) | 5.9 | 6.37E-23| 1.6 |
| rpsA      | PmVP161_1987    | Ribose-5-phosphate isomerase A                        | 8.6              | 2.36E-46| 2.7                   |

**FIG 2** Insertion site plots showing Himar1 insertions in *spoT* and *relA*. Insertion sites represent combined data from the rich media TraDIS libraries (black), LD2 cell layer TraDIS libraries (blue), and HD2 cell layer TraDIS libraries (green). Below the insertion site plots is a graphical representation of domains in RelA and SpoT. TGS, ThrRS GTPase domain; CC, coiled coil/zinc finger domain; ACT, aspartokinase, chorismate mutase domain. The horizontal red line in the RelA Hydrolase domain indicates that this domain is predicted to be non-functional based on similarity to the *E. coli* homolog.
in the relA mutant population recovered from the second gradient centrifugation, and these were predominantly in the 3′ regulatory region. This indicates that the majority of relA mutants present in the input library had a cell density different from that of both the LD2 and HD2 population and were therefore lost from the TraDISort analysis. However, the majority of the relA mutants still present in the HD2 cell layer (log2 fold change increase of 6.3 compared to those in the LD2 cell layer) had Himar1 insertion sites predominantly at nucleotide 1975 or nucleotide 1976, both located upstream of the region encoding the ACT domain.

FIG 3 Amount of hyaluronic acid (HA) capsule produced by P. multocida parent strains VP161 and VP161-Tn7, VP161 TargeTron mutants (hyd, ptsH, ppx, spoT), or Himar1 mutants (pgm, galU). Each mutant was provided with the appropriate vector (pAL99S or pAL99T) or an intact copy of the target gene (provided on the appropriate vector) to demonstrate that the gene inactivated in each strain was responsible for the acapsular phenotype. HA production by each strain was measured in biological quadruplicate using mid-exponential-phase growth cultures. Significant differences in the amount of capsule produced by different strains were determined using Mann-Whitney U test; *, P < 0.05; **, P < 0.01. Error bars represent mean ± standard deviation (SD).

FIG 4 Scanning electron microscopy (SEM) of P. multocida. The parent strain VP161 was provided with pAL99S vector, and the VP161 3′ spoT and hyd TargeTron mutants were provided with the pAL99S vector or with an intact copy of the appropriate gene on pAL99S (complemented mutant). SEM was performed using mid-exponential-phase growth cultures. (A) Electron micrographs of each strain showing the smooth surface of the 3′ spoT and hydD TargeTron mutants (capsulated cells) and the ruffled surface of the parent and complemented mutant strains (capsulated cells). (B) Cell widths of 50 individual cells of each strain as measured using the FUI imaging package in ImageJ. Significant differences in width (lower values indicative of reduced surface capsule) were determined using an unpaired t test. ****, P < 0.0001; error bars, mean ± SD.
The 5’ region of spoT, encoding the (p)ppGpp hydrolase and synthase domains, was deemed essential for the growth of *P. multocida* in vitro in HI media, as no mutants with transposon insertions in this region were recovered following growth in HI media (Fig. 2). However, there were a significant number of mutants with transposon insertions throughout the 3’ regulatory region of spoT (Fig. 2). TraDISort analysis revealed that although some of the 3’ spoT mutants were present in the LD2 cell layer, most were present in the HD2 cell layer (log2 fold change increase of 6.0 in the HD2 cell layer compared to the LD2 cell layer), correlating with the acapsular phenotype displayed by the 3’ spoT TargeTron mutant. The Himar1 transposon insertions in the HD2 mutants were distributed throughout the 3’ regulatory region between nucleotides 1416 to 2124 but particularly in the region encoding the helical domain, the CC/ZFD domain, and the ACT domain (Fig. 2).

Disrupting the 3’ regulatory region of relA in *E. coli*, or the equivalent region in the spoT homolog rel from *Mycobacterium smegmatis* and *Streptococcus equisimilis*, results in increased (p)ppGpp synthesis and/or decreased (p)ppGpp hydrolysis by the modified proteins (25–28). Moreover, increased concentration of ppGpp in *E. coli* results in downregulation of the gene encoding the nucleoid-associated regulatory protein Fis (29, 30). In *P. multocida*, Fis positively regulates the expression of genes within the capsule biosynthesis, and in the absence of Fis, the bacterium is unable to produce HA capsule (16). To determine if the *P. multocida* 3’ spoT TargeTron mutant (AL3391, Table S1) with an acapsular phenotype also had reduced expression of fis, quantitative reverse transcriptase PCR (qRT-PCR) was performed to measure the expression of fis and capsule biosynthesis gene hyaD relative to the expression of the housekeeping gene gyrB. Expression of these genes during mid-exponential-phase growth was measured in wild-type VP161 harboring empty vector, the spoT TargeTron mutant harboring empty vector, and the complemented spoT mutant. The spoT mutant harboring empty vector had significantly reduced hyaD expression compared to VP161, and expression of this gene was partially restored when the mutant was provided with an intact copy of hyaD in trans (Fig. 5). However, there was no significant difference in fis expression between these strains (Fig. 5), indicating that disrupting the C-terminal regulatory region of SpoT resulted in decreased capsule biosynthesis gene expression, but this regulatory change was independent of Fis.

**DISCUSSION**

In this study, we first used TraDIS analysis of a saturated transposon library to identify genes that were essential for in vitro growth of *P. multocida* strain VP161 in rich media. Following the identification of essential genes, we used TraDISort to identify genes that
when inactivated were associated with a high-density, acapsular cell phenotype. Capsule is an essential virulence factor, as it allows *P. multocida* to avoid complement-mediated killing and phagocytosis (13, 14).

TraDIS analysis identified 509 genes as essential for growth of strain VP161 in rich media, which was similar to the number of essential genes identified using transposon insertion sequencing methods in other bacterial species (19). Moreover, approximately 93% of the 473 *P. multocida* proteins identified as essential for growth in rich media were homologs of essential proteins in another bacterial species, the majority of which were also identified using cells grown in vitro in rich media. Thus, our TraDIS analysis successfully identified genes truly essential for *P. multocida* growth in rich media. Most of the essential *P. multocida* genes encoded proteins involved in housekeeping processes, with a large proportion involved in translation and cell wall biosynthesis. Essential genes included those required for glycolysis, pyruvate oxidation, and gluconeogenesis, for the biosynthesis of membranes and cell wall components (phosphatidylethanolamine, lipid A, peptidoglycan), for the synthesis of nucleotides, lysine, vitamins, and cofactors, and for housekeeping processes such as ribosome production, tRNA synthetases, DNA replication, cell division, and Sec protein export. Genes in several of these pathways have also been identified in other bacterial species as essential for growth in rich media (31–34).

Following essential gene analysis, TraDiSort was used to identify genes important for HA capsule production. Consecutive Percoll discontinuous gradient centrifugations were employed to isolate mutants with a high-density, acapsular phenotype. Genes identified as inactivated in these mutants included all genes within the capsule biosynthesis locus as well as genes encoding the global regulator Fis and the Hfq RNA chaperone that each play a role in the positive regulation of capsule in *P. multocida* serogroup A strains (15, 16). Five genes identified as important for capsule production were investigated further. Disrupting *ppx*, *ptsH*, *galU*, *pgm*, and the 3′ autoregulatory region of *spoT* resulted in reduced HA capsule production that was restored when each mutant was provided with the appropriate gene in trans. Moreover, SEM of the VP161 3′ *spoT* mutant and the complemented mutant demonstrated a clear correlation between inactivation of the regulatory region of *SpoT* and an acapsular phenotype.

Several genes that were identified as important for HA capsule production encoded proteins involved in carbohydrate metabolism and the production of HA capsule monomers UDP-α-D-glucosamine (UDP-GlcNAc) and UDP-glucuronic acid (UDP-GlcA). The genes *pgm* and *galU* encode homologs of phosphoglucomutase Pgm and glucose-1-phosphate uridylyltransferase GalU, which together with the UDP-glucose dehydrogenase HyaC synthesize UDP-GlcA from glucose (9, 35, 36). Both *pgm* and *galU* are required for UDP-GlcA synthesis and capsule production in *Streptococcus pneumoniae* (serogroups 1 and 3) and in *Klebsiella pneumoniae* strain NTUH-K2044 (17, 37, 38). In *P. multocida*, we predict that inactivation of *pgm* or *galU* would stop UDP-GlcA synthesis and hence stall HA capsule production. Enriched in the HD2 cell layer were mutants with insertions in *glmS*, encoding β-D-fructose-6-phosphate aminotransferase, which converts β-D-fructose 6-phosphate to α-D-glucosamine 6-phosphate. This substrate is required for the biosynthesis of one of the HA capsule monomers UDP-GlcNac, which is also essential for peptidoglycan and lipopolysaccharide biosynthesis. Also identified was the gene *pgl*, which encodes a protein with a NagB domain. In *E. coli*, NagB is a 6-phosphogluconolactonase/glucosamine-6-phosphate deaminase that converts α-D-glucosamine 6-phosphate to β-D-fructose. *GlmS* acts at the same position in the pathway but in the reverse direction (39). Experiments in *E. coli* strain K-12 showed that *glmS* mutants were viable so long as glucosamine-6-phosphate or N-acetylglucosamine-6-phosphate (converted to glucosamine by *nagA*) was available in the growth media (40). However, more recent TraDIS experiments identified *glmS* as an essential gene when strain K-12 was grown in vitro in Luria broth (41). Multiple studies have shown that the gene *glmM*, encoding a phosphoglucomutase mutase, and *glmU*, encoding a fused N-acetylglucosamine-1-phosphate uridylyltransferase, which are required for the last two steps in the UDP-GlcNac biosynthesis pathway, are essential for the viability of *E. coli* (41–43). Initial TraDIS analysis of the VP161-Tn7 Himar1 library grown in vitro in rich media
revealed that there were no VP161-Tn7 Himar1 mutants with insertions in glmM or glmU (Table S3), indicating that these genes are also essential in *P. multocida*.

The genes encoding RelA and SpoT, proteins involved in regulation of the stringent response, were both identified as important for capsule production. The stringent response is a bacterial stress response induced by nutrient starvation and is activated by increased concentration of the nucleotide alarmones (p)ppGpp (44, 45). The concentration of (p)ppGpp is controlled by RSH proteins together with small alarmone synthase (SAS) and hydrolase (SAH) proteins (24, 44). When nutrient availability is low, RSH, SAS, and SAH proteins respond by increasing (p)ppGpp synthase activity and/or decreasing (p)ppGpp hydrolase activity (46–48). The (p)ppGpp alarmone molecules activate the stringent response by binding to several protein targets and allosterically regulating their activity (49, 50). Targets of (p)ppGpp vary depending on the RSH enzyme and bacterial species but include proteins involved in DNA replication, translation, and nucleotide biosynthesis (51–54). The result is a halt in cell growth and the induction of amino acid biosynthesis and other stress responses to alleviate the nutrient limitation. In addition, (p)ppGpp can bind to RNA polymerase (RNAP) alone or with the RNAP-binding transcription factor DskA (55, 56), changing promoter affinity of RNAP that results in genome-wide changes in gene expression (57, 58). In other bacterial species, the stringent response positively regulates several virulence factors, including capsule production in *Staphylococcus aureus* (44, 59, 60). The two RSH proteins RelA and SpoT produced by *P. multocida* share a high level of identity with the equivalent proteins in other bacterial species, including the C-terminal regulatory region containing the TGS, helical, CC/ZFD, and ACT domains that have been shown in other bacterial species to autoregulate the (p)ppGpp synthase/hydrolase activity (61). No other RSH, SAS, or SAH proteins are present in the VP161 genome.

The TraDIS essential gene data indicated that insertions into the 5′ region encoding the (p)ppGpp hydrolase and synthase domains of spoT were lethal, and TraDISort data indicated that insertions in the 3′ regulatory region were associated with a high-density cell phenotype and an acapsular colony morphology. Further experiments showed that 3′ spoT mutants had reduced expression of hyaD, encoding the glycosyltransferase essential for HA capsule biosynthesis, and reduced amounts of HA capsule. We predict that mutants with insertions in the 3′ region of spoT produce functional truncated proteins that have lost autoregulatory function and have uncontrolled (p)ppGpp synthase activity leading to an increase in cellular levels of (p)ppGpp. Interestingly, the expression of the gene encoding the global regulator Fis is negatively regulated by the stringent response in *E. coli* (30, 62). If the *P. multocida* Fis were also negatively regulated by the stringent response, an increase in (p)ppGpp concentration would result in reduced fis expression in this species. However, our experiments showed that the expression of fis was unchanged in the *P. multocida* 3′ spoT mutant, indicating that the activation of the stringent response results in reduced expression of capsule genes in a Fis-independent manner. Other possible mechanisms of reduced capsule gene expression include reduced affinity of RNAP for capsule biosynthesis gene promoters or the hfg promoter when bound to (p)ppGpp. Alternatively, (p)ppGpp may allosterically regulate the activity of Hfq. Furthermore, as Hfq is an RNA chaperone known to interact with small RNAs to regulate gene translation (15), there may be Hfq-dependent small RNAs that regulate capsule biosynthesis gene expression in *P. multocida*. However, the density of the *P. multocida* VP161-Tn7 Himar1 insertions in our library was insufficient to identify any such small RNA genes (typically <250 nucleotides [nt] in length).

TraDIS data analysis of the VP161-Tn7 Himar1 mutant library following growth in rich media identified Himar1 insertions throughout the relA gene; however, following TraDISort analysis of the LD2 and HD2 cell layers, insertions in relA were identified only in the 3′ regulatory region of the gene. This suggested that relA mutants with insertions in the 5′ region of the gene, which would have disrupted (p)ppGpp synthase activity, were fully capsulated and partitioned only to the upper cell layer following the first gradient centrifugation. Moreover, the majority of the 3′ relA mutants recovered from the second gradient separated into the bottom, HD2 cell layer following centrifugation, correlating with an acapsular phenotype. In *E. coli*, removing the 3′ regulatory region of relA or disrupting the ACT domain
results in truncated proteins that have increased (p)ppGpp synthesis activity (25, 28). We predict that 3’ relA mutants have increased (p)ppGpp synthesis due to the loss of the autoregulatory ACT domain. As above, we predict this increase in (p)ppGpp results in altered protein activity or regulation of genes required for HA capsule production. Attempts were made to construct a directed TargeTron mutant in the 3’ end of relA, but these were unsuccessful despite multiple attempts.

Several other genes identified as important for capsule production in *P. multocida* were also associated with activation of the stringent response, including *ptsH*, *ppx*, *tufA_1*, and *tufA_2*. The gene *ptsH* encodes HPr, a component of the phosphoenolpyruvate:sugar phosphotransferase (PTS) system that imports and phosphorylates carbohydrate monomers. No other PTS system genes were identified as important for capsule production by the TraDISort analysis, suggesting that import and phosphorylation of carbohydrate monomers by the PTS system are not required for HA production in strain VP161. In *E. coli*, HPr indirectly regulates SpoT activity by binding to phosphorylated Rsd during carbon downshift, stopping Rsd from binding to SpoT and inducing (p)ppGpp synthesis (63). In *Caulobacter crescentus* during glutamine starvation, HPr is required for phosphorylating ElaA<sup>100</sup>, which then binds to SpoT (64). HPr in *P. multocida* may have a similar role, binding to or controlling the phosphorylation state of proteins that regulate the activity of SpoT and/or RelA. The *ppx* gene is also predicted to be involved in the stringent response, as it encodes an exopolysphatase that cleaves terminal phosphates from long-chain polyphosphate (polyP) (65). Phosphate starvation in *E. coli* results in an increase in (p)ppGpp concentration due to altered SpoT activity (66). Disrupting *ppx* likely reduces available phosphate within the cell and increases (p)ppGpp synthesis and/or decreases (p)ppGpp hydrolysis by SpoT. The genes *tufA_1* and *tufA_2* both encode elongation factor Tu (EF-Tu), which delivers charged tRNAs to the ribosome by forming a complex with the tRNA and GTP (67). The EF-Tu-tRNA-GTP complex competes with RelA bound to uncharged tRNAs for binding to the ribosome A site. Reduced EF-Tu levels from inactivation of either *tufA_1* or *tufA_2* may result in an increased number of RelA-uncharged tRNA complexes interacting with the ribosome, and this would result in increased (p)ppGpp synthesis.

In addition to *tufA_1* and *tufA_2*, other genes encoding translation factors were identified by TraDISort as important for HA capsule biosynthesis in *P. multocida* strain VP161, including *efp*, *empA*, and *empB*. The gene *efp* encodes elongation factor P (EF-P), which restores ribosomes stalled during translation of polyproline stretches (68, 69). The genes *empA* and *empB* encode an EF-P lysine l-lysyltransferase that adds R-lysine to Lys34 within EF-P and a lysine 2,3-aminomutase that converts l-lysine to R-lysine, respectively (70, 71). Both *empA* and *empB* are required for correct EF-P function in *E. coli* (70). Mutation of *efp*, *empA*, or *empB* in *E. coli* leads to reduced production of several capsule-associated genes, some of which are homologs of genes identified as important for *P. multocida* capsule production by TraDISort analysis, including Rnb and several ATP synthase subunit proteins (72). Therefore, EF-P is likely important for capsule production in *P. multocida*, as it facilitates the production of other proteins crucial for HA capsule production.

Overall, we have identified genes required for growth in rich media and comprehensively characterized HA capsule production in *P. multocida* strain VP161. To our knowledge, this is the first time a whole-genome saturation mutagenesis approach has been used to identify essential genes and those associated with the production of extracellular capsule in *P. multocida*, an important virulence factor for this pathogen. Importantly, the stringent response was shown to be a negative regulator of capsule biosynthesis genes. The RSH proteins, SpoT and RelA, critical for the synthesis and control of the levels of the alarmone nucleotides (p)ppGpp and the associated regulatory proteins, TufA<sub>1</sub>, TufA<sub>2</sub>, HPr, and PXX, were all identified by TraDISort as important for HA capsule production. Further work is required to elucidate the exact mechanism by which these stringent response proteins precisely control capsule production in this pathogen.

**MATERIALS AND METHODS**

**Bacterial strains, plasmids, and culturing conditions.** Bacterial strains and plasmids used in this study are listed in Table S1 in the supplemental material. *P. multocida* strains were cultured in HI broth (Oxoid). *E. coli* strains were cultured in lysogeny broth (LB). *P. multocida* and *E. coli* were grown at 37°C.
with liquid broth cultures shaken at 200 RPM. Solid media were produced by adding 1.5% (wt/vol) agar to the media. When required, media were supplemented with appropriate antibiotics at the following concentrations: kanamycin (50 μg/mL), spectinomycin (50 μg/mL), and tetracycline (2.5 μg/mL).

**DNA manipulations, Sanger sequencing, and oligonucleotides.** Genomic DNA was extracted using the HiYield genomic DNA minikit (RBC bioscience) and plasmid DNA was extracted using the NucleoSpin plasmid kit (Macherey-Nagel), as per the manufacturer’s instructions. DNA samples were quantified using a NanoDrop 1000 Spectrophotometer or Qubit fluorometer (Thermo Fisher Scientific). PCRts were performed using Taq DNA polymerase (Roche), Phusion high-fidelity DNA polymerase (NEB), or KAPA HiFi HotStart DNA polymerase (Roche) as per the manufacturer’s instructions. Restriction digestion and ligation reactions were performed using enzymes from Roche or New England Biolabs, as per the manufacturer's instructions. Oligonucleotides used in this study were synthesized by Sigma-Aldrich and are listed in Table S4 in the supplemental material. Sanger sequencing reactions were performed using plasmid or genomic DNA as the template with BigDye Terminator version 3.1 (Applied biosystems) as described previously (73). Sanger sequencing data were analyzed using Geneious Prime (Biomatters).

**Transformation and conjugation.** Transformation of *P. multocida* was performed as described previously (14), with the electroporation conditions modified to 2 kV, 600 Ω, and 25 μF. Conjugation experiments were performed by filter mating with *E. coli* donor and *P. multocida* recipient VP161-Tn7 (kanamycin-resistant parent, see supplemental material for construction details) as follows: cultures were each grown to mid-exponential stage (OD600 of ~0.5) and washed in 1x phosphate-buffered saline (PBS), and then 1 mL of each donor and recipient culture was mixed and filtered through a 0.22-μm nylon filter using a syringe and Swinnex filter holder. The filter containing the cells was then aseptically placed (cell-side up) onto an HI plate without antibiotics and incubated at 37°C for 1 h. Following incubation, we resuspended the cells in 1 mL of HI by vortexing the filter and then plated them onto HI agar plates containing spectinomycin, or spectinomycin and kanamycin to select for growth of donor, recipient, or transconjugants, respectively. Following overnight incubation at 37°C, viable counts were performed to determine CFU/mL of donor, recipient, and transconjugants.

**Site-directed mutagenesis of selected genes using TargeTron.** Site-directed mutagenesis of specific genes in the VP161 genome was performed using the *P. multocida* TargeTron plasmid pAL953 targeted to the appropriate gene using the intron site finder Clostron (74) as described previously (16, 73). A full list of rearget primers is shown in Table S4. TargeTron intron insertion into the target gene was confirmed by PCR amplification of the target region using two PCRs. The first reaction used primers flanking the gene of interest and the second used the TargeTron-specific primer paired with the appropriate primer flanking the target gene (Table S4). Direct Sanger sequencing was also conducted using genomic DNA as the template with the TargeTron-specific outward-facing primer BAP6544 to confirm that only a single intron insertion was present in the genome at the specific target site.

**Construction of complementation vectors.** We complemented selected *P. multocida* mutants by providing a wild-type copy of the target gene on the *P. multocida* expression plasmid pAL99S, which conferred spectinomycin resistance and was suitable for the TargeTron mutants, or pAL99T, which contained tetracycline resistance and was suitable for the *Himar1* mutants (Table S1). To produce each of the complementation vectors, we PCR-amplified the target gene using Phusion polymerase with *P. multocida* VP161 genomic DNA as the template, a forward primer containing an appropriate restriction site followed by a ribosome binding site, and a reverse primer with an appropriate restriction site (Table S4). Purified PCR products were digested with the appropriate restriction enzymes and ligated into similarly digested pAL99S or pAL99T. Sanger sequencing was confirmed to show that each plasmid complementation vector contained a wild-type copy of the target gene. To produce each complementation strain, we used the appropriate plasmid (Table S1) to transform the appropriate *P. multocida* mutant strain by electroporation. To produce mutant strains harboring empty vector (control strains), pAL99S or pAL99T was used to separately transform each mutant (Table S1).

**Generation of the *P. multocida Himar1* transposon mutant library and growth in rich media.** *P. multocida* transposon mutant libraries were produced by delivery of the Himar1 transposon-containing suicide vector pAL614 into *P. multocida* by conjugation between donor *E. coli* strain AL2972 and *P. multocida* strain VP161-Tn7 (Table S1). Following overnight growth, Himar1 transconjugants were scraped from HI agar plates supplemented with both kanamycin and spectinomycin, pooled in 15% glycerol broth, and then stored at −80°C in 1-mL aliquots containing ~6.45 × 107 VP161-Tn7 Himar1 mutants. For identification of genes essential for growth in rich media, two aliquots of the VP161-Tn7 Himar1 mutant library (named A and B) were grown overnight in 50 mL of HI broth, subcultured into 50 mL of fresh HI broth, and grown until mid-exponential growth phase (OD600 of ~0.5). Two 1-mL aliquots of surviving mutants were recovered from each mutant library culture (A1, A2, B1, B2) and used for TraDIS analysis.

**Discontinuous Percoll gradient centrifugation.** Separation of capsular and acapsular mutant strains in the VP161-Tn7 Himar1 mutant library was performed using Percoll density gradient centrifugation as described previously (17, 75) with the following modifications. For each discontinuous gradient, an 80% Percoll (in PBS wt/vol) bottom layer, a 40% Percoll middle layer, and a 10% Percoll top layer were added to a polyallomer centrifuge tube (17 mL, 16 mm by 102 mm, Beckman). Cells were grown to mid-exponential-phase growth, and 20 mL of each culture was centrifuged, resuspended in 1 mL of PBS, and gently loaded onto the top of each gradient. Gradients were then centrifuged in a fixed angle rotor for 30 min at 3,000 × g. Cell layers of interest were recovered by syringe aspiration. To remove excess Percoll, the cell pellet was washed with 1 mL of PBS, centrifuged, and resuspended in 1 mL of PBS. Three replicates of the Himar1 library were subjected to discontinuous gradient centrifugation as described above. Following the first centrifugation, the top layer of cells consisting of the capsulated population of the mutant library was not examined further. Cells recovered from the bottom cell layer were grown to
mid-exponential growth stage for 4 h in HI broth (37°C with shaking at 200 rpm) and subjected to a second discontinuous Percoll gradient centrifugation using the same conditions as described above. The top (LD2) cell layer and the bottom (HD2) cell layer generated by the second gradient centrifugation were collected separately and used for TraDISort analysis.

**TraDIS library production, sequencing, and data analysis.** TraDIS libraries for Illumina sequencing were produced as described previously (76) with minor modifications. A summary (Fig. S1) and full details of TraDIS library production, Illumina MiSeq sequencing, and data analysis methods are provided in the supplemental material.

TraDIS libraries were submitted to the Micromanon Genomics facility at Monash University and sequenced on an Illumina MiSeq v2 using 150-bp single-end sequencing with the custom Himar1 sequencing oligonucleotide BAP0842 (Table S4). Poor-quality reads and Illumina adapter sequences were removed from the sequencing data using Trimmomatic v0.38. The sequencing reads were then analyzed using Bio-TraDIS v1.4.1 and modified scripts as described previously (17, 76). All reads beginning with a correct Himar1 transposon tag (5'-CAACCTGT-3') were aligned to the reference *P. multocida* VP161 genome sequence, with the first base of each alignment deemed to be a transposon insertion site (VP161 NCBI database accession no. CP048792.1). Genes essential for growth in *in vitro* in HI were defined as those with an insertion index (number of unique insertions relative to gene length) of less than 0.0134, with data from all rich media TraDIS libraries (A1, A2, B1, B2) combined for analysis. Specific metabolic pathways for each of the *P. multocida* genes predicted to be essential were identified by comparison to the Kyoto Encyclopedia of Genes and Genomes (KEGG) database by BlastKOALA (77) and to the UniProt database by BLASTp (78). Genes associated with high-density/reduced capsule production were identified by comparing normalized read counts for each gene in TraDIS libraries generated from the upper, low-cell-density layer (LD2 cell layer) with TraDIS libraries generated from the lower, high-density cell layer (HD2 cell layer). Any gene with a significant increase in the number of Himar1 insertion sites identified in mutants recovered from the HD2 cell layer (log2 fold change > 2, q value of <0.001, insertion index ratio of >2.0) compared to that in mutants recovered from the LD2 cell layer was identified as important for capsule production. Homologs of genes identified by TraDISort were identified by comparison to the Kyoto Encyclopedia of Genes and Genomes (KEGG) database by BlastKOALA (77) and to the UniProt database by BLASTp (78).

**Measurement of hyaluronic acid capsule production.** Capsular polysaccharide was extracted from *P. multocida* cells grown in broth and measured as described previously (14) with minor modifications. Briefly, cells were harvested from 1 mL of mid-exponential culture using centrifugation at 13,000 × g for 10 min, washed with PBS, and resuspended in 1 mL of PBS. The cells were then heated at 42°C for 1 h and pelleted by centrifugation at 13,000 × g for 10 min, and the supernatant containing capsular polysaccharide was recovered. For each sample, 100 μL of supernatant was mixed with 900 μL of stains-all solution (14) and the absorbance at 640 nm was measured. The absorbance of each sample was compared to hyaluronic acid standards (between 2.5 μg/mL and 40 μg/mL) to determine the concentration of hyaluronic acid.

**Scanning electron microscopy.** SEM was performed on *P. multocida* strains grown to mid-exponential growth phase (OD600 of 0.5 to 0.6). Cells were filtered onto a 0.4-μm aperture polycarbonate membrane and prepared for imaging using lysine acetate-ruthenium red-osmium fixation, critical point drying, and sputter coating as previously described with minor modifications (79). For the fixation solution, 100 μL of 16% formaldehyde was used in place of 80 μL of 25% formaldehyde solution. Cells were imaged using a field emission scanning electron microscope Nova NanoSEM 450 with an accelerating voltage of 5 kV in secondary electron mode with a working distance of ~4.5 mm. Images were taken under immersion mode through the lens detector. Cell widths were measured from electron micrographs generated by SEM using the Fiji package for ImageJ. Significant differences in cell widths were determined using an unpaired t test.

**RNA extraction and qRT-PCR.** RNA was isolated from *P. multocida* cells grown to mid-exponential growth phase (OD600 of ~0.4) in biological quadruplicate using Trizol reagent (ThermoFisher Scientific) as per the manufacturer’s instructions. RNA samples were DNase treated using the RNeasy kit (Qiagen) as per the manufacturer’s instructions, and then RNA was purified using phenolchloroform extraction with 5 PRIME phase lock gel tubes (Quanta Biosciences) following the manufacturer’s instructions followed by sodium acetate-ethanol precipitation. cDNA was synthesized for qRT-PCR using the AffinityScript cDNA synthesis kit (Agilent Technologies) as per the manufacturer’s instructions. Each sample was used in cDNA synthesis reactions with (+ RT) or without (− RT) reverse transcriptase. The qRT-PCRs were performed using Brilliant II SYBR green qPCR master mix (Agilent Technologies) per the manufacturer’s instructions with primers specific to *P. multocida* strain VP161 gyrB, hyaD, or fis (Table S4), with reactions performed in an Aria Mx real-time PCR system (Agilent). Biological quadruplicate cDNA samples were each measured in technical duplicate. The − RT samples were analyzed to ensure no products were amplified before 35 cycles, and the melt curves from +RT reactions were analyzed to ensure only a single product was amplified in each reaction. Significant differences in gene expression were determined using a Mann-Whitney U test.

**Data analysis.** Sequencing data and analysis output files have been uploaded to the NCBI Gene Expression Omnibus database under accession number GSE198087.

### SUPPLEMENTAL MATERIAL

Supplemental material is available online only.

**SUPPLEMENTAL FILE 1** PDF file, 1.2 MB.
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