MICROBIAL PATHOGENICITY

Distribution of type III secretion gene clusters in *Burkholderia pseudomallei*, *B. thailandensis* and *B. mallei*

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*Burkholderia pseudomallei*, the causative agent of melioidosis, carries a cluster of genes closely related in organisation to the type III secretion (TTS) system gene clusters of the plant pathogens *Ralstonia solanacearum* and *Xanthomonas* spp. The TTS gene cluster (TTS1) is present only in *B. pseudomallei* and not in avirulent *B. thailandensis*. Adjacent to the gene cluster encoding putative secreton structural proteins lie a number of open reading frames (ORFs) encoding putative proteins with little or no homology to known proteins, with the exception of one predicted protein with homology to *Pseudomonas syringae* HrpK. In both *R. solanacearum* and *Xanthomonas* spp., genes in this location encode secreted effector proteins. RT-PCR analysis indicated that TTS genes, including two of these ORFs, are expressed in broth at 37°C. Analysis of genome sequence data identified a second cluster of TTS genes (TTS2) present in both *B. pseudomallei* and *B. mallei* (99% identity). However, *B. mallei* appears to lack the TTS1 gene cluster. PCR assays indicated that TTS2 was also present in *B. thailandensis*. TTS1 and TTS2 are similar in gene organisation, but nucleotide sequences are sufficiently divergent to suggest that the two TTS systems may have different roles.

Introduction

*Burkholderia pseudomallei* is the causative agent of melioidosis, a disease that can manifest itself as an acute, subacute or chronic infection [1, 2]. The acute form of the disease accounts for a significant proportion of often fatal, community-acquired septicaemia in areas of south-east Asia and Australia. *B. pseudomallei* also causes a subacute febrile illness leading to systemic abscess formation involving various organs. The chronic form of the disease is evident from discovery at autopsy or by subsequent activation causing other forms of the disease. It is known that *B. pseudomallei* can enter a dormant state following initial infection or recovery from clinical disease [3], and that dormant cells can be triggered, leading to relapse or the onset of acute melioidosis [4]. The mechanisms underlying these different clinical manifestations and the triggers causing the more serious forms of the disease are poorly understood.

It has been suggested that environmental *B. pseudomallei* isolates can be separated on the basis of a number of factors into two biotypes, best defined by the ability to assimilate arabinose. There is evidence that *B. thailandensis* (Ara⁺ strains) [5] are avirulent whereas Ara⁻ strains constitute melioidosis-causing ‘true’ *B. pseudomallei* [6]. It is possible to discriminate between the two biotypes by molecular approaches such as *fliC* variation [7, 8] or multiplex PCR assays [9]. Furthermore, differences in genomic macrorestriction patterns between the two biotypes have been reported [10]. As the mechanisms of *B. pseudomallei* virulence are poorly understood, there is considerable interest in the identification of the specific factors determining the greater virulence of *B. pseudomallei*. Indeed, subtractive hybridisation has been used to identify some virulence genes present in *B. pseudomallei* but absent from *B. thailandensis* [11].

Differences in virulence between closely related bacterial pathogens are often attributable to the presence of pathogenicity islands (PIs) in virulent strains. Type III secretion (TTS) systems have been implicated in the pathogenicity of several gram-negative bacterial pathogens, including intracellular bacteria such as *Shigella*...
spp. [12, 13]. The genes encoding TTS systems and secreted effector proteins are associated with PIs and, therefore, are prime candidates for involvement in the greater virulence of some strains over other related strains.

A previous study identified, in B. pseudomallei, a cluster of putative genes homologous to those encoding HpaP, HrcQ (HrpQ), HrcR (HrpT), HrcS (HrpU) and HrpV; all TTS-associated proteins in the plant pathogen Ralstonia solanacearum [14]. A further study, based on the detection of a single locus, reported a link between the presence of putative TTS system genes and the Ara− phenotype, the more virulent biotype of B. pseudomallei [15]. There was one exception to the link, a B. thailandensis strain (E27) which was positive by both PCR and dot-blot assay.

This study reports the nucleotide sequence of the entire putative structural region of the B. pseudomallei TTS1 gene cluster, identifies ORFs that may encode potential effectors and provides evidence that TTS genes are expressed. The study also presents more comprehensive evidence of the link between TTS1 genes and the Ara− biotype and reports evidence for a second TTS gene cluster (TTS2) common to B. pseudomallei and B. mallei, the causative agent of glanders.

Materials and methods

Bacterial strains

The strains used in this study have been described previously [15] and are listed in Table 1. B. pseudomallei and B. thailandensis strains were maintained on nutrient agar.

Nucleotide sequence of TTS gene PI

Cosmid clones c503-23, c503-5 and c503-V1 were previously identified as containing a TTS gene cluster [14], for which we propose the name TTS1. Fragments generated by HindIII digestion were subcloned from these cosmid clones into HindIII-digested pUC19 (Helena Biosciences). These and other subclones were sequenced by primer-walking with vector and internal oligonucleotide primer sequences.

Computer analyses

Nucleotide sequence alignments, G + C mol% values, determination of amino acid composition, predicted protein mass and alignments of predicted proteins with other related proteins (retrieved from EMBL, GenBank, PIR or SwissProt [16]) were performed with BESTFIT, GAP, PILEUP, PRETTY, COMPOSITION, PEPTIDE-SORT and FASTA from the GCG sequence analysis software package (Genetics Computer Group, University of Wisconsin). Hydrophobicity profiles and predictions of membrane-spanning regions were obtained with the PEPTIDESTRUCTURE and PEPPLOT programmes, employing the hydropathy measure of Kyte and Doolittle [17] and the measure of helical hydrophobic moment [18]. The programme MOTIFS was used to identify motifs within the predicted protein sequences. BLAST searches were conducted via the site http://www.ncbi.nlm.nih.gov/blast/blast.cgi [19]. Genome sequence data for B. pseudomallei, used to complete orf13, and for TTS2, were obtained from the Sanger Centre website http://www.sanger.ac.uk/Projects/B_pseudomallei/. These sequence data were produced by the B. pseudomallei Sequencing Group at the Sanger Centre and can be obtained from ftp://ftp.sanger.ac.uk/pub/pathogens/bps/. Preliminary sequence data for the genome sequence of B. mallei were obtained from the Institute for Genomic Research website at http://www.tigr.org. Searches of these data were conducted via the BLASTX or TBLASTN facilities provided at these websites.

Dot-blot and PCR assays

Genomic DNA extraction from B. pseudomallei and dot-blot hybridisation of genomic DNA (c. 0.25 μg) were performed as described previously [15]. Digoxigenin-11-2′-dUTP (DIG) (Roche)-labelled probes were made either by PCR amplification with c503-5 or c503-23 DNA as template in the presence of 60 μM DIG, as described previously [15], or by labelling amplicons

<table>
<thead>
<tr>
<th>Strains</th>
<th>orf11 PCR</th>
<th>orf7 PCR</th>
<th>bpscJ PCR</th>
<th>orf4/orf5 PCR</th>
<th>orf1-bpscQ gap PCR</th>
</tr>
</thead>
<tbody>
<tr>
<td>B. pseudomallei</td>
<td>E503, E504, E505, E506, 204 (E955), 576 (E957), E25 (E958), E8 (E960)</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>B. thailandensis</td>
<td>E27 (E956)</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>+/-</td>
</tr>
<tr>
<td>E82 (E959), E32, E111, E125, E132, E135, E202, E216, E251, E253, E254, E255, E260</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Neisseria meningitidis</td>
<td>C311</td>
<td>ND</td>
<td>ND</td>
<td>-</td>
<td>ND</td>
</tr>
</tbody>
</table>

ND, not done.
RNA extraction and RT-PCR

*B. pseudomallei* E503 (also known as strain 10-705) and *B. thailandensis* E27 were cultured in Luria broth at 37°C with and without the addition of 10 mM nitrilotriacetic acid (NTA) until they reached exponential growth (A600 of 0.5). Cells were harvested by centrifugation and RNA was extracted with an RNaseasy kit (Qiagen). Total RNA was treated with DNAse following the manufacturer's instructions (Life Technologies) and reverse transcribed into cDNA with M-MLV reverse transcriptase (Promega) and 100 μM of each primer, 200 nM of each primer, 1250 μM of each nucleotide (dATP, dCTP, dGTP, dTTP). Amplifications were performed in an Eppendorf MasterCycler thermal cycler for 30 cycles consisting of 30 s at 95°C, annealing temperature (1 min) and 72°C (2 min) with an additional extension time at 72°C (10 min) following completion of the 30 cycles. At the end of the amplification, 5-μl samples were subjected to electrophoresis on a standard agarose 1.0% w/v gel to confirm the presence of an amplified product.

RT-PCR products visualised after agarose 1% w/v gel electrophoresis were often faint. To ensure that there was convincing discrimination between positive and negative PCR assays, gels were Southern blotted by standard procedures and probed with equivalent labelled PCR products generated from genomic DNA. After purification through Microspin S-400 HR columns (Amersham Pharmacia Biotech), PCR products were labelled with DIG with a DIG-High Prime labelling system (Roche).

Accession number

The *B. pseudomallei* TTS1 system gene cluster nucleotide sequence reported in this paper can be found under the GenBank accession no. AF074878.

Results

Map of the TTS genes

A map of the strain E503 region sequenced from three overlapping cosmid clones is presented in Fig. 1. It has been possible to extend further to complete *orf13* with additional sequence data produced by the *B. pseudomallei* Sequencing Group at the Sanger Centre (obtained from ftp://ftp.sanger.ac.uk/pub/pathogens/bps). The region spanning *bpsscC* to *orf3* bears a close resemblance in gene organisation to the TTS gene clusters encoding the Hrp secretions of the plant pathogens *R. solanacearum* [20, 21] and *Xanthomonas* spp. (*X. campestris* pv. *vesicatoria* or *X. oryzae* pv. *oryzae*) [22–24].

Protein sequence alignments and motifs

Table 3 indicates the sequence similarities of the equivalent predicted proteins from *B. pseudomallei* and *R. solanacearum*. In most cases there was good alignment between the predicted proteins of the two organisms. BLASTP searches indicated that most of

<table>
<thead>
<tr>
<th>Target DNA (amplicon size)</th>
<th>Oligonucleotide primer sets for PCR</th>
<th>Annealing temperature (°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>orf11</em> (706 bp)</td>
<td>PM122: 5'-ATGCACCAATTGCGGCCGTTTC</td>
<td>55</td>
</tr>
<tr>
<td><em>orf10</em> (219 bp)</td>
<td>PM121: 5'-AGGCCTGTACACCGTCCAAG</td>
<td>60</td>
</tr>
<tr>
<td><em>orf7</em> (1059 bp)</td>
<td>PM127: 5'-CGACAGGACTGGAGAAAG'</td>
<td>60</td>
</tr>
<tr>
<td><em>bpjc</em> (481 bp)</td>
<td>PM123: 5'-TGCCATCCATTACATCAAAAG</td>
<td>55</td>
</tr>
<tr>
<td><em>orf4/orf5</em> (646 bp)</td>
<td>PM125: 5'-CGACAGGACTGGAGAAAG'</td>
<td>50</td>
</tr>
<tr>
<td><em>orf1-bpsscQgap</em> (490 bp)</td>
<td>PM124: 5'-TAAGACAGGACCCGAAAG</td>
<td>65</td>
</tr>
<tr>
<td><em>orf13</em> (722 bp)</td>
<td>PM126: 5'-TGCCATCCATTACATCAAAAG</td>
<td>48</td>
</tr>
<tr>
<td><em>bpsscC2</em> (TTSS) (488 bp)</td>
<td>PM127: 5'-CGACAGGACTGGAGAAAG'</td>
<td>63</td>
</tr>
<tr>
<td><em>sopE</em> homologue (TTSS) (608 bp)</td>
<td>PM128: 5'-TAAGACAGGACCCGAAAG</td>
<td>55</td>
</tr>
</tbody>
</table>

*PMXBA3 and PMBAM contain terminal restriction sites that are not relevant to this study.*
the B. pseudomallei predicted TTS1 proteins matched best with homologues from either R. solanacearum or Xanthomonas spp. The exceptions to this were Orf2, Orf3, Orf6 and Orf7, for which the BLAST searches identified no significant similarity. Similarity/identity values were particularly high for the 11 proteins belonging to families whose members are known to be conserved between TTS systems (BpscC, BpscD, BpscN, BpscL, BpscI, BpscQ, BpscR, BpscS, BpscT, BpscU and BpscV). There was also strong homology between Orf8 and R. solanacearum HrpB, a transcriptional activator responsible for regulating TTS genes in R. solanacearum. Further analysis with the NCBI Conserved Domain Database at http://www.ncbi.nlm.nih.gov/Structure/cdd/cdd.shtml indicated a highly significant match with bacterial regulatory helix-turn-helix proteins of the AraC family in the C-terminal domain (Orf8 residues 394–483). Members of the AraC-XylS family are basic. Whereas HrpB has a pI value of 8.56, Orf8 is only weakly basic, with a pI of 7.06. Even amongst those proteins exhibiting lower similarity/identity values there are similarities in predicted pI and mol. wt values (data not shown). The least convincing match between the two TTS systems was found with Orf6, which does not align well with HrpH, although the predicted pl values are similar.

Table 3. Properties of predicted TTS1 proteins and comparison with equivalent R. solanacearum and TTS2 proteins

<table>
<thead>
<tr>
<th>Protein*</th>
<th>R. solanacearum % similarity/% identity†</th>
<th>TTS2 % similarity/% identity</th>
<th>Versinia homologue</th>
<th>Predicted location/properties</th>
</tr>
</thead>
<tbody>
<tr>
<td>BpscC (HrpA/HrcC)</td>
<td>56.2/45.8</td>
<td>66.4/54.1 (C2b)</td>
<td>YscC</td>
<td>Outer-membrane secretin</td>
</tr>
<tr>
<td>Orf8 (HrpB)</td>
<td>47.7/36.1</td>
<td>50.7/42.1</td>
<td>None</td>
<td>Transcriptional activator</td>
</tr>
<tr>
<td>BpscT (HrpC/HrcT)</td>
<td>50.6/41.4</td>
<td>56.5/48.5</td>
<td>YscT</td>
<td>Inner membrane</td>
</tr>
<tr>
<td>Orf7 (HrpD)</td>
<td>41.4/29.9</td>
<td>40.7/33.5</td>
<td>None</td>
<td>Unknown</td>
</tr>
<tr>
<td>BpscN (HrpE/HrcN)</td>
<td>71.5/63.5</td>
<td>79.9/72.2</td>
<td>YscN</td>
<td>ATPase</td>
</tr>
<tr>
<td>Orf6 (HrpH)</td>
<td>37.0/33.0</td>
<td>40.9/35.6</td>
<td>None</td>
<td>Unknown</td>
</tr>
<tr>
<td>BpscI (HrpI/HrcI)</td>
<td>61.5/53.2</td>
<td>64.5/59.7</td>
<td>YscI</td>
<td>Lipoprotein</td>
</tr>
<tr>
<td>Orf5 (HrpI)</td>
<td>35.5/25.8</td>
<td>40.2/31.5</td>
<td>None</td>
<td>Unknown</td>
</tr>
<tr>
<td>Orf4 (HrpK)</td>
<td>41.0/30.8</td>
<td>42.9/35.4</td>
<td>None</td>
<td>Unknown</td>
</tr>
<tr>
<td>BpscU (HrpN/HrcU)</td>
<td>59.8/49.3</td>
<td>59.5/47.3</td>
<td>YscU</td>
<td>Inner membrane</td>
</tr>
<tr>
<td>BpscV (HrpO/HrcV)</td>
<td>69.3/61.1</td>
<td>77.6/71.2</td>
<td>YscV (LcrD)</td>
<td>Inner membrane</td>
</tr>
<tr>
<td>Orf1 (HpaP)</td>
<td>37.6/29.4</td>
<td>43.7/39.2</td>
<td>None</td>
<td>Unknown</td>
</tr>
<tr>
<td>BpscQ (HrpQ/HrcQ)</td>
<td>40.4/29.8</td>
<td>47.5/42.0</td>
<td>YscQ</td>
<td>Translocase</td>
</tr>
<tr>
<td>BpscR (HrpR/HrcR)</td>
<td>74.4/66.0</td>
<td>82.0/73.7</td>
<td>YscR</td>
<td>Inner membrane</td>
</tr>
<tr>
<td>BpscS (HrpU/HrcS)</td>
<td>64.0/55.8</td>
<td>70.1/57.5</td>
<td>YscS</td>
<td>Inner membrane</td>
</tr>
<tr>
<td>Orf2 (HrpV)</td>
<td>33.8/26.8</td>
<td>35.3/29.9</td>
<td>None</td>
<td>Unknown</td>
</tr>
<tr>
<td>BpscD (HrpW)</td>
<td>40.9/32.7</td>
<td>49.8/43.8</td>
<td>YscD</td>
<td>Inner membrane</td>
</tr>
<tr>
<td>Orf3 (HrpX)</td>
<td>36.4/29.9</td>
<td>47.6/37.8</td>
<td>None</td>
<td>Unknown</td>
</tr>
</tbody>
</table>

*Protein designations shown in parentheses relate to R. solanacearum Hrp proteins. Both the original and currently accepted designations for these proteins are given.
†% similarity/% identity values were calculated by GAP/BESTFIT with the following parameters: gap creation penalty 8, gap extension penalty 2.
‡Orf6 and HrpH do not align well. The figures presented were calculated by BESTFIT over a sequence of 113 amino acids.
§BpscJ data were calculated for the whole predicted peptide sequence not for the protein predicted after cleavage of the lipoprotein signal sequence.

Fig. 1. Map of the TTS1 and TTS2 gene cluster. The location and lengths of predicted genes are shown. Arrows indicate the direction of transcription. The location of an imperfectly repeated sequence in TTS1 is also indicated. TTS2 genes have been labelled according to comparison with equivalent TTS1 genes. 12* and 8* indicate ORFs with homology to TTS1 orf12 and orf8 respectively. Genes with homology to histidine kinases (hk) and response regulators (rr) from known two-component regulatory systems are also indicated. Only genes with predicted protein sequences showing significant homology to proteins in the database are shown. The gap between orf8 and hk is to scale but may contain other ORFs. The organisation of TTS2 genes is based on data obtained from the unfinished B. pseudomallei and B. mallei genome sequence projects.
The predicted properties of BpscQ, BpscR, BpscS, Orf1 and Orf2 have been discussed previously [14]. BpscN had one of the highest sequence identities with its R. solanacearum homologue (HrcN) and contains an ATP-binding 'A' consensus sequence (P loop; 182-APAVGAVKS) and an ATP synthase αβ signature (362-PAID/LGSL) typical of TTS-associated ATPases. BpscJ has a prokaryotic membrane lipoprotein lipid attachment site (23-VLALSVLLAGC) similar to those found in the equivalent R. solanacearum (HrcJ) and Yersinia (YscJ) proteins. As was the case with their R. solanacearum homologues [20], hydrophy analysis predicted that Orf7 and BpscL were highly hydrophilic.

Analysis with the programme MOTIFS indicated that the predicted protein BpscC contains a protein D signature common to type II and type III secretion outer-membrane secretin proteins (536-QGSLLIAGYSTDKRANGVAGVPWLSKIPPLLLGF). BpscC also contained a putative signature characteristic of bacterial regulatory proteins of the LysR family. Proteins of the LysR family possess a potential 'helix-turn-helix' DNA-binding motif in their N-terminal section. The amino acid sequence pattern used to detect these proteins is a consensus sequence derived from the complete helix-turn-helix motif and the next 10 residues. In BpscC, the consensus sequence was not detected in the N-terminal region and therefore is probably not significant.

It has been predicted that the N-terminal half of secretins faces the periplasm and this region is not well conserved [25]. The C-terminal region (β domain) is more conserved and is predicted to contain several transmembrane β-strands whose likely location is the outer membrane. BpscC aligns strongly with homologues in the conserved C-terminal region and contains the consensus motif (V,I)PXL(S,G)XIPXXGXF common to proteins of this family [26]. BpscV aligns particularly strongly with the N-terminal region of R. solanacearum HrcV. In HrcV and homologues this region of the protein is predicted to be hydrophobic, containing a number of membrane-associated helices.

Downstream of bpscC lies a region containing a 264-bp imperfect triple direct repeat sequence that marks the boundary between the gene cluster with homology to R. solanacearum and Xanthomonas spp. and several putative ORFs (Table 4) likely to include genes encoding secreted proteins if the B. pseudomallei TTS gene cluster continues its resemblance to that of the plant pathogens. It is possible to identify an ORF within this repeated region. If genuine, this ORF, transcribed in the same direction as bpscC and orf9, would encode a glycine-rich 461 residue peptide with predicted mol. wt and pl values of 45764 Da and 4.16, respectively, and no significant homology to known proteins. As the repeated sequences are in-frame with each other, any such peptide would contain regions of repeated amino acid sequences.

Orf9 is predicted to be a largely hydrophilic protein with a C-terminal hydrophobic α-helical domain. BLAST searches identified homology with P. syringae HrpK (32% identity/46% positives between residues over 338 amino acid residues (94–418); accession no. AF232004), a hydrophilic protein for which a function has yet to be ascribed. However, P. syringae HrpK does not contain a C-terminal hydrophobic domain. BLAST searches indicated that Orf10 shared best homology with acetyltransferases. The highest match was 35% identity and 51% positives over 65 amino acid residues (Orf10 73–132) against a ribosomal-protein-alanine N-acetyltransferase of Bacillus halodurans (accession no. AP001509). BLAST searches failed to identify any significant homology between Orf11 and known proteins. MOTIFS indicated that Orf11 contains a putative actin-binding domain signature (E,Q)x2(A,T,V)(F,Y)x2WxN). In most proteins identified as possessing this motif, including α-actinin, the signature resides within the N-terminal region. An exception to this rule is fimbrin, where the signature is located in duplicate in the C-terminal region. In Orf11, the signature is located between residues 551 and 560. Orf11 also contains a putative ATP/GTP-binding site – P-loop; (A,G)x4GK(S,T) – typical of kinases, ADP-ribosylating factors and a number of proteins involved in secretion.

Downstream of Orf3 lies other potential genes. The predicted protein of Orf12 shows strong homology to HpaB (accession no. BAB07868), a protein of the

Table 4. Predicted properties of putative proteins encoded by ORFs downstream of bpscC and orf3

<table>
<thead>
<tr>
<th>ORF</th>
<th>Length</th>
<th>Mol. wt (Da)</th>
<th>pl</th>
<th>Properties/BLASTP matches</th>
</tr>
</thead>
<tbody>
<tr>
<td>9</td>
<td>602 (641)</td>
<td>61412 (67678)</td>
<td>4.33 (4.98)</td>
<td>Homology with Pseudomonas syringae HrpK (properties indicated in parentheses) (338 residues; E = 1 × e⁻²⁹)</td>
</tr>
<tr>
<td>10</td>
<td>161</td>
<td>17852</td>
<td>7.10</td>
<td>Some homology with acetyltransferases</td>
</tr>
<tr>
<td>11</td>
<td>1287</td>
<td>136278</td>
<td>9.34</td>
<td>Actin-binding domain signature; glycine rich; ATP/GTP-binding site motif A (P-loop)</td>
</tr>
<tr>
<td>12</td>
<td>151</td>
<td>16847</td>
<td>4.18</td>
<td>Homology with Xanthomonas oryzae pv. oryzae HpaB (151 residues; E = 1 × e⁻³⁶)</td>
</tr>
<tr>
<td>13</td>
<td>371</td>
<td>40079</td>
<td>10.40</td>
<td>Homology with Serratia entomophila SepC (327 residues; E = 2 × e⁻²⁷)</td>
</tr>
</tbody>
</table>
Xanthomonas spp. TTS system. These proteins are small and acidic, properties suggesting that they may function as chaperones [12]. Further downstream lies Orf13, which has homology to SepC, a potential insect toxin of Serratia entomophila [28].

**Dot-blot and PCR assays**

Table 1 summarises the data obtained by dot-blot and PCR assays to determine the presence of different TTS genes in various strains of *B. pseudomallei* and *B. thailandensis*. Five TTS1 loci were targeted by dot-blot and PCR assays. Only dot-blot data are presented for two of the loci because PCR amplification with the primers PM84/PM95 (for *orf4*/orf5) and PMXBA3/PMBAM (for the gap between *orf1* and *bpscQ*) with the various genomic DNAs proved to be difficult. However, in both cases it was possible to produce amplicons for DIG-labelling to make probes for dot-blot analysis. Fig. 2 shows examples of PCR assay results. In all cases the *B. pseudomallei* strains tested positive for the presence of TTS-related DNA whereas *B. thailandensis* strains were negative. Strain E27, which had previously proved positive for *orf2*/bpscD by both PCR and dot-blot assays [15], was negative for the loci tested in this study. Occasionally, a faint amplicon of the correct size was visible when primers PM98/PM99 (for *bpscJ*) were used to amplify strain E27 genomic DNA. However, dot-blot analysis suggested that this gene is not present in strain E27, or is significantly different from the *bpscJ* genes present in *B. pseudomallei* strains.

**RT-PCR**

The conditions required for expression of TTS1 genes in *B. pseudomallei* were not known. In *Pseudomonas aeruginosa*, the addition of NTA to growth media induces TTS genes [29]. The initial approach in the present study involved culturing the bacteria at 37°C in the presence or absence of NTA. Two primer combinations were used to screen for gene expression from putative genes either side of the repeat sequence region of the gene cluster. The three genes targeted were *bpscI*, *orf10* and *orf11*. The *fliC* gene was targeted as a positive control. In both strains E503 and E27, grown with and without NTA, the *fliC* gene could be amplified from cDNA. All of the controls where reverse-transcriptase was not used were negative, indicating that there was no residual genomic DNA in the cDNA samples (Fig. 3). RT-PCR amplicons for *bpscI*, *orf10* and *orf11* were obtained with strain E503 samples only. This occurred with and without NTA, indicating that NTA is not required for the expression of these genes in *B. pseudomallei*.

**TTS2 and comparisons with *B. mallei***

Comparisons of *B. pseudomallei* E503 TTS1 sequences with those available through the *B. pseudomallei* genome sequence project indicated that there are sequence variations (typically 1–2%). Such comparisons also suggested that there was a second TTS gene cluster in this bacterium. A cluster of genes homologous to those encoding BpscU, BpscV, Orf1, BpscQ, BpscR, BpscS, Orf2, BpscD, Orf3 and Orf4, could be identified on the same 9315-bp sequence contig. A comparison of this cluster (TTS2) with *B. mallei* genome sequence project data indicated the presence of an equivalent gene cluster (99% sequence identity), and further suggested that genes encoding Orf5, BpscJ, Orf6, BpscL and BpscN were also present in the cluster. Further analysis of another *B. pseudomallei* sequence...
contig, overlapping with the *B. mallei* sequence, suggests that the TTS2 gene cluster resembles TTS1 in gene organisation from *bpscT-orf12*, but that an equivalent gene to *orf8* lies downstream of the TTS2 *orf12* and that the equivalent gene to *bpscC* is several kilobases further downstream. Indeed, there are two possible *bpscC* equivalents (*bpscC2a* and *bpscC2b*) separated by a cluster of genes with homology to genes involved in type IV pilus production (Fig. 1). PCR assays were used to screen three *B. pseudomallei* and three *B. thailandensis* strains, and indicated that the TTS2 gene *bpscU2* is present in both *B. pseudomallei* and *B. thailandensis* (data not shown). There was no evidence for the presence of the TTS1 gene cluster in *B. mallei* from *orf11* to *orf12* (at the time the search was conducted the data were estimated to represent at least three-fold coverage of the genome). However, a BLASTX search with Orf13 indicated its presence in *B. mallei* (99% sequence identity). Further sequence analysis confirmed that *orf13* and DNA directly downstream are present in *B. mallei* but upstream of *orf13* the sequence similarity ends (between *orf12* and *orf13*). A PCR assay for *orf13* was conducted on a selection of *B. pseudomallei* and *B. thailandensis* strains. The results indicate that unlike *B. mallei* and *B. pseudomallei*, *B. thailandensis* strains lack *orf13* (data not shown).

Fig. 3. RT-PCR results for detection of expression of *orf10* (a), *bpscJ* (b), *orf11* (c) and *fliC* (d) are shown. The figure comprises Southern blot hybridisation data from four gels. In each case the lanes refer to the following: 1, strain E503 grown at 37°C in the absence of NTA; 3, strain E503 grown at 37°C in the presence of NTA; 5, strain E27 grown at 37°C in the absence of NTA; 7, strain E27 grown at 37°C in the presence of NTA; 2, 4, 6, 8, controls lacking reverse-transcriptase for equivalent reactions to preceding lanes; 9, genomic DNA controls.
Discussion

The TTS1 sequenced region of the B. pseudomallei E503 genome reported in this paper contains numerous putative genes with homology to TTS system genes reported previously and is highly likely to encode such a system. The overall G+C content (63 mol%) is similar to the value reported previously for the region spanning orf1–orf2 [14] and is lower than the value estimated for this organism by the B. pseudomallei sequencing group (65 mol%; ftp://ftp.sanger.ac.uk/pub/pathogens/bps).

However, the difference in G+C content is marginal compared with some reported for TTS PIs in other bacteria [12]. Although the genes encoding TTS systems are generally clustered together, the order of genes varies between bacteria. Whereas proteins involved in the formation of the secretion structure, or secreton, tend to be conserved throughout TTS systems, other proteins are apparent only in related groups of bacteria, or are unique to a particular TTS system. Often genes are conserved in relative location between bacteria whilst exhibiting low sequence homologies [12, 13]. The organisation of secreton genes in the B. pseudomallei TTS1 cluster strongly resembles that seen in the plant pathogens R. solanacearum and Xanthomonas spp. The only significant difference stems from the fact that in Xanthomonas spp., hrpXv, the equivalent of the genes encoding the putative transcriptional regulators HrpB (in R. solanacearum) and Orf8 (B. pseudomallei), is not located between the genes encoding BpscC and BpscT or homologues, but lies in an unlinked position [30].

The regions downstream of bpscC and its equivalents differ significantly between B. pseudomallei TTS1, R. solanacearum and Xanthomonas spp. In R. solanacearum and Xanthomonas spp., this domain marks the boundary between genes encoding proteins with largely structural roles in the formation of a secreton and those that encode the secreted products. Therefore, it would seem reasonable to postulate that ORFs in B. pseudomallei lying in an equivalent location may encode potential type III-secreted proteins or accessory proteins associated with secretion. In B. pseudomallei, the boundary is marked by the presence of a repeated sequence, which may be indicative of a recombination event that has linked the structural genes of the TTS system with genes encoding effector or other proteins specific to B. pseudomallei.

PCR and dot-blot assays in this study confirmed the association between TTS genes and an Ara− phenotype reported previously [15]. The earlier observations that one B. thailandensis strain E27 was carrying some TTS1 system-related DNA were not further substantiated. The results obtained previously for the orf2/ bpscD region, and the reproducible presence of a faint band following PCR amplification with the PM98/PM99 primer set (for bpscJ) cannot simply be explained by the presence of TTS2 in B. thailandensis because none of the primers match the TTS2 sequence sufficiently well. There may be DNA in B. thailandensis strain E27 that is related to B. pseudomallei TTS1 DNA. If this is the case, the equivalent genes in strain E27 are sufficiently different from those in B. pseudomallei to ensure that detection is not possible by dot-blot hybridisation in most cases. The alternative explanation is that strain E27 contains DNA equivalent to some but not all the B. pseudomallei TTS1 gene cluster. The RT-PCR assays used in this study did not lead to the detection of any expression of TTS1 genes in E27. The problems associated with two of the PCR assays in this study may well be related to minor sequence variations between strains, causing primer mismatching.

In many cases, although TTS is triggered in situ by cell contact, growth conditions can induce expression of TTS genes. Secretion of ExoS and other proteins by P. aeruginosa can be induced artificially by growth in the presence of the chelator NTA [29]. The present observations indicate that expression of the B. pseudomallei TTS system is detectable during growth in Luria broth at 37°C. The addition of NTA to growth media was not required for TTS gene expression. The data presented here do not include quantification of gene expression, and there may be growth conditions that lead to higher levels of expression. However, it is clear that there was some expression of TTS genes in nutrient-rich media at 37°C. There are several examples of artificial induction of TTS, including the TTS systems of Yersinia spp. (induced by growth in media containing low levels of calcium ions [31]), Shigella spp. (in which chemical compounds, including Congo red, Evans blue and direct orange, induce secretion of Ipa proteins [32]) and Salmonella enterica serovar Typhimurium (in which SPI2 genes can be induced by Mg^{2+} deprivation and phosphate starvation [33]). However, temperature induction of TTS without the requirement for additional media supplements is not unprecedented. In Bordetella bronchiseptica, TTS is under the control of the BvgAS regulon, and can be induced in response to growth at 37°C (Bvg^+). In contrast, growth at or below 26°C, or in the presence of high concentrations of nicotinic acid or MgSO_4 (Bvg^−), excludes the production of almost all of the known virulence factors in Bordetella spp., including TTS [34].

There is considerable homology between the predicted Orf8 protein and R. solanacearum HrpB, the positive regulator responsible for regulating expression of TTS operons [20, 21], and a member of the AraC family. Putative binding sites for such regulatory proteins are difficult to identify from sequence data. A consensus sequence (TTTTaGYCtTt at, in which nucleotides conserved in ≈60% of sequences are shown in capital letters) has been reported for VirF binding sites in Yersinia spp. [35]. The present study found a sequence conforming to this consensus upstream of the orf8
gene. The presence of a putative VirF-like binding site suggests that, as in *R. solanacearum*, the *B. pseudomallei* TTS gene cluster may encode a transcriptional activator that is autoregulated. In *Xanthomonas* spp. HrpXv, the equivalent AraC-like positive regulatory protein, is not autoregulated [30].

In *R. solanacearum* the *hrp* genes are expressed in minimal media but repressed in nutrient-rich media [36], which suggests that the regulatory system differs considerably from that of *B. pseudomallei* TTS1. The fact that the latter system is induced in nutrient-rich media at 37°C may be indicative of a system designed for activity in mammals rather than in plants. A number of candidate ORFs for secreted proteins have been identified by relative location compared to the TTS PIs of *R. solanacearum* and *Xanthomonas* spp. RT-PCR indicated that orf10 and orf11 are expressed, suggesting that this region does contain functional, expressed genes. There are putative ribosome-binding sites (AGGA) upstream of orf9 and orf10. The end of orf9 is separated from the beginning of orf10 by only 24 bp. Therefore, it seems likely that orf9, orf10 and orf11 form an operon. Although there is a potential ORF within the repeat region, and there are potential ribosome-binding sequences upstream of this ORF (5’-GGAGGCGGAA); without evidence to the contrary it seems unlikely that this does encode a functional protein. Any such protein would contain the same sequence repeated imperfectly three times with short intervening gaps.

Only one of the predicted protein sequences derived from these ORFs was significantly homologous to proteins identified in other TTS systems. Orf9 had homology to the HrpK protein of *P. syringae*, a protein of unknown function. This homology did not include the C-terminal region, which suggests that there may be functional differences between these proteins. Although Orf10 shared some homology with a number of bacterial acetyltransferases, *E* values were much higher than the level generally considered as indicating significance (*E* ∼ 1 × 10⁻³). Orf11 is a large protein, which shares no significant homology to any proteins in the database, but contains a number of potential motifs. There is evidence that *B. pseudomallei* is a facultative intracellular pathogen, capable of multiplying within various cell types [37]. It has been demonstrated that, after internalisation, the organism can escape from membrane-bound phagosomes into the cytoplasm [37–39]. More recently it has been shown that *B. pseudomallei* can induce cell fusion leading to the formation of multinucleated giant cells and actin-associated membrane protrusion in phagocytic and non-phagocytic cell lines, potentially enabling the pathogen to spread between cells [40]. It is tempting to speculate that type III-secreted proteins may be implicated in such interactions with actin. However, neither ActA, SipA nor SipC contain the actin-type actin-binding domain signature sought by the programme MOTIFS, and the significance of the signature identified in Orf11 is difficult to assess without evidence of activity. A number of protein families have the ATP/GTP-binding site motif identified in Orf11. Amongst the most interesting of these are members of the ADP-riboseylation factors family and kinases.

Downstream from orf11, following a gap of 310 bp, there is DNA which when translated in the opposite orientation to orf11 gives a partial predicted protein sequence with strong homology (50% identity/65% similarity over 251 amino acids; BLASTP *E* value of 3 × 10⁻²⁸) to a transposase of *R. metallidurans* (AF236817). Residual transposon or IS sequences are a feature of PIs [12] and the presence of such a sequence strongly suggests that the TTS-associated gene cluster ends at this point, and that a putative transposase may have played a role in the acquisition of this gene cluster by *B. pseudomallei* E503. Interestingly, the high levels of sequence identity between strain E503 TTS1 and the equivalent region of the strain chosen for genome sequencing, K96243, ends between orf11 and the putative transposase gene. Indeed, BLAST searches failed to identify the strain E503 transposase gene in K96243 or *B. mallei*, although there are sequences with homology to other transposases downstream of orf11 in K96243 (transposase of *Tn*1721/*Tn*501, *E* = 3 × 10⁻²², over 177 amino acids). However, the K96243 sequence contains a stop codon within the matching region, suggesting either that the genome data is erroneous or that a mutation has occurred that would render the gene inactive. This observation does suggest that there may be DNA sequence differences between strains at the edge of the TTS1 gene cluster, and that there are variations in transposon or IS sequences between strains of *B. pseudomallei*.

With TTS1 predicted proteins as probes in silico, a second TTS gene cluster (TTS2) was identified in both *B. pseudomallei* and *B. mallei*. Analysis indicates that *B. pseudomallei* and *B. mallei* both carry TTS2 genes whereas only *B. pseudomallei* carries the TTS1 gene cluster. In addition, a PCR assay for detection of *bpscU2* indicated that at least part of TTS2 is present in *B. thailandensis*. Recently, it has become apparent that there is a third TTS system gene cluster (tentatively named TTS3). TTS3 includes genes with homology to some of the *Salmonella* Inv/Spa secreted virulence factors [41] and is present in *B. pseudomallei* and *B. mallei*. A PCR assay to screen for the putative gene encoding a protein with homology to *Salmonella* SopE showed evidence to suggest that TTS3 DNA is also present in *B. thailandensis* (data not shown). Thus, only the TTS1 system defines a major difference between *B. pseudomallei* and both *B. thailandensis* and *B. mallei*. One boundary marking the variation between *B. mallei* and *B. pseudomallei* lies between orf12 and orf13 in TTS1. At this stage, it is not clear whether orf13 should be considered part of the TTS1 gene cluster. Orf13 is homologous to the C-terminal region
of the putative insect toxin SepC. Hurst et al. [28] demonstrated homology between SepC and other insect toxins. However, those regions common to insect toxins were not the same as the region of best homology with Orf13. Whether Orf13 is involved in TTS or not, it may be a gene of interest because of its presence in B. mallei and B. pseudomallei but absence from B. thailandensis. The TTS2 genes are organised in a similar fashion to the TTS1, R. solanacearum and Xanthomonas spp. TTS gene clusters suggesting common ancestry for all these systems. Although the B. pseudomallei TTS1 and TTS2 genes are most similar to each other, there has been considerable divergence suggesting that the two TTS systems may have different functions. Although the actual role of TTS during infection with B. pseudomallei has yet to be resolved, the likelihood is that, as in all other cases, TTS makes an important contribution to the pathogenicity of this bacterium.

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