Different patterns of integral membrane protein localization during cell division in *Bacillus subtilis*

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INTRODUCTION

Cell division in rod-shaped bacteria is an intensively studied process involving the coordinated action of a suite of proteins that assemble at the mid-cell (Errington et al., 2003). Although not all cell-division proteins are conserved, the tubulin homologue FtsZ is present in virtually all bacteria, some chloroplasts and mitochondria from ancient eukaryotes (Gilson & Beech, 2001). FtsZ self-polymerizes to form a ring at mid-cell, driving the invagination of membrane and cell wall throughout cell division (Errington et al., 2003). It is also the first known protein to assemble at new division sites (Errington et al., 2003) and so represents an excellent early marker for the location of cell division. Although it is a cytoplasmic protein, FtsZ assembles as a polymer adjacent to the inside surface of the cytoplasmic membrane in order to lead the coordinated assembly of the division complex comprising a series of cytoplasmic and integral membrane proteins (Errington et al., 2003). FtsZ placement is dependent on the action of the Min and Noc (nucleoid occlusion) systems, which ensure that division occurs at mid-cell and will not bisect a replicating chromosome (Errington et al., 2003; Wu & Errington, 2004; Rothfield et al., 2005).

The mature *Bacillus subtilis* division complex (the divisome) comprises eight different proteins (Errington et al., 2003; Hamoen et al., 2006; Harry et al., 2006) and many thousands of protein molecules. For example, it is estimated that there are around 50,000 molecules of DivIC per cell during exponential growth (Katis et al., 1997). All of these proteins have been shown to localize to the sites of cell division, and are integral membrane proteins, apart from FtsZ and FtsA (Errington et al., 2003; Hamoen et al., 2006; Harry et al., 2006).

The cytoplasmic membrane is also thought to be important in cell division (see Milejkovskaya & Dowhan, 2005; Matsumoto et al., 2006 for reviews). The major phospholipids found in *B. subtilis* are phosphatidylglycerol (PG) and phosphatidylethanolamine (PE), although the minor component cardiolipin (CL) is also considered an important membrane lipid due to its packing and non-bilayer formation properties (de Mendoza et al., 2002; Kawai et al., 2004; Matsumoto et al., 2006). Sites of cell division become enriched in CL and PE; this is thought to be related to their ability to form non-bilayer structures that would be important in septation events (Matsumoto et al., 2006). There is now ample evidence that the mid-cell and polar regions have a different lipid profile to the longitudinal regions of the membrane (Binenbaum et al., 1999; Fishov & Woldringh, 1999; Milejkovskaya & Dowhan, 2000; Kawai et al., 2004; Nishibori et al., 2005) and so proteins with specific lipid preferences may show altered localization during cell division. It is also possible that due to the large amount of division-specific protein at mid-cell, other integral membrane proteins could be physically excluded from the mid-cell region during division due to crowding effects.

We have investigated the localization of integral membrane proteins in *B. subtilis* and found that some, but not all, are transiently diminished/absent from mid-cell around the time of FtsZ-ring assembly. At this stage of the cell cycle,
the amount of known division-specific integral membrane protein present at mid-cell would be low. All the proteins investigated were present at mid-cell during the later stages of division when the level of division-specific proteins was at its highest. The specific reduction of proteins from the mid-cell region was also shown to be dependent on division ring formation and not on other processes such as DNA replication, which is also known to occur around mid-cell (Lemon & Grossman, 1998; Migocki et al., 2004). Consistent with studies on the localization of specific lipids, we interpret our data as suggesting that integral membrane proteins migrate into division septa at different rates dependent on their local lipid environment.

**METHODS**

**Bacterial strains and media.** All cloning was carried out in *Escherichia coli* DH5α (Gibco-BRL). *B. subtilis* strains used and constructed in this work are listed in Table 1. Transformation of *B. subtilis* was carried out by the method of Anagnostopoulos & Spizizen (1961), as modified by Jenkinson (1983). Transformants constructed in this work are listed in Table 1. Transformation of *B. subtilis* was performed by growing cells grown at 37°C in CH medium as described previously (Davies et al., 2005). FtsZ-depletion experiments were performed by growing cells to control expression of genes, xylose and IPTG were used at 0.5% (w/v) and 0.5 mM, respectively, unless otherwise stated. For DNA manipulations, *p16.7* was amplified from *B. subtilis* chromosomal DNA (Table 1) using the primers Forward 5’-AACAGCCTGG-CCTAGGATGGAAGCTTATTG-GATG-3’ and Reverse 5’-CAACA-GCCTTTTTCAATTGACCCCGGATATG-3’. The resulting product was digested with *AvrII* and *HindIII* (sites underlined in primer sequences) and inserted into similarly cut pSG1192 to generate pNG510 (Table 1). *secDF* was amplified from strain 168 chromosomal DNA using the primers Forward 5’-GAGTACTCGATGG-AAAAAGACG-3’ and Reverse 5’-GCCTGATATGAATTCTTGCGCCGA-3’. The resulting product was digested with *XhoI* and *EcoRI* and inserted into similarly cut pSG1729 to give pNG534 (Table 1). The integrity of inserts was confirmed by DNA sequencing. These plasmids were used to transform competent *B. subtilis* strains as listed in Table 1.

**Microscopy, image acquisition and analysis.** Microscopy was performed as detailed previously (Davies et al., 2005) by mounting cells on 1.2% (w/v) agarose pads, followed by imaging with a Zeiss Axioscop 2 fluorescence microscope fitted with a Photometrics Quantix 1401E cooled CCD camera. Cyan fluorescent protein (CFP) fluorescence was visualized with filter set 3104x4v2, green fluorescent protein (GFP) with set 41018, and yellow fluorescent protein (YFP) with set 41029 (Chroma Technology). Image processing and analysis was performed as described by Johnson et al. (2004) using MetaMorph v6.1.02 software (Molecular Devices). When performing linescans, a line was drawn right around the cell periphery starting at

### Table 1. Strains and plasmids used in this study

<table>
<thead>
<tr>
<th>Strain/plasmid</th>
<th>Genotype</th>
<th>Source/construction</th>
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<tbody>
<tr>
<td>168</td>
<td>trpC2</td>
<td>Laboratory stock</td>
</tr>
<tr>
<td>BB11</td>
<td>trpC2 Chr::pJSIZApble (Ppuc-ftsZ ble)</td>
<td>Beall &amp; Lutkenhaus (1991)</td>
</tr>
<tr>
<td>BS125</td>
<td>trpC2 spoA3 su amyE (spc Pspc-P16.7-gfp)</td>
<td>Meijer et al. (2001)</td>
</tr>
<tr>
<td>BS133</td>
<td>trpC2::pNG25 (atpA-cfp Pspc-atpA OpCm:Sp (cat&lt;sup&gt;+&lt;/sup&gt; spe&lt;sup&gt;+&lt;/sup&gt;) chr::pNG115 (sdhA-yfp Pspc-sdhA cat)</td>
<td>Johnson et al. (2004)</td>
</tr>
<tr>
<td>BS351</td>
<td>trpC2 Chr::pJSIZApble (Ppuc-ftsZ ble) amyE (spc Pspc-fisZ-cfp) chr::pNG115 (sdhA-yfp Pspc-sdhA cat)</td>
<td>This work, BS1059 transformed with BS116</td>
</tr>
<tr>
<td>BS352</td>
<td>trpC2 Chr::pJSIZApble (Ppuc-ftsZ ble) amyE (spc Pspc-fisZ-yfp) chr::pNG25 (atpA-cfp Pspc-atpA cat)</td>
<td>This work, BS1060 transformed with BS24</td>
</tr>
<tr>
<td>BS426</td>
<td>trpC2 Chr::pJSIZApble (Ppuc-ftsZ ble) chr::pNG25 [atpA-cfp Pspc-atpA OpCm:Sp (cat&lt;sup&gt;+&lt;/sup&gt; spe&lt;sup&gt;+&lt;/sup&gt;) chr::pNG115 (sdhA-yfp Pspc-sdhA cat)</td>
<td>This work, BS133 transformed with BB11 DNA</td>
</tr>
<tr>
<td>BS441</td>
<td>trpC2 Chr::pNG115 (sdhA-yfp Pspc-sdhA cat) amyE (spc Pspc-P16.7-cfp)</td>
<td>This work, BS116 transformed with pNG510</td>
</tr>
<tr>
<td>BS451</td>
<td>trpC2 amyE (spc Pspc-gfp-secDF)</td>
<td>This work, 168 transformed with pNG534</td>
</tr>
<tr>
<td>BS462</td>
<td>trpC2 amyE (spc Pspc-gfp-secDF) chr::pNG25 (atpA-cfp Pspc-atpA cat)</td>
<td>This work, BS24 transformed with pNG534</td>
</tr>
<tr>
<td>BS1059</td>
<td>trpC2 Chr::pJSIZApble (Ppuc-ftsZ ble) amyE (spc Pspc-fisZ-cfp)</td>
<td>Feucht &amp; Lewis (2001)</td>
</tr>
<tr>
<td>BS1060</td>
<td>trpC2 Chr::pJSIZApble (Ppuc-ftsZ ble) amyE (spc Pspc-fisZ-yfp)</td>
<td>Feucht &amp; Lewis (2001)</td>
</tr>
<tr>
<td>pSG1729</td>
<td>bla amyE' spc Pspc-P16.7-cfp amyE'</td>
<td>This work, PCR p16.7 from BS125 and insert into pSG1192</td>
</tr>
<tr>
<td>pNG530</td>
<td>bla amyE' spc Pspc-gfp-secDF amyE'</td>
<td>This work, secDF inserted into pSG1729</td>
</tr>
</tbody>
</table>
one of the cell poles. In the resulting intensity profile, the mid-cell sites would therefore correspond approximately to the one-quarter and three-quarter points on the x-axes. Final images were prepared for publication using Adobe Photoshop.

RESULTS

Colocalization of integral membrane proteins

In previous studies we investigated the distribution of succinate dehydrogenase and ATP synthase complexes using fluorescent protein fusions to the AtpA and SdhA subunits, respectively, which showed that both complexes were distributed around the cytoplasmic membrane (Johnson et al., 2004). The expression of both fluorescent fusions was driven by the activity of the wild-type promoters, and polar effects of insertion of the fusions into the chromosome were avoided through expression of downstream genes from the xylose-inducible promoter (see Johnson et al., 2004 and Fig. 1A). The pattern of localization for both SdhA and AtpA fusions was heterogeneous, with multiple small regions of fluorescence of varying levels of intensity (see Johnson et al., 2004). Due to this highly varied level of fluorescence around the cytoplasmic membrane, no obvious pattern of low signal at mid-cell was apparent in either of the single-labelled strains. However, we subsequently noticed that a proportion of the cells in the atpA-cfp sdhA-yfp dual-labelled strain BS133 (Table 1) showed much lower levels of signal at the mid-cell site (Fig. 1B). As is clear in the linescan of the cell perimeter shown in Fig. 1(D), the distribution of each fusion was heterogeneous as marked by a series of peaks and troughs, but at the mid-cell region (light and dark grey arrows, Figs 1C, D) there was a coincident drop in signal intensity for both fusions. These areas, corresponding to simultaneous dips in the signal of dual fluorescent labels at mid-cell, were subsequently termed cleared regions (CRs) and were found to be present in 9.4 % of exponentially growing cells (Table 2). It should be noted that whilst there are many peaks and troughs for the ATP synthase and succinate dehydrogenase signals, the coincident drops in signal representing CRs only occurred at mid-cell (see arrowed cells, Fig. 1B, C).

In order to determine how widespread CRs were, we examined the localization of some additional integral membrane proteins. Strain BS441 contains a xylose-inducible p16.7-cfp fusion along with the sdhA-yfp fusion (Fig. 2A; Table 1). p16.7 is an integral membrane protein from the B. subtilis phage φ29 that is involved in phage DNA replication, and has previously been shown to localize around the cell membrane when ectopically expressed (Meijer et al., 2001; Johnson et al., 2004). This protein plays no role in the metabolism of uninfected B. subtilis and so was used to determine if the appearance of CRs was a non-specific effect on integral membrane proteins at the onset of cell division. Strain BS462 containing an N-terminal gfp fusion to secDF along with the atpA-cfp fusion was also constructed (Fig. 2E; Table 1). SecDF, as part of the Sec secretion complex, is involved in protein secretion (Dalbey & Chen, 2004) and so is not directly associated with the same metabolic activities as ATP synthase and succinate dehydrogenase. Therefore, this strain would inform us whether CRs were a phenomenon that occurred with all B. subtilis integral membrane proteins, or just a subset.

We were unable to obtain a fluorescent YFP fusion to SecDF and so the dual-labelled BS462 strain contained a CFP and GFP fusion. The level of spectral crossover of GFP signals into CFP channels and vice versa was determined with single-labelled strains BS24 and BS451 (Table 1) and shown to be insignificant (not shown). Previous work has

![Fig. 1. Colocalization of the integral membrane proteins ATP synthase and succinate dehydrogenase. (A) Schematic of the chromosomal construction of strain BS133, carrying an atpA-cfp and sdhA-yfp fusion, that was produced by Johnson et al. (2004). Transcription of each of the fluorescent protein fusions was driven by the wild-type atp or sdh promoters, whilst expression of genes downstream from the sites of insertion was controlled by the xylose-inducible promoter P_xyl (B, C) Overlays of AtpA-CFP (red) and SdhA-YFP (green) fluorescent signals in cells of strain BS133 growing exponentially in CH medium. The white arrow in (B) indicates a cell in which there is a well-defined mid-cell region of low red and green fluorescent signals. The grey arrows indicate the mid-cell regions in a chain of cells where there is a lowering of fluorescent signal at mid-cell, but to a lesser degree than for the cell indicated with the white arrow. A linescan of the arrowed cell in (C) is displayed in (D). The light and dark grey arrows on the micrograph indicate regions in the linescan where there is a reciprocal drop in fluorescent signal from both fusions at the mid-cell point. Scale bars, 2 µm.](image-url)
also shown that CFP and GFP fusions can be unambiguously localized with little or no signal crossover (Davies & Lewis, 2003) and so strain BS462 was used to colocalize SecDF and ATP synthase.

**Table 2.** Frequency of mid-cell clearing in strains carrying fluorescent protein fusions to integral membrane protein complexes

<table>
<thead>
<tr>
<th>Time (min)*</th>
<th>BS133</th>
<th>BS426</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean cell length (μm)</td>
<td>Frequency of CRs (%)†</td>
</tr>
<tr>
<td>0</td>
<td>3.3</td>
<td>9.4</td>
</tr>
<tr>
<td>45</td>
<td>6.0</td>
<td>1.9</td>
</tr>
<tr>
<td>90</td>
<td>9.1</td>
<td>0</td>
</tr>
<tr>
<td>150</td>
<td>17.2</td>
<td>0</td>
</tr>
<tr>
<td>180</td>
<td>23.3</td>
<td>0</td>
</tr>
</tbody>
</table>

*Number of minutes following removal of IPTG from the medium (for strain BS426 only).
†CRs, ‘cleared regions’, corresponding to coincident drops in fluorescence on opposite sides at regular intervals along the length of the cell. At least 250 cells were counted for each sample.

Mid-cell clearing of membrane proteins

Fig. 2(B, C, D) shows the results of examination of strain BS441. Surprisingly, there was no evidence that p16.7 formed CRs, and instead a p16.7 signal was seen to concentrate at division sites concomitant with a drop in the SdhA-YFP signal (arrows, Fig. 2B, D). At early stages of cell division around the time that septal invagination was just beginning, p16.7-CFP signals appeared as slight concentrations of signal at mid-cell (red arrows, Fig. 2B, C), that became more intense as the divisome matured (blue arrows, Fig. 2B, D).

Similarly, no evidence for CRs could be found with strain BS462, and as with p16.7, SecDF appeared to localize to septal regions/membrane ingrowths at an earlier stage of cell division than ATP synthase (Fig. 2F, arrows). Thus, the phenomenon of CRs appears to be restricted to a subset of integral membrane proteins currently comprising the ATP synthase and succinate dehydrogenase complexes.

**Cleared regions occur early in the division cycle**

The occurrence of CRs only at mid-cell sites indicated that they could be connected to the cell division cycle. In order to investigate the role of cell division in the clearing of integral membrane proteins from the mid-cell site, further strains were constructed to contain dual fluorescent protein fusions to either *ftsZ* and *sdhA* (BS351; Table 1, Fig. 3A), or *ftsZ* and *atpA* (BS352; Table 1). In these strains expression of wild-type *FtsZ* was dependent on the

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presence of IPTG, whereas expression of fluorescently tagged FtsZ was dependent on the presence of xylose. Production of wild-type FtsZ was necessary in these strains as the fluorescently tagged protein is not fully functional (Feucht & Lewis, 2001). The strains also contained a fusion to either *sdhA* or *atpA*, both of which lie within operons and require the presence of xylose for the expression of genes downstream from the sites of insertion, and the FtsZ-CPF fusion was controlled by the xylose-inducible promoter P<sub>xyl</sub>. Expression of wild-type FtsZ was under the control of the IPTG-inducible P<sub>spac</sub> promoter. In the left-hand micrograph in (B), SdhA-YFP fluorescence has been pseudocoloured green and the FtsZ-CFP signal, red. The FtsZ-CFP signal is shown in the middle micrograph, and the SdhA-YFP signal in the right-hand micrograph. The white arrow indicates a cell containing a coincident Z-ring and CR, whilst the grey arrow indicates a cell containing a Z-ring, but no CR. Scale bar, 2 μm. (C) Linescan around the perimeter of the white-arrowed cell, (D) linescan of the grey-arrowed cell from (B); the black arrows correspond to the mid-cell sites (and FtsZ peaks). (E) Bar chart illustrating the distribution of cells at different stages of the division cycle and the correlation of CRs with the morphology of FtsZ division rings for strain BS351. The cartoons in (E) indicate the appearance of FtsZ rings at different stages of the division cycle; mean cell sizes are shown above each cartoon. Classes of cells are as follows: without, no FtsZ ring apparent; early, the FtsZ ring appears as two dots on opposite sides of the mid-cell representing the early stages of cell division when other division proteins are still being recruited to the mid-cell site; ring, a mature FtsZ ring is present which appears as a band across the middle of the cell and represents a cell undergoing initial stages of division; dot, represents the appearance of the FtsZ ring at later stages of the division cycle when invagination of the cell wall is clearly visible by phase-contrast microscopy. Black columns represent cells in which no CR could be seen. Grey columns represent cells in which CRs could be observed. (F) Western blot of whole-cell extracts immunoblotted with anti-FtsZ antibody. Equivalent amounts of protein were loaded in each lane. WT corresponds to strain 168 (wild-type), 351 and 352 to the dual-labelled strains containing FtsZ fluorescent protein fusions (BS351 and BS352) grown under the conditions detailed in the text.

Cultures of strains BS351 (*sdhA*, *ftsZ* fusions; Table 1) and BS352 (*atpA*, *ftsZ* fusions; Table 1) were grown and examined microscopically. An image of a chain of cells from strain BS351 along with linescans around the perimeter of the marked cells is shown in Fig. 3(B, C, D). For both strains, regions of low membrane signal corresponded with FtsZ division rings (white arrow Fig. 3B, and black arrows Fig. 3C). However, CRs were not present in all cells with visible FtsZ rings (grey arrow Fig. 3B, and black arrows Fig. 3D). In these cells, the SdhA signal can be

![Fig. 3. Protein-free mid-cell regions correspond to the early stages of cell division. (A) Schematic of the chromosomal construction of strain BS351, carrying an *sdhA*-yfp and *ftsZ*-cpf fusion. Transcription of the *sdhA*-yfp fusion was driven by the wild-type *sdh* promoter, whilst expression of genes downstream from the sites of insertion, and the *ftsZ*-cpf fusion was controlled by the xylose-inducible promoter P<sub>xyl</sub>. Expression of wild-type FtsZ was under the control of the IPTG-inducible P<sub>spac</sub> promoter. In the left-hand micrograph in (B), SdhA-YFP fluorescence has been pseudocoloured green and the FtsZ-CFP signal, red. The FtsZ-CFP signal is shown in the middle micrograph, and the SdhA-YFP signal in the right-hand micrograph. The white arrow indicates a cell containing a coincident Z-ring and CR, whilst the grey arrow indicates a cell containing a Z-ring, but no CR. Scale bar, 2 μm. (C) Linescan around the perimeter of the white-arrowed cell, (D) linescan of the grey-arrowed cell from (B); the black arrows correspond to the mid-cell sites (and FtsZ peaks). (E) Bar chart illustrating the distribution of cells at different stages of the division cycle and the correlation of CRs with the morphology of FtsZ division rings for strain BS351. The cartoons in (E) indicate the appearance of FtsZ rings at different stages of the division cycle; mean cell sizes are shown above each cartoon. Classes of cells are as follows: without, no FtsZ ring apparent; early, the FtsZ ring appears as two dots on opposite sides of the mid-cell representing the early stages of cell division when other division proteins are still being recruited to the mid-cell site; ring, a mature FtsZ ring is present which appears as a band across the middle of the cell and represents a cell undergoing initial stages of division; dot, represents the appearance of the FtsZ ring at later stages of the division cycle when invagination of the cell wall is clearly visible by phase-contrast microscopy. Black columns represent cells in which no CR could be seen. Grey columns represent cells in which CRs could be observed. (F) Western blot of whole-cell extracts immunoblotted with anti-FtsZ antibody. Equivalent amounts of protein were loaded in each lane. WT corresponds to strain 168 (wild-type), 351 and 352 to the dual-labelled strains containing FtsZ fluorescent protein fusions (BS351 and BS352) grown under the conditions detailed in the text.](https://www.microbiologyresearch.org)
clearly seen to overlap the FtsZ signal at mid-cell. These results suggested that CRs may occur at a specific stage of the division cycle and so a detailed analysis of CR appearance with FtsZ rings and cell length was conducted (Fig. 3E). Cells were classified as having no detectable FtsZ ring (Fig. 3E) (‘without’, mean cell length 3.12 ± 0.52 μm), an early-stage division ring that appeared as two dots on opposite sides of mid-cell (morphologies first characterized for DivIC by Katis et al. (1997); ‘early’, 3.59 ± 0.49 μm), a mature division ring (‘ring’, 4.70 ± 0.74 μm), or to be in the late stages of cell division (‘dot’, 6.12 ± 0.42 μm). Both the sdhA ftsZ and atpA ftsZ dual-labelled strains had similar proportions of each class, and similar proportions of cells in which CRs were observed and so results are only shown for the sdhA ftsZ strain BS351 in Fig. 3(E). The majority of cells containing CRs were in the early stage of cell division (Fig. 3E, grey columns) indicating that formation of CRs occurred prior to the full assembly of the division complex, which occurs at the ‘ring’ stage of the division cycle. This effect was not observed in all the cells. It may be a specific response to a very short-lived event in the division cycle not fully resolved in our analysis, or the lowering of mid-cell signal may have been much less in some cells than others. Repeated attempts were made to perform time-lapse imaging on strains BS351 and BS352 to definitively establish the timing of CR formation with cell division, but due to the rapid photobleaching of the FtsZ fusion we were unable to obtain usable data.

The division complex represents a significant number of protein molecules and it is possible to envisage a situation whereby the assembly of a large number of proteins involved in synthesis of the division septum might exclude other integral membrane proteins, so creating the lowering of signal we observed. However, since CRs occurred at an earlier stage of the division cycle than the full assembly of the division complex, and were not observed for all integral membrane protein fusions, there must be another explanation.

**Mid-cell clearing is not due to DNA replication or chromosome segregation cycles**

Due to the overlapping nature of the bacterial cell cycle, DNA replication, chromosome segregation and cell division may all be ongoing in the same cell simultaneously. DNA replication occurs in replication factories positioned approximately around the mid-cell point (Lemon & Grossman, 1998; Migocki et al., 2004), and it was possible that CRs coincided with some aspect of the replication or segregation cycles. To investigate this possibility, strain BS426 was constructed (Fig. 4A; Table 1), in which the expression of ftsZ was dependent on induction with IPTG. Thus, by removing IPTG from the medium, FtsZ levels in the cells would drop, resulting in the formation of filaments. In filamentous cells, it is well established that DNA replication and segregation occur normally until the cells grow too large and lyse (e.g. see Fig. 5 in Wu et al., 1995; Lewis et al., 2000). So, if mid-cell clearing was dependent on aspects of the replication and segregation cycles independent of Z-ring formation, CRs should still appear at regular intervals along filaments in this strain.

Strain BS426 was grown in CH medium containing 0.5 mM IPTG to OD600 0.3 prior to resuspension in IPTG-free medium (see Methods). Cells were then collected for analysis at various time points following resuspension, and the frequency of CRs and cell length measurements were determined. The results were compared against the data for strain BS133, which is the
parent strain of BS426 with wild-type ftsZ expression (Table 2). Immediately following resuspension ($t_0$) cells of strain BS426 had a mean cell length of 4.3 μm which is slightly longer than that of the parent strain BS133 (mean 3.3 μm). Titration of FtsZ levels in strain BS426 was attempted to achieve a mean cell length closer to that of BS133, but increasing or decreasing IPTG levels (0.1–1 mM) did not result in improved mean cell lengths, and the most homogeneous cell-length distributions were obtained using 0.5 mM (not shown).

The frequency of CRs was slightly lower in BS426 compared to BS133 (63% and 94%, respectively; Table 2). In addition, the mean cell length for cells containing CRs was determined as well as the relative position of the CR along the long axis of the cell, where a measurement of 0.5 indicates the mid-point. The mean length at which CRs appeared in BS133 was 4.2 ± 0.49 μm with a relative position of 0.52 ± 0.03, and in BS426 was 5.6 ± 0.9 μm with a relative position of 0.51 ± 0.04. Therefore, although cells of strain BS426 were slightly longer than the parent strain, CRs still formed in the same relative position, indicating that the mechanism affecting this phenomenon was operating normally in these cells.

Following resuspension in IPTG-free medium, strain BS426 continued to grow as indicated by the exponential increase in mean cell length over the 180 min time-course when cells increased in mean length from 4.3 to 23.3 μm (Table 2). As the mean cell length increased, the frequency of CRs (classified for these cells as concomitant drops in fluorescent signals at regular intervals along filaments rather than a single region at mid-cell) decreased so that by 90 min (mean cell length 9.1 μm) no more could be detected at mid-cell or any other position along the cell length, indicating the dependence of CR formation on ftsZ expression and that this phenomenon was not dependent on DNA replication or segregation.

**DISCUSSION**

In this work we have shown that certain integral membrane proteins are not observed at the mid-cell membrane region during the early stages of cell division. This process is not ubiquitous, but restricted to a subset of proteins, currently comprising the ATP synthase and succinate dehydrogenase complexes. Additional proteins tested were not absent from mid-cell at any stage of the cell cycle. The phage protein p16.7 is small (16.7 kDa) and so the presence of this protein at mid-cell could be attributed to a higher diffusion rate within the membrane compared to the larger ATP synthase and succinate dehydrogenase complexes. Also, since p16.7 is not a component of the normal cellular proteome, it may not respond to cytoplasmic signals in the same way as normal cellular proteins. However, no evidence of CRs was observed using the SecDF fluorescent fusion. SecDF is part of the normal proteome and is a component of the SecYEG/SecDFYajC (YrbF in *B. subtilis*) protein secretion complex (Dalbey & Chen, 2004). Thus, it does not appear that size is the criterion for temporary absence of certain proteins at mid-cell during cell division. Nevertheless, it was observed that septa containing GFP-SecDF were visible before the AtpA-CFP fusion (see arrows, Fig. 2F), which may indicate that although both complexes are large, the ATP synthase complex diffuses more slowly than SecYEG/SecDFYajC.

We also investigated the possibility that CRs may be due to some aspect of DNA metabolism, as DNA replication has also been shown to occur around the mid-cell region. However, in strains depleted of FtsZ, in which cells became filamentous but underwent normal DNA replication and segregation, CR frequency rapidly dropped to zero, indicating that CRs were dependent in some way on cell division.

Quantitative analysis showed that CRs only occurred at the early stages of the division cycle, corresponding approximately to the time at which FtsZ rings first assemble. It had seemed possible that the physical crowding by division-specific proteins could be responsible for a passive exclusion of other proteins from the mid-cell. Two lines of evidence suggest this is not the case. Firstly, during the later stages of cell division when the level of division proteins is highest at mid-cell, CRs are not observed. They are only seen at the early stages of division when the mature divisome is still being assembled. Secondly, CRs were not observed with all integral membrane proteins suggesting they may represent a specific response by a subset of proteins.

It was also possible that the lipid composition of the membrane plays a role in CR formation, and there does appear to be a connection between local changes in lipid composition and cell division (Mileykovskaya & Dowhan, 2005). A number of studies in both *E. coli* and *B. subtilis* using membrane- and CL-specific dyes indicate that mid-cell and polar regions have a different staining profile to cell-cylinder membrane, and are particularly rich in CL and PE (Fishov & Woldringh, 1999; Mileykovskaya & Dowhan, 2000; Kawai et al., 2004; Nishibori et al., 2005). There is some evidence that ATP synthase is associated with PG domains (Ksenzenko & Brusilow, 1993), which are less abundant at sites of cell division. It could be that due to interaction with different subsets of lipids/lipid domains, some proteins/complexes are more mobile than others in the membrane and can diffuse into newly synthesized membranes before others (e.g. p16.7 and SecDF). Also, the enzymes responsible for synthesis of PE and CL localize to the division septum, where lipid synthesis is thought to occur (Nishibori et al., 2005), and this may inhibit the rate of diffusion of proteins embedded within PG-rich domains into the septal regions.

So, are CRs due to active exclusion or passive diffusion? The former is unlikely, as if that was the case, we would have expected to see them clearly in every cell early during
the division cycle, whereas they were only seen in about 40% of cells at the early stages of cell division. It seems more likely that CRs are due to a slower rate of diffusion of certain complexes into newly synthesized membrane at the onset of cell division. The juxtaposition of membrane domains to centre cell at the onset of division could affect how easily detected a CR is, hence they are only seen in about 40% of cells early in the division cycle. Nevertheless, CRs could be indicative of local changes in the membrane composition at mid-cell, which are important in the formation of an active divisome that may be important in targeting division proteins to that region of the cell.

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