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# The anchoring of the polysaccharide II is essential for Clostridioides difficile survival



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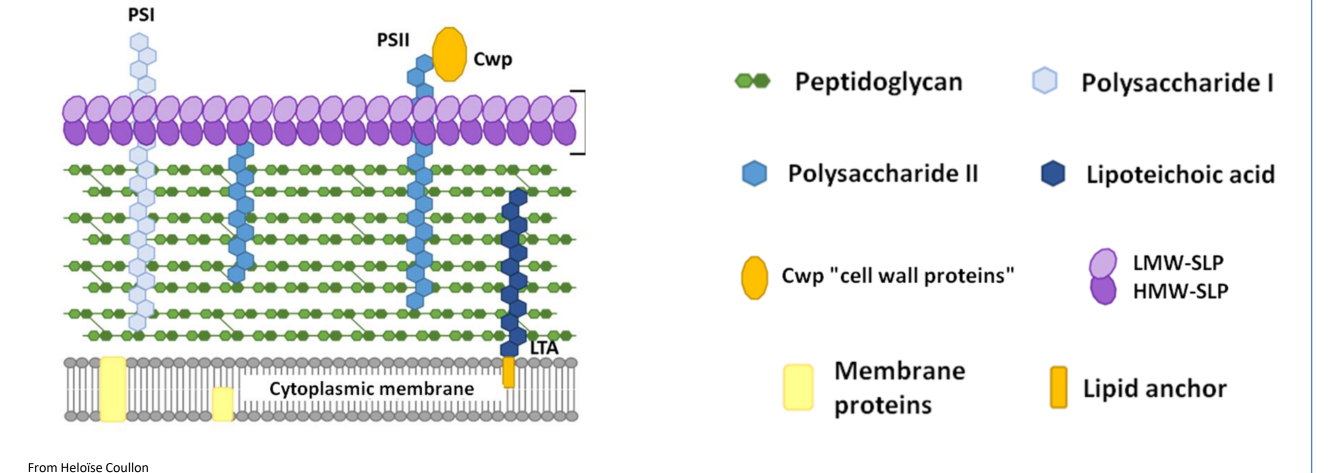
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culture supernatant of the mutant

### INTRODUCTION

C. difficile is an anaerobic, motile and spore-forming bacterium, responsible for 15 to 25% of post-antibiotic diarrhea and 95% of pseudomembranous colitis. While its toxins are described to be the major virulence factors in *C. difficile* infections, there is an increasing interest in the role of non-toxin factors in pathogenesis and virulence. In many other pathogens, cell wall glycopolymers influence the virulence. In C. difficile, three major carbohydrates are described : the polysaccharide I (PSI), the polysaccharide II (PSII) and the lipoteichoic acid (LTA). In the synthesis model suggested before<sup>1</sup>, the polysaccharide II is synthetized in the cytoplasm, then exported to the surface by a flippase and finally anchored into the peptidoglycan by two proteins of the LytR family : LcpA and LcpB. The two *lcp* genes are supposed to be non-redundant and were previously disrupted by insertional inactivation<sup>1</sup>.



## **AIMS OF THE STUDY**

The aims of the study were first to investigate the role of the polysaccharide II and its correct anchoring for growth, surface set-up and survival of C. difficile and also to develop an innovative technique to study essential genes in C. difficile.

# RESULTS

### **CONDITIONAL-LETHAL MUTANT TECHNIQUE TO STUDY ESSENTIAL GENES**

Single and double mutation of the *lcp* genes

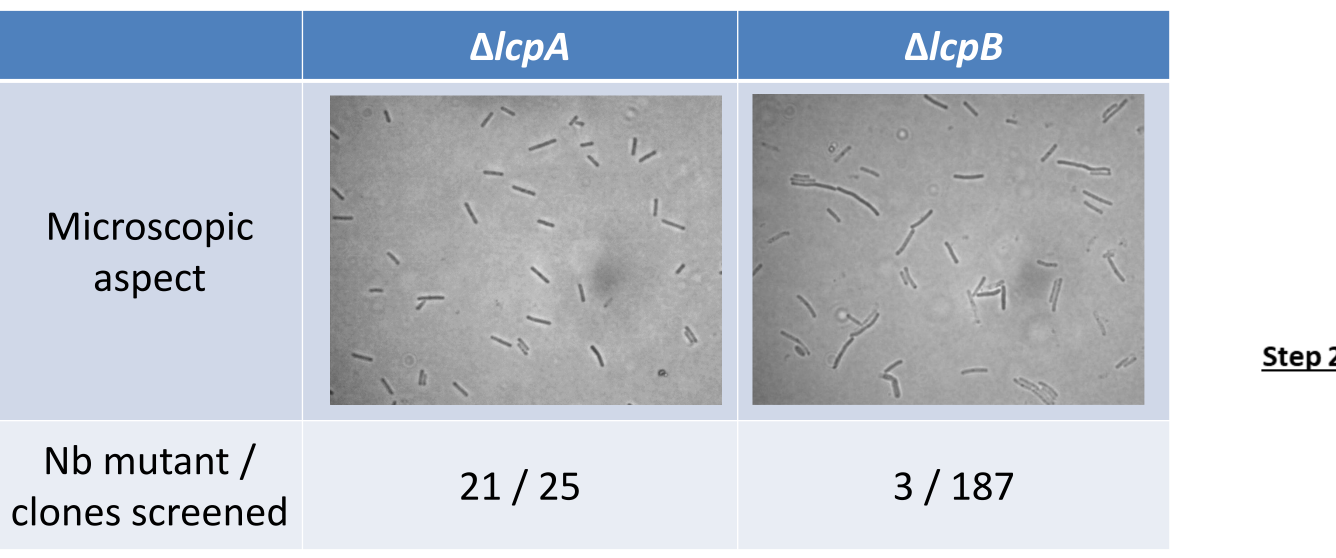
Δ*IcpA* and Δ*IcpB* : constructed by allele-coupled exchange technique

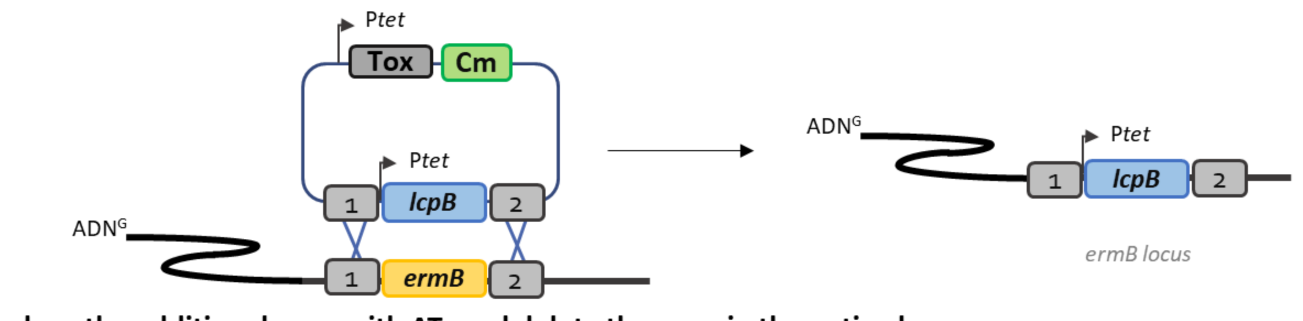
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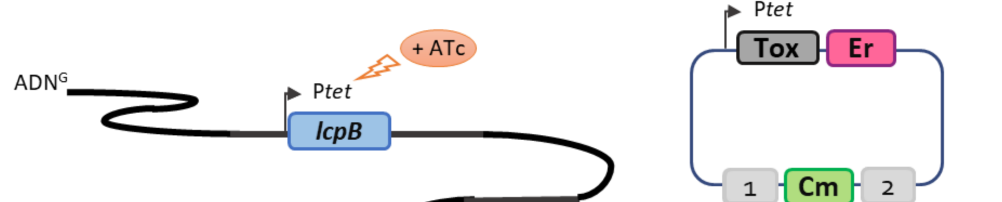


<u>Step 1</u> : insert the supplementary copy into the *ermB* region of the chromosome by ACE and verify by PCR





Step 2 : induce the additional copy with ATc and delete the gene in the native locus

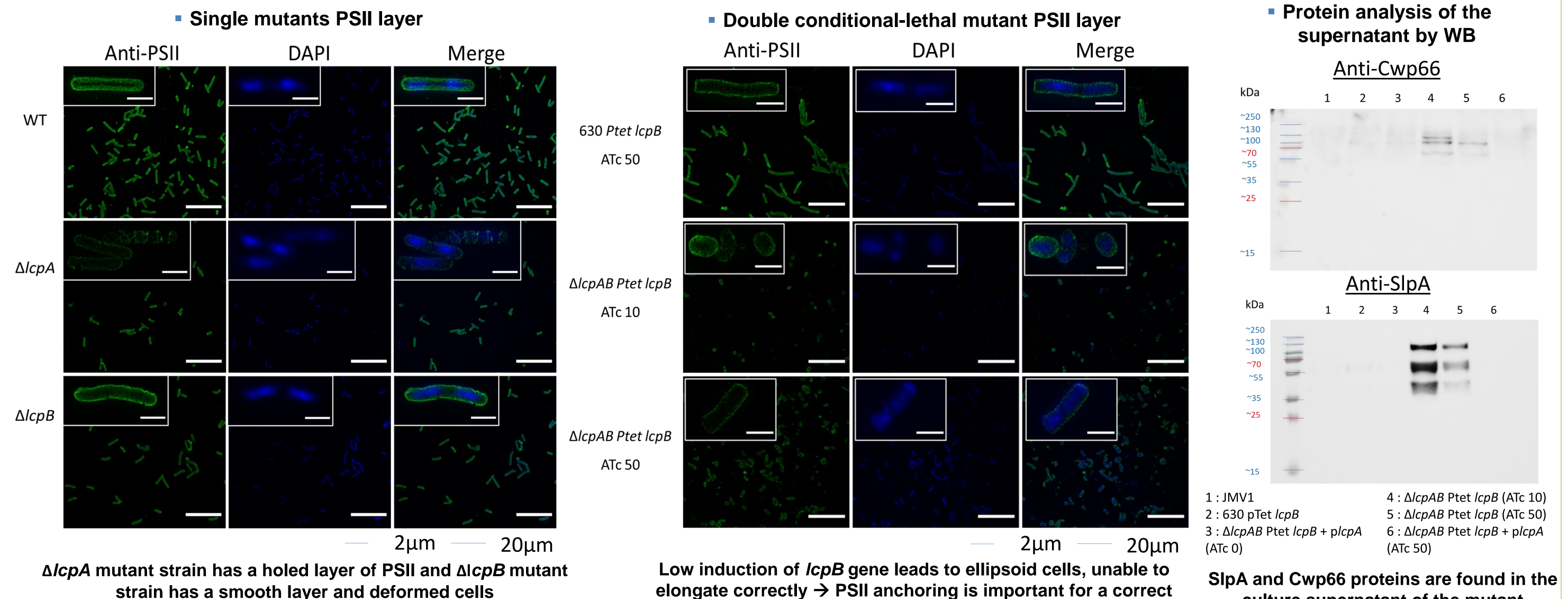


 $\geq \Delta lcpA \Delta lcpB$ : unsuccessful to obtain double mutant Only WT revertants found in PCR over 457 clones screened IcpA IcpB Native locus

 $\rightarrow$  It suggests that LcpA and LcpB have redundant functions and the activity of at least one Lcp seems to be essential for survival

 $\rightarrow$  Modulation of the gene expression to allow survival and observe phenotype  $\rightarrow$  Double *lcp* mutant is not able to grow without ATc  $\rightarrow$  LcpA and LcpB are redundant  $\rightarrow$  PSII anchoring is essential

### MUTANTS ANALYSIS BY IMMUNOFLUORESCENCE MICROSCOPY AND WESTERN BLOT



strain has a smooth layer and deformed cells

# **CONCLUSION - PERSPECTIVES**

bacterial elongation

Thanks to the construction of the single *lcp* mutants by allelic exchange, we confirmed the phenotypes previously observed<sup>1</sup>. The impossibility to isolate a double *lcp* mutant clone tends to confirm the essentiality of the PSII anchoring into the peptidoglycan. This was confirmed using to the conditional-lethal mutant of both *lcp*, which is completely unable to grow without induction of the supplementary copy of *lcpB* added in the chromosome. When *lcpB* expression is very low, we observe ellipsoid cells, unable to elongate. Our results show the critical role of polysaccharide II anchoring in growth, elongation, and correct surface set-up of C. difficile. Our technique provides new opportunities to study essential genes in *C. difficile*.

<sup>1</sup> Chu M *et al*, Plos Pathogens, 2016