

Comparative subproteomic analysis of experimentally induced fluoroquinolone resistance and ciprofloxacin stress in Salmonella Typhimurium DT104B

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Comparative subproteomic analysis of experimentally induced fluoroquinolone resistance and ciprofloxacin stress in Salmonella Typhimurium DT104B

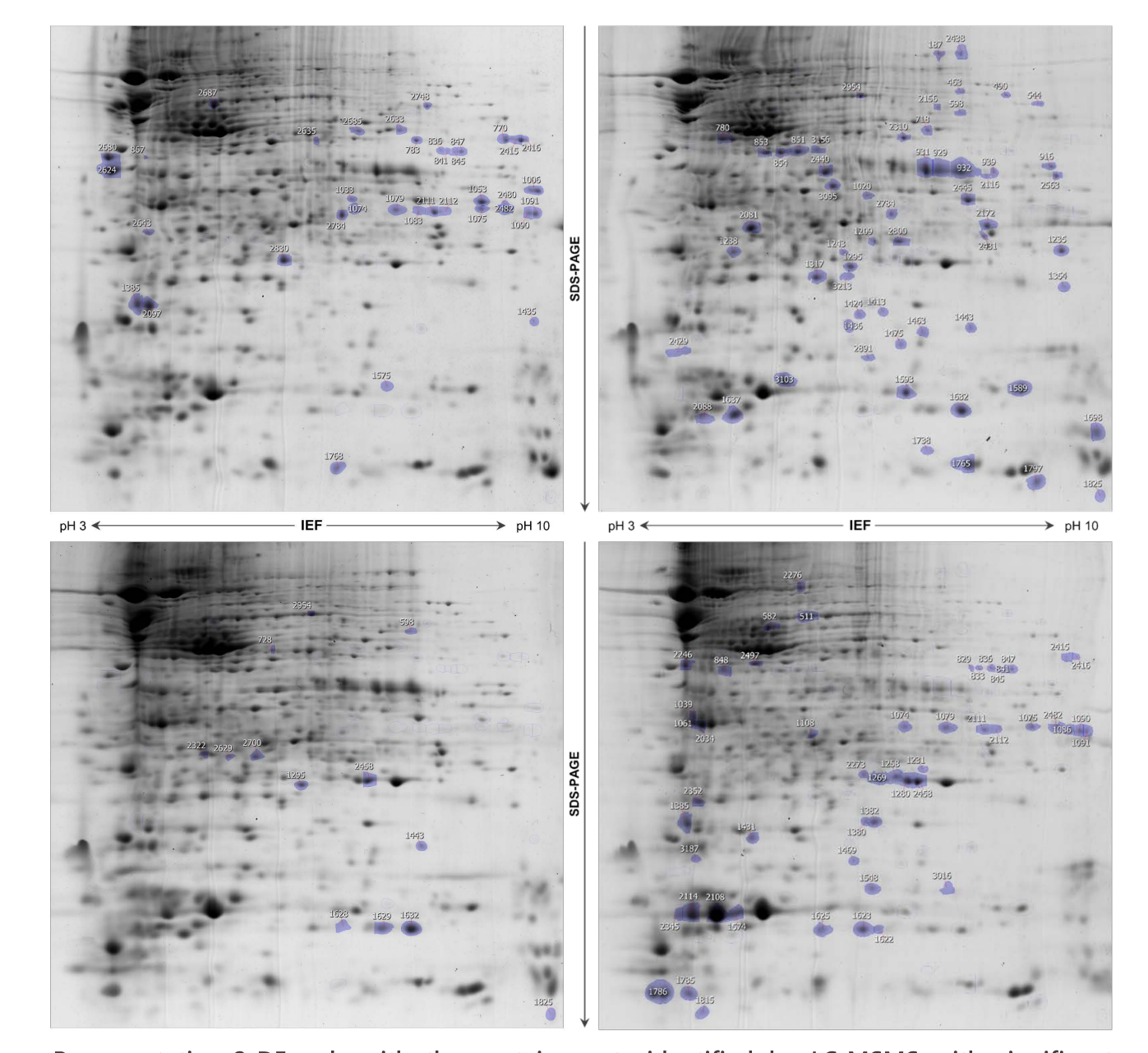
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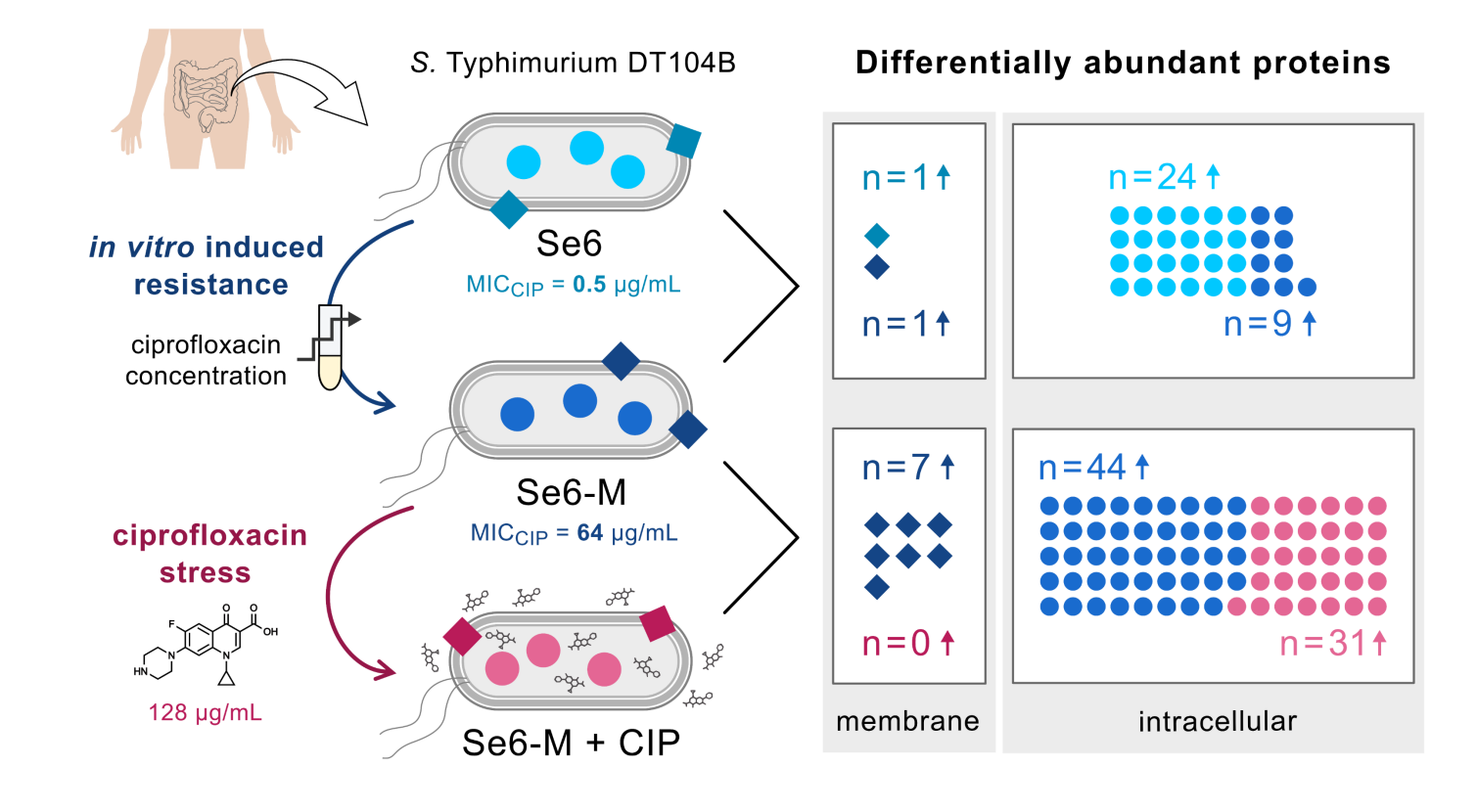
Fluoroquinolone-resistant Salmonella spp. is a global high priority pathogen for research and development of new antibiotics (World Health Organization, 2017) and Salmonella Typhimurium DT104B multiresistant strains with additional quinolone resistance have been responsible for global outbreaks and high mortality. Multiple factors are known to be involved in fluoroquinolone resistance; however, additional determinants continue to be described and the exact contribution of many mechanisms remains unexplained and/or need better understanding.

To give new insights into the mechanisms involved in fluoroquinolone resistance in Salmonella Typhimurium DT104B, this work aimed to evaluate subproteome changes between a laboratory-induced fluoroquinoloneresistant strain (Se6-M) and its parent strain (Se6) and also in Se6-M under ciprofloxacin (CIP) stress.

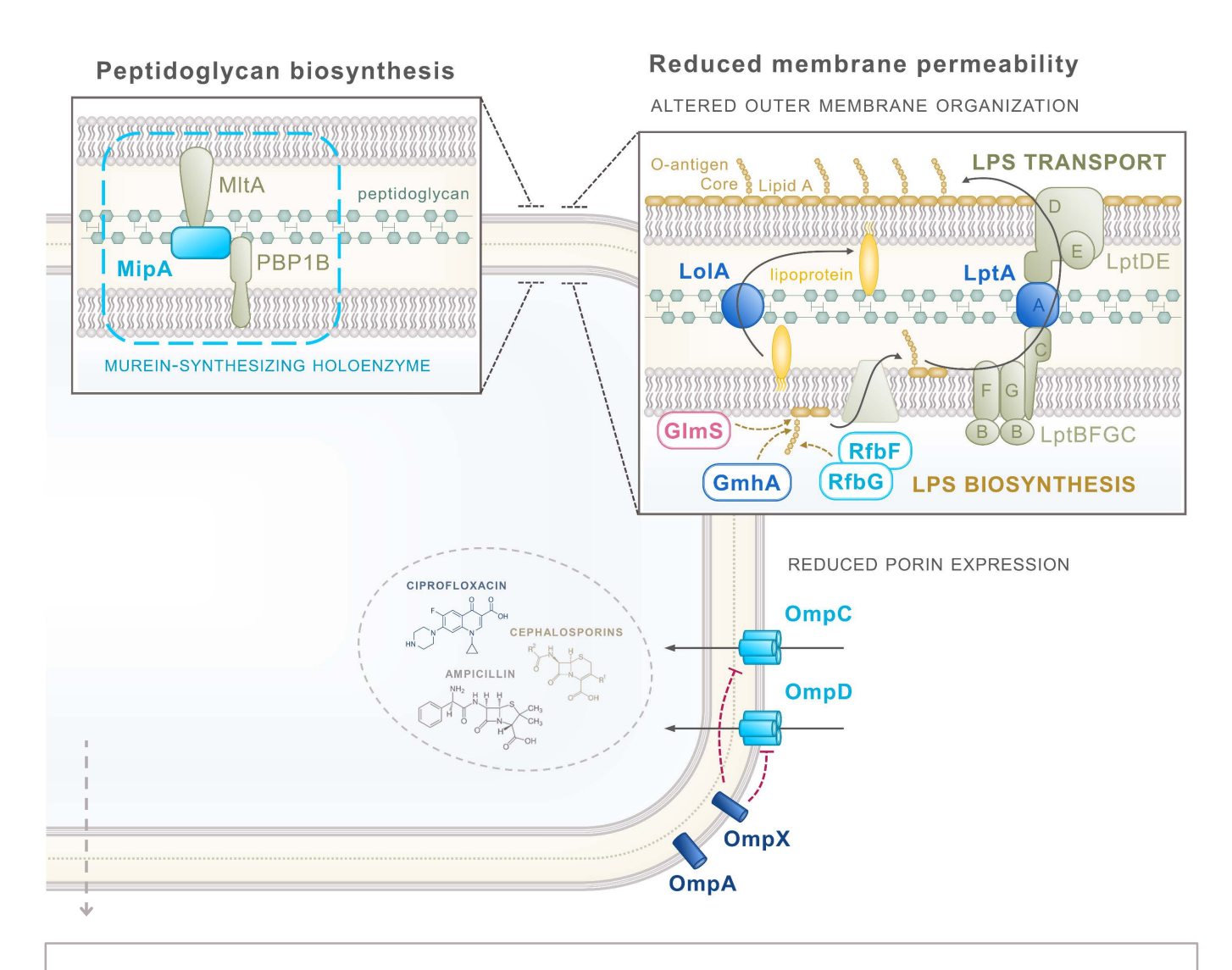
- A Salmonella Typhimurium DT104B isolate (Se6) was previously recovered from a hospitalized elderly patient with acute gastroenteritis before CIP treatment [1].
- An in vitro selected fluoroquinolone-resistant mutant (Se6-M) was experimentally obtained from the parental Se6 strain by a multi-step selecting process in the presence of increasing concentrations of CIP [2].
- Se6 and Se6-M were proven to be clonally related by PFGE.
- The different comparative subproteomic analyses were performed by combining 2-DE~LC-MSMS and shotgun LC-MSMS identification approaches for intracellular and membrane subproteomes, respectively.
- Proteomics data was validated by performing quantitative real time PCR for selected proteins of interest.



Representative 2-DE gels with the protein spots identified by LC-MSMS with significant differential abundance between the intracellular subproteomes of the S. Typhimurium DT104B strains (SameSpots software). Left: the protein spots more abundant in Se6 are represented on top and the more abundant in Se6-M are shown on the bottom. Right: the protein spots more abundant in Se6-M grown without ciprofloxacin stress are represented on top and the more abundant in Se20+CIP are shown on the bottom.



SCHEMATIC REPRESENTATION OF THE MAIN RESISTANCE MECHANISMS IDENTIFIED



Periplasmic substrate-binding proteins of ABC importers: Artl, FliY, Gltl/Gltl, GsiB, LsrB, MalE, ModA, MppA, Oxd-6a, PstS, RbsB, UgpB, YrbC

Global regulators of bacterial metabolism and bacterial stress response proteins: CspA, CspC, H-NS, IhfA, LexA, NusG, PhoP, RecA, Rho, SeqA, StpA,

Roles as determinants of antimicrobial resistance and/or targets for the development of new antimicrobial drugs

The proteomic signatures obtained provide valuable information about mechanism-related differential protein expression, supporting the current knowledge and leading to new testable hypotheses on the mechanisms of bacterial resistance to fluoroquinolones and other drugs.

- [1] de Toro M, et al (2010). *In vivo* selection of aac(6')-*Ib-cr* and mutations in the *gyrA* gene in a clinical *gnrS1*-positive *Salmonella* enterica serovar Typhimurium DT104B strain recovered after fluoroquinolone treatment. J Antimicrob Chemother 65(9): 1945-1949.
- [2] Correia S, et al (2016). Impacts of experimentally induced and clinically acquired quinolone resistance on the membrane and intracellular subproteomes of Salmonella Typhimurium DT104B. J Proteomics 145: 46-59.











