**Super model of the world – Enterococcus faecalis**

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*Enterococcus faecalis* is a Gram-positive bacterium from the lactic acid group (figure 1). This natural inhabitant of the mammalian gastrointestinal tract is also an opportunistic pathogen responsible for urinary tract infections, nosocomial infections, bacteremia and infective endocarditis [1]. *E. faecalis* normally grows as a commensal organism in the human gut, but it possesses very subtle virulence traits that are not easily identified. Additionally, its intrinsic physiological properties such as inherent antibiotic resistance and exceptional ability to adapt to harsh conditions, provides this organism with an enormous advantage in the infection processes.

Since the metabolic reconstruction of *Haemophilus influenzae* was published in 1999 [2], over 80 organisms have seen their sequenced genome being used to reconstruct a genome scale metabolic model. Of these genome-scale reconstructions, 57 belong to bacteria alone. Recognizing the medical importance of this bacterium, the first genome-scale metabolic model of *E. faecalis* was reconstructed. This model consists of 682 reactions, 655 metabolites, 516 genes (with associated metabolic function) and 584 reactions without gene association. The reconstruction process is elucidated in Figure 2 and followed the protocol described in reference 3. To support the in silico reconstruction, high throughput omic techniques (genomics, proteomics, fluxomics, metabolomics and transcriptomics) provided data to validate, test and sustain the robustness of the model. The first metabolic model generated for this bacterium allows correlating metabolite levels and fluxes and better understanding the key control points in its metabolism. As it has been previously shown for other organisms, the metabolic network reconstruction may serve as a valuable tool to predict the phenotypic behaviour of the microbe under various genetic and environmental conditions. With so many capabilities and strengths, *Enterococcus faecalis* is a good candidate to be the next super model of the bacterial world?


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