PT029 Optimizing locked nucleic acid/2’-o-methyl-rna fluorescence in situ hybridization (lna/2omez-fish) procedures for bacterial detection

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Background: Despite the successful application of locked nucleic acid/2'-O-methyl-RNA fluorescence in situ hybridization (LNA/2’OMe-FISH) procedures for bacteria detection, there is a lack of knowledge on the properties that affect hybridization. Such information is crucial to find the more suitable hybridization conditions for bacteria detection either in an individual or in a multiplex assay.

Objectives: This work aimed to evaluate the effect of three essential factors on the LNA/2’OMe hybridization step - hybridization temperature, [NaCl] and type/concentration of denaturant (formamide, ethylene carbonate and urea).

Methods: The optimization was performed for 3 Gram-negative bacteria (Escherichia coli CECT 515, Citrobacter freundii SGSC 5345 and Pseudomonas aeruginosa PAO1) and 2 Gram-positive bacteria (Enterococcus faecalis CECT 184 and Staphylococcus epidermidis RP61A), using an Eubacteria LNA/2’OMe probe (5’-mTIGlCmCITmCmCICmTAmGmGlA3’; “l” - LNA; “m” – 2’OMe). The signal quantification was evaluated by flow cytometry and Response Surface Methodology was used to model the interaction between the 3 parameters.

Results: It was observed that a high NaCl concentration is beneficial (2M-5M), regardless of the denaturant used. Urea, formamide and ethylene carbonate are suitable denaturants for LNA/2’OMe-FISH applications; but urea provides higher fluorescence intensities among the different bacteria. The results indicate that a hybridization solution with 2M of urea and 4M of NaCl would be a proper starting point for multiplex LNA/2’OMe-FISH procedures. Furthermore, a hybridization temperature around 62°C, for 14bp probes with LNA monomers at every third position of 2’OMe might be use in initial optimizations.