

Azoreductase activity of dye-decolorizing bacteria isolated from the human gut microbiota

Amal Emad, Marwa Tammam, Sara Zahran, Abdelgawad M. Hashem, Ramy K. Aziz

Abstract

The gut microbiota enriches the human gene pool and contributes to xenobiotic metabolism. Microbial azoreductases modulate the reduction of azo-bonds, activating produgs and azo polymer-coated dosage forms, or degrading food additives. Here, we aimed to screen the healthy human gut microbiota for food colorant-reducing activity and to characterize factors modulating it. Four representative isolates from screened fecal samples were identified as *E. coli* (AZO-Ec), *E. faecalis* (AZO-Ef), *E. avium* (AZO-Ev) and *B. cereus* (AZO-Bc). Both AZO-Ef and AZO-Ev decolorized amaranth aerobically and microaerophilically while AZO-Ec and AZO-Bc had higher aerobic reduction rates. The isolates varied in their activities against different dyes, and the azo-reduction activity mostly followed zero-order reaction kinetics, with a few exceptions. Additionally, the isolates had different pH dependence, e.g., AZO-Ec was not affected by pH variation while AZO-Bc exhibited variable degradation kinetics at different pH levels. Cell-free extracts showed NADH-dependent enzymatic activities 3663 ; " times higher than extracellular fractions. FMN did not affect the reducing activity of AZO-Ef cell-free extract, whereas AZO-Ec, AZO-Ev and AZO-Bc had significantly jki jgt"tgfwewkqp"tcvgu"kp"kvu"rtgugpeg"R xcnewgu ? 2024. "202223"cpf"2024." respectively). Using Degenerate primers allowed the amplification of azoreductase genes, whose sequences were ; :ó; ; ' "similar to genes encoding FMN-dependent-NADH azoreductases.

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