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Impact of two disinfectants on detachment of Enterococcus faecalis from polythene in aquatic microcosm

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After cell adhesion processes in microcosm, the impact of sodium hypochlorite (NaOCl) and hydrogen peroxide (H₂O₂) on the detachment of Enterococcus faecalis from polythene fragments immersed in water under stationary and dynamic conditions was assessed. The abundance of planktonic cells was also evaluated. The density of E. faecalis adhered in absence of disinfectant fluctuated between 2 and 4 units (Log CFU/cm²). After living in disinfected water, the density of E. faecalis remained adhered to polythene sometimes reached 2 units (Log CFU/Cm²). This highest abundance of cells remained adhered was recorded with cells coming from the lag, exponential and stationary growth phases in water treated with 0.5‰ NaOCl. In H₂O₂ disinfected water, the highest value was recorded at all cells growth phases with 5‰ H₂O₂ concentration. Adhered E. faecalis cells have been sometimes completely or partially decimated respectively by NaOCl and H₂O₂ treated water. Considering separately each experimental condition, it was noted that increasing the concentration of disinfectant caused a significant decrease (P≤0.01) in abundance of cells stay adhered after living in water disinfected by the two disinfectants. Changes in disinfectant concentrations in different experimental conditions had an impact on the detachment of E. faecalis cells from the substrates.

Keywords: Disinfectants, Water treatment, E. faecalis, Cell growth phase, Adhesion, Detachment

The distribution network of drinking water is often the scene of many physicochemical and biological reactions resulting from interactions between disinfectants, pipe walls and the free and fixed biomasses. Microbiology of drinking water distribution networks is of great importance because of health risks in the short term and general degradation of water quality or its products (Dukam et al., 1995). Microorganisms are usually found in distribution networks and in storage tanks...
and cause deterioration in the organoleptic water quality (Mouchet et al., 1992; Dogett, 2000).

To remedy this situation, recourse to the use of disinfectants is common. The effectiveness of disinfectants in drinking water distribution network varies from one application to another. While almost 100% of cells found in planktonic state can be eliminated, the yield is lower on adherent cells (Allion, 2004). Disinfectants used in the treatment of drinking water presents a wide spectrum of action if nothing hinders their activity. They have a preferential action on a type of microorganisms, and if they are not applied in conditions where their lethal activity can be expressed, they favor the selection and even sometimes dangerous proliferation of surviving species. This justifies sometimes the formation of biofilms in water pipes walls despite treatment with disinfectants. Microorganisms in the pipe are either of internal origin (microorganisms that escaped water disinfection processes) or of external origin by water return during intervention work on the network, through cracks caused by aging pipes) (Van der Kooij and Hijnen, 1988).

Quality control analyses of drinking water are based not only on physical, chemical, but also microbiological parameters. The purpose of drinking water treatment is not to produce sterile water, but water that does not present a risk from a public health viewpoint. These analyses showed that Enterococcus faecalis cell is a minority commensal germ of the gastrointestinal and genito-urinary tracts of humans and warm-blooded animals. However, it may become pathogenic in immuno-compromised individuals and elderly persons where it causes more than 10% of nosocomial infections (Jett et al., 1994; WHO, 2004). Previous work has clearly shown that E. faecalis’s adaptability to harsh environments makes it possible to find it in different kinds of waters (Manero and Blanch, 1999). It is an indicator of fecal contamination usually involved in the occurrence of nosocomial infections. It is found in human feces as well as those of warm-blooded animals with concentrations of up to $10^{12}$ cells per gram (Beaupoil et al., 2010). In waste water, they are found at a concentration of $10^8$ CFU/l (Manero and Blanch, 1999).

In aquatic microcosm containing E. faecalis cells and solid particles, the abundances of planktonic E. faecalis cells decreased with incubation periods. This decrease in abundance of cells remained planktonic helped to note that some cells adhere to the support when it is immersed (Hoiby et al., 2010). On this basis, it was thought that the same phenomenon can occur in water distribution networks and cause the formation of biofilms. Then was posed the question, what would be the impact of disinfectants on the physiological and metabolic state of cells detached from substrates?

In a biofilm, microorganisms can develop, and due to their protection in the shell of biofilm, they could potentially become pathogens and pose a public health problem. In fact, studies have shown that bacteria biofilm are frequently observed on the inner walls of the drinking water pipes. These bacteria dropped in water pipes are sources of contamination and health risks to consumers (Allion, 2004; Boutaleb, 2007). In addition, biofilms are escape routes of microorganisms faced with disinfectants.

Although previous studies helped to understand the formation and development of biofilms on rocky substrates and the inside walls of drinking water pipes (Moungang et al., 2013a; Moungang et al., 2013b), the impact of the presence of disinfectants on microorganisms adhered to these walls so far have been paid very little attention. Some authors worked on the physiological and
metabolic state of the microorganisms and their detachment in the presence of sodium hypochlorite (NaOCl) or hydrogen peroxide (H₂O₂) (Lontsi Djimeli et al., 2013; Lontsi et al., 2014a). This study aims at evaluating in microcosm conditions with respect to incubation periods and different physiological conditions, the impact of sodium hypochlorite (NaOCl) or hydrogen peroxide (H₂O₂) on the detachment of *E. faecalis* cells from polythene fragments immersed in water under stationary and dynamic regimes.

**Materials and methods**

**Collection and identification of *Enterococcus faecalis***

*Enterococcus faecalis* strain was provided by the Laboratory of Microbiology and Environment of Centre Pasteur of Cameroon. After subculture on plate count agar (PCA), Gram staining was performed on the smear of *E. faecalis*. Identification tests were performed by standard biochemical methods (Holt *et al.*, 2000; APHA, 2005; Marchal *et al.*, 1991; Rodier, 2009). These identifications revealed that *Enterococcus faecalis* is oxidase positive generally catalase negative and quickly reduces triphenyl tetrazolium chloride (TTC). It is a facultative anaerobic Gram positive bacterium. The bacterium *Enterococcus faecalis* was highlighted on Bile-Esculin Azide (BEA) medium whose colonies is translucent and surrounded by a black halo. The black halo surrounding colonies reflects the production of H₂S and the hydrolysis of esculin to esculetin that binds with iron (Holt *et al.*, 2000).

**Assessment of *Enterococcus faecalis* cell growth phases**

*E. faecalis* cell growth phases were evaluated using growth curve (Rubio, 2002). Three replicates of 15 test tubes each containing 10 ml sterile peptone (Biokar) solution was used. Tubes of each set were labelled t₀, t₂, t₄, t₆, t₈, t₁₀, t₁₂, t₁₄, t₁₆, t₁₈, t₂₀, t₂₂, t₂₄, t₂₆ and t₂₈. Prior to the experiment, *E. faecalis* strains previously freezer stored in glycerol vials were defrosted at room temperature in the laboratory. The culture (300 μl) was then transferred into 10 ml of nutrient broth (Oxford) and incubated at 37°C for 24 hours and cells latter collected by centrifugation (8000 rpm for 10 min at 10°C) and washed twice with sterile NaCl (8.5 g/l) solution. The sediment was then diluted in 10 ml of sterile NaCl solution. After dilution, 100 μl of the suspension was added to each of the 15 tubes containing 10 ml of sterilized peptone solution. The cell suspensions in the 3 tubes labelled t₀ were immediately analyzed. Those in the tubes labelled t₂, t₄, t₆ … t₂₈ were incubated for 2, 4, 6… 28 hours at 37°C.

After each incubation period, analyses were carried out using spread plate method on BEA culture medium, incubated for 48 hours at 37°C. The colony forming units (CFUs) were then counted. Mean CFUs were calculated from the results of the triplicates and Log (CFU) also calculated. The straight Log (CFUs/ml) curve against storage period was plotted and compare to the cell growth curve. The cell growth phases of *E. faecalis* were then determined.

**Absorbing substrate used**

The adsorbing substrate used was high dense polythene. It differed from low radical dense polythene and low linear dense polythene by sparsely branched chains of its molecular structure, and its relatively high resistance to shocks, high temperatures and ultraviolet rays (Moungang *et al.*, 2013a; Coeyrehourcq, 2003). It is a plastic piping material obtained directly from the manufacturer and used in drinking water distribution networks. High dense polythene results from polymerization of macromolecules of polyolefin family. This
polymerization is obtained from gaseous ethylene according to the following equation (Ratner, 1993; Ratner, 1995):

\[ n \ H_2C=CH_2 \rightarrow \left( \begin{array}{c} H \\ C \\ H \\ H \end{array} \right)_n \]

The polythene used in this study is commercialized by Goodfellow SARL (France).

**Disinfectants used**

Two disinfectants were used: NaOCl (Colgate-Palmolive, USA), which belongs to the group of halogen derivatives, and H₂O₂ (Gilbert, France) which belongs to the group of oxidants. The ease with which these two disinfectants are generally used for the treatment of drinking water justifies their choice for this study. Concentrations of NaOCl used ranged from 0.5 to 1.5‰, while those of H₂O₂ oscillated between 5 and 15‰. These concentrations were evaluated by simple method of dilution of crude solution obtained directly from the manufacturer. To count the surviving microorganisms after disinfection, the sterile NaCl solution was used as a diluent.

**Adhesion test of E. faecalis on polythene**

On the basis of previous studies (Lontsi Djimeli et al., 2013; Lontsi Djimeli et al., 2014a; Lontsi Djimeli et al., 2014b), parallelepipedic shaped fragments of polythene with 13.28 cm² of total surface area suspended to a wire of 0.1 mm diameter were immersed in triplicate in two sets named A and B. Set A contained four subsets each having three Duran’s 250 ml flasks labelled as follows: A1, A1', A1'', A2, A2', A2'', A3, A3', A3'', and A4, A4', A4''. Same for set B with labelling as follows: B1, B1', B1'', B2, B2', B2'', B3, B3', B3'' and B4, B4', B4''. Each flask contained 99 ml of NaCl solution. Meanwhile, controls were made and coded A01, A02, A03, A04 and B01, B02, B03, B04 (Noah Ewoti et al., 2011). The whole was then autoclaved. Prior to the experiment, E. faecalis strains previously freezer stored in glycerol vials were defrosted at room temperature. The culture (300 μl) was then transferred into 10 ml of nutrient broth (Oxford) and incubated at 37°C for 24 hours and the cells latter collected by centrifugation at 8000 rpm for 10 min at 10°C, then washed twice with sterile NaCl solution. The sediment was then diluted in 10 ml NaCl solution.

After serial dilutions, the initial concentration of bacterium cells (concentration at the initial moment t=0) in each mother solution was adjusted to 8.77815 units (Log (CFU/ml)). This was performed by reading the optical density (OD) at 600 nm using a spectrophotometer (DR 2800) followed by culture on BEA medium (Marchal et al., 1991).

Afterwards, a volume of 1 ml of the suspension was added to 99 ml of sterile physiological water contained in each flask. Erlenmeyer flasks labelled A1, A1', A1'', A2, A2', A2'', A3, A3', A3'', and A4, A4', A4'' were incubated with stirring at a speed of 60 revolutions/minutes, using a stirrer (Rotatest brand). Erlenmeyer flasks, labelled B1, B1', B1'', B2, B2', B2'', B3, B3', B3'' and B4, B4', B4'' were incubated under static condition. Erlenmeyer flasks labelled A1, A1', A1'' and B1, B1', B1'' were incubated for 180 min. Those labelled A2, A2', A2'' and B2, B2', B2'' B1'' were incubated for 360 minutes. Erlenmeyer flasks labelled A3, A3', A3'' and B3, B3', B3'' were incubated for 540 minutes. Those labelled A4, A4', A4'' and B4, B4', B4'' were incubated for 720 min. All these incubations were made at room temperature in the laboratory (25±1 °C).

**Determination of abundance of E. faecalis cell remained planktonic**

E. faecalis cell adhesion to polythene was performed in sterile physiological water
contained in the Erlenmeyer flasks. This led to assess the abundance of *E. faecalis* cells remained planktonic at the initial time (T0) and densities of cells remained planktonic after 180, 360, 540 and 720 minutes of incubation. The collection and enumeration of cells remained planktonic were carried out after removal of each polythene fragment from Erlenmeyer by monitoring the cells growth on BEA medium followed by incubation at 37 °C for 24 to 48 hours.

**Impact of NaOCl and H\textsubscript{2}O\textsubscript{2} on the detachment of *E. faecalis* cells from the polythene fragments**

The contaminated substrates allowed to highlight the impact of NaOCl and H\textsubscript{2}O\textsubscript{2} concentrations on cells detachment for 25 to 30 minutes. The disinfecting effect was stopped by introducing substrates in 10 ml of sterile saline. Cells detachment and appreciation of the reduction of the bacteria load were assessed after culture of surviving bacteria.

Depending on the type of disinfectant tested, the fragments removed from Erlenmeyer flasks A1, A2, A3, A4, B1, B2, B3 and B4 after 30 minutes from water treatment with 0.5‰ NaOCl or 5‰ H\textsubscript{2}O\textsubscript{2}. The fragments removed from flasks A1', A2', A3', A4', B1', B2', B3' and B4' after 30 minutes living in water disinfected with 1‰ NaOCl or 10‰ H\textsubscript{2}O\textsubscript{2}. Similarly, those removed from Erlenmeyer flasks A1'', A2'', A3'', A4'', B1'', B2'', B3'' and B4'' after 30 minutes living in water treated 1.5‰ NaOCl or 15‰ H\textsubscript{2}O\textsubscript{2}. After this disinfection test of 30 minutes at room temperature under static condition, each fragment was released and drained sterile and introduced into 10 ml sterilized NaCl solution.

The unhooking of adherent cells was performed by vortex agitation at increasing speeds for 30 seconds in three consecutive series of 10 ml sterilized NaCl solution. This technique allowed the unhooking of maximum adhered cells (Dukam *et al.*, 1995; Noah Ewoti *et al.*, 2011). The total volume of the suspension containing unhooked *E. faecalis* cells was 30 ml. Collection and numbering of unhooked cells were performed using the spread plate method on BEA medium, followed by the incubation in Petri dishes at 37°C for 24 to 48 hours. The disinfectant was not evaluated after incubation.

**Data analysis**

Variations in abundance of planktonic and adhered *E. faecalis* cells in each experimental condition were illustrated by semi-Logarithmic curves. Standard deviations were not considered because the curves were too close. Spearman "r" correlation test was used to assess the degree of relation between the abundance of adhered *E. faecalis* cells and incubation periods for each concentration of disinfectant on one hand, and between the abundance of the cells remained adhered and concentrations of disinfectants for each incubation period and for each cell growth phase on the other hand. To compare the mean abundance of *E. faecalis* remained adhered from one experimental condition to another, Kruskal-Wallis H test and Mann-Whitney U test were used using the statistical software package SPSS 17.0. A *P*-value of 0.05 was assumed to be statistically significant.

**Results and discussion**

**Enterococcus faecalis growth curve**

The growth of *E. faecalis* in non renewed sterile tryptone liquid medium (Biokar) exhibited a hyperbolic curve of 4 phases (Fig. 1): a lag growth phase of 2 hours, an exponential growth phase from the 5\textsuperscript{th} to the 11\textsuperscript{th} hour of incubation, a stationary growth phase of 12 hours and a decline growth phase which began from the 22\textsuperscript{nd} hour of incubation (Fig. 1). Bacterial growth is an orderly increment of all the components of
the bacterium (Rubio, 2002). It leads to an increase in the number of bacteria. During growth, there is, on one hand, a depletion of nutrients in the culture medium and, on the other hand, an enrichment of products of metabolism, that are toxic. During the lag growth phase, the growth rate is nil. Bacteria adapt and synthesize the enzymes necessary to metabolize new substrates. The exponential growth phase corresponds to the period of nutrient utilization and duplication of cell number. The stationary growth phase is the period when the growth rate becomes nil. In fact, the bacteria multiply compensating those who die. The decline growth phase is the time when all food resources are exhausted. There is accumulation of toxic metabolites. There is a decrease of viable organisms and an occurring of cell lysis by the action of endogenous proteolytic enzymes. However, there is a persistant growth leading to the release of substances during lysis (cryptic growth) (Rubio, 2002).

![Growth curve of E. faecalis](image)

**Figure 1.** Growth curve of *E. faecalis* (LP: Lag phase; EP: Exponential growth phase; SP: Stationary growth phase; DP: Decline growth phase).

**Assessment of *E. faecalis* cell abundance remained planktonic**

The densities of *E. faecalis* cells remained planktonic ranged from 8.77776 to 8.77807 units (Log CFU/Cm²). The maximum value was recorded after 180 minutes with cells coming from the lag growth phase. The minimum value was registered after 720 minutes with cells coming from the stationary growth phase (Fig. 2). When considering either under stationary or dynamic regime, a relative decrease was noted with abundance of *E. faecalis* cells remained planktonic at each incubation period and each cell growth phase. This decrease in abundance of cells remained planktonic revealed that, some cells adhered to polythene fragment follow the reversible and irreversible adhesion mechanisms. According to Pouneh (2009), bacterial adhesion to supports takes place in two main stages: adherence and adhesion, which
respectively correspond to a reversible adhesion and an irreversible fixation of bacteria. The reversible adhesion is usually non-specific and short-term (5-10 hours) (Gauthier and Isoard, 1989). The irreversible adhesion is a step slower than the former. The irreversibility of adhesion is due to the fact that the bacterium secretes a matrix of exopolymers forming an envelope around it called glycocalyx, and other specific molecules (adhesins) that allow it to consolidate its adhesion to the substrate (Rubio, 2002).

**Figure 2.** Temporal variations of abundance of *E. faecalis* cells remained planktonic at each cell growth phase under static and dynamic conditions.

**Impact of NaOCl and H₂O₂ on the detachment of *E. faecalis* cells adhered to polythene**

After the adhesion process of *E. faecalis*, the impact of these disinfectants on the detachment of cells remained adhered to polythene was assessed by allowing to stay for the specific periods, the contaminated materials in water disinfected with increasing of NaOCl and H₂O₂ concentrations. After the adhesion test, followed by the disinfection test, some *E. faecalis* cells remained adhered to the polythene fragments. These cells remained adhered were themed "cells remained".
Figure 3. Temporal evolution of *E. faecalis* cells remained adhered to polythene under static condition in NaOCl and H$_2$O$_2$ treated water.

When using NaOCl under static condition, the density of *E. faecalis* cells remained adhered to polythene varied with different cells growth phases and different disinfectant concentrations (Fig. 3). The abundance of adhered *E. faecalis* cells in the absence of disinfectant ranged from 1 to 3 units (Log CFU/Cm$^2$). The lowest value was recorded with cells coming from decline cell growth phase after 180 minutes incubation period.
The highest value was registered with cells harvested from lag and exponential growth phases after 540 and 720 minutes incubation durations. In water treated with NaOCl, the abundance of cells remained adhered often reached 2 units (Log CFU/Cm²). This highest abundance of *E. faecalis* cells remained adhered was recorded with cells coming from the lag, exponential and stationary growth phases in water treated with 0.5‰ NaOCl. Adhered *E. faecalis* cell has been sometimes completely decimated by NaOCl treated water under dynamic condition (Fig. 3).

**Figure 4.** Temporal evolution of *E. faecalis* cells remained adhered to polythene under dynamic condition in NaOCl and H₂O₂ treated water.
Table 1. Spearman "r" Correlation coefficients between abundance of *E. faecalis* cells remained adhered to polythene and incubation periods for each concentration of disinfectant and each experimental condition.

<table>
<thead>
<tr>
<th>Experimental conditions</th>
<th>Disinfectants and concentrations</th>
<th>NaOCl</th>
<th>H2O2</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0.5‰</td>
<td>1‰</td>
<td>1.5‰</td>
</tr>
<tr>
<td>Static</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>540 min</td>
<td>-0.601*</td>
<td>-0.517*</td>
<td>-0.461</td>
</tr>
<tr>
<td>360 min</td>
<td>-0.759**</td>
<td>-0.534*</td>
<td>-0.479</td>
</tr>
<tr>
<td>Dynamic</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

** : P ≤0.01  * : P ≤0.05  df=15

Table 2. Spearman "r" Correlation coefficients between cells abundance of *E. faecalis* remained adhered to polythene and disinfectant concentrations in each incubation period and each experimental condition.

<table>
<thead>
<tr>
<th>Experimental conditions</th>
<th>Disinfectants and incubation durations</th>
<th>NaOCl</th>
<th>H2O2</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>180 min</td>
<td>360 min</td>
<td>540 min</td>
</tr>
<tr>
<td>Static</td>
<td>-0.415</td>
<td>-0.444</td>
<td>-0.415</td>
</tr>
<tr>
<td>Dynamic</td>
<td>-0.401</td>
<td>-0.490</td>
<td>-0.562</td>
</tr>
</tbody>
</table>

** : P≤0.01  * : P≤0.05  df=11

Table 3. Comparison between mean abundance remained adhered on polythene in disinfected water and different cell growth phases.

<table>
<thead>
<tr>
<th>Disinfectants</th>
<th>Cell growth phases</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Lag</td>
</tr>
<tr>
<td>NaOCl</td>
<td>P=0.019*</td>
</tr>
<tr>
<td>H2O2</td>
<td>P=0.008*</td>
</tr>
</tbody>
</table>

*: P≤0.05  df=22

Under dynamic condition, the abundance of *E. faecalis* cells remained adhered to polythene also varied with different cell growth phases and different concentrations of NaOCl (Fig. 4). The density of *E. faecalis* adhered in absence of disinfectant under dynamic condition fluctuated between 2 and 4 units (Log(CFU/cm²)). The highest value was recorded with cells harvested from lag and exponential growth phases after 720 minutes incubation duration. The lowest value was registered with cells coming from all cell growth phases after 180 minutes incubation period. The abundance of *E. faecalis* remained adhered to polythene in NaOCl treated water sometimes reached 2 units (Log CFU/Cm²). The highest value was registered at all cells growth phases after living in water disinfected with 0.5‰ NaOCl. Under dynamic condition, adhered *E. faecalis* cell has been sometimes completely decimated by NaOCl treated water (Fig. 4).

It was observed that under dynamic condition, the abundance of *E. faecalis* cells remained adhered to polythene was relatively higher than that observed under static condition. In general, when the concentration of NaOCl in water increased, the abundance of *E. faecalis* cells remained adhered decreased with time (Figs 3 and 4).

When using H2O2 and under static condition, it is clear from figures 3 and 4 that the cells of *E. faecalis* remained adhered to polythene generally varied with different cell growth phases and concentrations of H2O2. The density of *E. faecalis* adhered in absence of H2O2 under static condition ranged from 2
to 4 units (Log CFU/Cm²). The highest value was recorded with cells coming from exponential cell growth phase after 720 minutes incubation duration. The lowest value was registered with cells coming from all cell growth phases after 180 and 360 minutes incubation periods. The abundance of *E. faecalis* remained adhered to polythene in H₂O₂ disinfected water often reached 2 units (Log CFU/Cm²). Under static condition, adhered *E. faecalis* cell has been sometimes partially decimated by H₂O₂ treated water. The highest value was recorded at all cells growth phases in water disinfected with 5‰ H₂O₂ after 360, 540 and 720 minutes incubation periods (Fig. 3).

Under dynamic condition, the density of adhered *E. faecalis* cells in the absence of H₂O₂ oscillated between 2 and 4 units (Log CFU/Cm²). The highest value was registered with cells harvested from lag and exponential growth phases after 720 minutes incubation duration. The lowest value was recorded with cells coming from all cell growth phases after 180 minutes incubation period. The abundance of cells remained adhered to polythene often reached 2 units (Log CFU/cm²) in H₂O₂ treated water. This value was recorded with cells coming from the lag, exponential and stationary growth phases in water treated with 5 and 10‰ H₂O₂. Adhered *E. faecalis* cell has been sometimes partially decimated by H₂O₂ treated water under dynamic condition (Fig. 4).

The abundance of *E. faecalis* cells remained adhered to polythene was generally more important under dynamic condition than under static condition. In most cases, the abundance of *E. faecalis* cells remained adhered to polythene decreased with time as the concentration of H₂O₂ increased in water (Figs 3 and 4).

Detachment of adhered *E. faecalis* cells from polythene depends on the incubation period, cell growth phase and concentrations of disinfectant. Increasing the duration of the disinfection process in most cases leads to a significant decrease in abundance of *E. faecalis* cells remained adhered to polythene in each solution treated with NaOCl or H₂O₂ under static and dynamic conditions. In fact, *E. faecalis* cells fixed on a substrate and produce extracellular polymers have an altered phenotype compared to corresponding remaining planktonic cells, particularly with regards to growth, gene transcription, protein production and intercellular interaction (Hoiby et al., 2010). In addition, the reduction of penetration of molecules including disinfectants due to changes in cell density and production of exopolysaccharides, slow growth, modulation of stress response and other metabolic processes are the main causes of the reduction of the sensitivity of cells adhered to disinfectants (Keren et al., 2004).

It is noted that the abundance of *E. faecalis* remained adhered in all cell growth phases was relatively lower in general, in the presence of NaOCl or H₂O₂ than in the absence of these two disinfectants, whether under static or dynamic conditions. NaOCl is the best chlorine molecule used as a disinfectant. Its biocidal efficacy is based on the penetration of the chemical substance and its action on the essential oxidative enzymes of the cell (Lomander et al., 2004).

Relationships amongst the considered parameters

The Spearman "r" correlation coefficient between abundance of adhered cells and incubation periods for each concentration of disinfectant and experimental condition was calculated and are presented in Table 1. It is noted that increasing the duration of adhesion caused a significant decrease of cell abundance remained adhered to polythene in each disinfectant solution (P≤0.01).

Compared to low concentrations, increasing the concentration of disinfectants caused a significant decrease (P≤0.01) in cells
abundance of *E. faecalis* remained adhered after a stay in water disinfected with NaOCl and H₂O₂. This resulted in a relative increase in the efficacy of NaOCl and H₂O₂ on *E. faecalis* cells remained adhered to polythene as the disinfectants concentration increased in the medium (Table 2). Indeed, biofilms are a protection means for growth, allowing microorganisms to survive in hostile environments and conferring reduced susceptibility to dehydration, antibiotics and disinfectants. The cells attached to surfaces (supports) have reduced sensitivity to disinfectants compared to cells remained planktonic (Smith and Hunter, 2008). A biofilm can develop within hours and thus allow bacteria therein to become resistant to external agents causing contamination (Beech and Coutinho, 2003; Beech and Sunner, 2004). In addition, adhered cells often induce increased resistance to disinfecting agents compared to planktonic cells (Smith and Hunter, 2008; Parot, 2007). Prolonged contact between cells and the support leads to the secretion in situ of proteins which enhance their interactions. These interactions are thereafter inhibited by the disinfectants (Guillemot, 2006). On the other hand, for each incubation period, the action of NaOCl and H₂O₂ on *E. faecalis* cells remained adhered to polythene increased significantly (Ps0.01). The action of these disinfectants is explained by the action of disinfectant molecules that are chemically more reactive on biofilms (Ntsama et al., 1995). Furthermore, this variation of reaction of *E. faecalis* cells in the face of disinfectants may be related to changes in their surface following a change in their cell growth phase (Briandet et al., 1999).

Table 3 revealed a significant difference (Ps0.05) between the mean abundance of *E. faecalis* remained adhered on polythene after a stay in treated water and the different concentrations of NaOCl at different cell growth phases. This difference is due to a nutritional limitation experimented with these cells and which is the cause of variation in the growth rate (Yasuda et al., 1993; Yasuda et al., 1994; Suci et al., 1994). Similarly, at each cell growth phase, there was a significant difference (Ps0.05) between the mean abundance of *E. faecalis* cells remained adhered after a stay in waters treated with different concentrations of NaOCl (Table 3). The effectiveness of any disinfection method depends on biotic factors such as physiological state and the intrinsic microbial resistance to lethal agents (Parot, 2007). *E. faecalis* cells remained adhered to polythene under dynamic condition are more sensitive to NaOCl and H₂O₂ than those remained adhered under static condition. According to (Klausen et al., 2006) this could be explained by the structure of cells that remained adhered which depends on a hydrodynamic regime. Moreover, NaOCl and H₂O₂ act on several components of the bacteria and biofilm while other disinfectants (chlorhexidine) act only on the cell wall (Leung et al., 2012).

**Conclusion**

This study focused on assessing in microcosm condition the impact of NaOCl and H₂O₂ on the detachment of *E. faecalis* cells adhered to polythene fragments immersed in water under static and dynamic conditions. It appeared from this study that, after the adhesion process, *E. faecalis* cells remained adhered to polythene revealed temporal variations. These changes are related to biotic and abiotic properties of *E. faecalis* cells and polythene.

Considering separately each experimental condition, it was noted that increasing the duration of the adhesion process causes a significant decrease of abundance of *E. faecalis* cells remained adhered to polythene in each solution treated.
with NaOCl or H₂O₂. Similarly, increasing the concentration of disinfectants caused a significant decrease in abundance of E. faecalis cells remained adhered after a stay in water disinfected with two disinfectants. No significant differences were recorded between the mean densities of E. faecalis cells remained adhered under static condition and those obtained under dynamic condition. Changes in disinfectant concentrations under different experimental conditions have an impact on the detachment of E. faecalis cells adhered to substrates. The same disinfectant could have a different impact on adhered cells in different contexts.

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Conflict of Interests
The authors declare that they have no conflict of interests that could inappropriately influence this work.

References


Guillemot G. 2006. Compréhension des mécanismes à l’origine de l’adhésion de


