Wastewater Treatment by Catalytic Wet Peroxidation Using Nano Gold-Based Catalysts: A Review

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Abstract: Nowadays, there is an increasing interest in the development of promising, efficient, and environmentally friendly wastewater treatment technologies. Among them are the advanced oxidation processes (AOPs), in particular, catalytic wet peroxodation (CWPO), assisted or not by radiation. One of the challenges for the industrial application of this process is the development of stable and efficient catalysts, without leaching of the metal to the aqueous phase during the treatment. Gold catalysts, in particular, have attracted much attention from researchers because they show these characteristics. Recently, numerous studies have been reported in the literature regarding the preparation of gold catalysts supported on various supports and testing their catalytic performance in the treatment of real wastewaters or model pollutants by CWPO. This review summarizes this research; the properties of such catalysts and their expected effects on the overall efficiency of the CWPO process, together with a description of the effect of operational variables (such as pH, temperature, oxidant concentration, catalyst, and gold content). In addition, an overview is given of the main technical issues of this process aiming at its industrial application, namely the possibility of using the catalyst in continuous flow reactors. Such considerations will provide useful information for a faster and more effective analysis and optimization of the CWPO process.

Keywords: catalytic wet peroxidation; radiation; gold-based catalysts; wastewater treatment; advanced oxidation processes

1. Introduction

The world’s population growth and increasing industrial development led to the intense usage of natural resources with the water bodies being used as a final destination for wastewater containing pollutants [1–3]. The discharge of untreated wastewater introduces persistent contaminants into the environment, some examples being metals, organic, and inorganic compounds [4–6], which have harmful effects on ecology and public health [7].

In an attempt to minimize the impacts of effluent discharges, the European Union Water Framework Directive (EU-WFD), in 2000, imposed maximum permissible values for ecotoxic or possibly ecotoxic substances [8]. Thus, it is mandatory to adopt practical, efficient, and low-cost effluent purification technologies [9,10], which will allow the complete elimination or, at least, reduction of the contaminants.
concentration up to the limit values imposed by legislation [10,11], before wastewaters are discharged into water bodies.

The wastewaters can be treated by physical-chemical processes, such as sedimentation, coagulation/flocculation, filtration, adsorption, ultrafiltration, reverse osmosis, ion exchange, or chemical precipitation [12,13], by biological degradation [13–15], and/or by conventional oxidative processes, which degrade the pollutant by the action of oxygen or other oxidants, such as hydrogen peroxide, ozone, and permanganate [16–18]. Physical-chemical processes are not very appealing because the pollutants are concentrated at another phase, which requires a subsequent treatment [10]. Biological degradation, although economically advantageous, is inefficient since the compounds present in effluents are very often toxic and/or non-biodegradable [19,20]. Moreover, conventional oxidative processes might not have enough capacity to completely oxidize refractory compounds with high chemical stability and, therefore, there is a high risk of intermediate products being formed during oxidation, which can be even more toxic than the initial ones [21–23].

Advanced oxidation processes (AOPs) are emergent and attractive treatment technologies to degrade compounds with high chemical stability, toxicity, and non-biodegradability [10,24]. AOPs generate the hydroxyl radical (HO•), responsible for oxidizing refractory organic compounds into non-toxic products, such as CO₂ and H₂O [10,25–27]. Given the high efficiency of the hydroxyl radical, the AOPs have been widely used, not only in wastewater treatment [9,19,28–30], but also in soil and sediment remediation [31,32], decontamination of gaseous effluents containing volatile organic compounds and elimination of odors [33–36], water and groundwater treatment [37–39], and conditioning of municipal sludge [40,41].

Several AOPs are available, as will be detailed in the next section, that use different oxidants, with or without catalysts, in the presence of absence of radiation. Herein, we will focus on the catalytic wet peroxidation (CWPO) process using nano gold-based catalysts for wastewater treatment. This process presents several advantages compared to other AOPs, namely: it uses environmentally friendly reagents, does not require sophisticated equipment, and is operated under mild conditions of pressure and temperature. Moreover, catalysis by gold presents additional advantages, such as non-leaching of the metal to the treated effluent and efficient and stable performance, which are important for industrial applications.

A survey of the catalyst properties, operating conditions, and their effect on the efficiency of the process will be discussed. To the best of the authors knowledge, such review has not yet been reported in the literature.

2. Advanced Oxidation Processes

As mentioned above, AOPs are based on the formation of the hydroxyl radical. This radical has a high oxidation potential (2.8 eV [42,43]), being immediately below the fluorine (see Table 1), and exhibits high oxidation reaction rates, compared to traditional oxidants, such as chlorine, hydrogen peroxide, or potassium permanganate [11].

| Chemical Species          | Oxidation Potential (eV) |
|---------------------------|--------------------------|
| Fluorine                  | 3.03                     |
| Hydroxyl radical          | 2.80                     |
| Atomic oxygen             | 2.42                     |
| Ozone                     | 2.07                     |
| Hydrogen peroxide         | 1.77                     |
| Potassium permanganate    | 1.67                     |
| Hypobromous acid          | 1.59                     |
| Chlorine dioxide          | 1.50                     |
| Hypochlorous acid         | 1.49                     |
| Chlorine                  | 1.36                     |
| Bromine                   | 1.09                     |
| Iodine                    | 0.54                     |
The hydroxyl radicals are able to react with almost every type of organic compounds [27], leading, in some cases, to their complete oxidation into CO$_2$ and H$_2$O [10,25,26]. However, partial oxidation can be the main route, usually leading to more biodegradable products [25,26]. The oxidation of organic matter (RH) by hydroxyl radicals occurs by three mechanisms [19,42]: radical addition (Equation (1)), electron transfer to radicals (Equation (2)), and hydrogen abstraction (Equation (3)) that generates organic radicals, which yield peroxyl radicals by addition of molecular oxygen (Equation (4)).

\[
\begin{align*}
\text{HO}^\bullet + \text{RH} & \rightarrow \text{RH}_2\text{O} \tag{1} \\
\text{HO}^\bullet + \text{RH} & \rightarrow \text{RH}^{\bullet^+} + \text{OH}^- \tag{2} \\
\text{HO}^\bullet + \text{RH} & \rightarrow \text{R}^\bullet + \text{H}_2\text{O} \tag{3} \\
\text{R}^\bullet + \text{O}_2 & \rightarrow \text{RO}_2^\bullet \tag{4}
\end{align*}
\]

A large number of technologies are responsible for the generation of the hydroxyl radicals. Most of them use a combination of oxidants, such as ozone or and hydrogen peroxide, semiconductors (like titanium dioxide or zinc oxide) or catalysts (e.g., transition metal ions), and irradiation (ultraviolet and/or visible, sunlight or ultrasounds), as shown in Figure 1 [11,19,24,44]. Processes in which the catalyst is dissolved in the effluent are called homogeneous, but when the catalyst is supported on a solid matrix they are designated as heterogeneous.

Figure 1. Some relevant advanced oxidation processes (AOPs) for wastewater treatment (adapted from Poyatos et al. [24]).

The benefits of AOPs are: the possibility of degrading pollutants in a wide range of concentrations, the easiness in combining with other processes, such as biological and adsorption, and also the fact that some of them are conducted at near ambient pressure and temperature [9,44]. Nevertheless, each AOP has characteristic drawbacks associated. As an example, when using ozone-based processes, sophisticated equipment is required, such as an ozone generator, a cooling system, pre-and post-treatment setups to dry the air fed to the ozonator and to reduce the residual ozone in the gas off, respectively [45], which increases the implementation and operation costs. The processes that use radiation have high expenses of energy consumption, in addition to the costs of the installation and equipment. However, when effective photocatalysts are used, this charge can be null in countries with high incidence of solar radiation, which can replace artificial sources. Another disadvantage
of these processes is that the compounds present in the effluent can filtrate or absorb the radiation, limiting its reaction with the oxidant [46], which decreases the treatment efficiency.

The CWPO process is known for its simplicity, as it does not require any sophisticated equipment, involves safe and easy to handle reactants, has high efficiency, and low investment cost [47,48]. This review is focused on the wastewater treatment by CWPO using nano gold-based catalysts. The use of gold catalysts shown benefits as there is no loss of metal into solution and materials are stable and efficient, as will be further explained below.

3. Catalytic Wet Peroxidation

In the end of the 19th century, the CWPO process was firstly observed by H.J.H. Fenton, who described the highly oxidative properties of hydrogen peroxide in presence of iron ions during oxidation of tartaric acid [49]. Later, Haber and Weiss [27] discovered that the hydroxyl radical was the responsible for the degradation of the organic compounds. So, CWPO is based on the catalytic hydrogen peroxide decomposition by transition metallic cations (M) that generates hydroxyl radicals (see Equation (5)) in mild reaction conditions [27,50–53]. In this process, the catalyst is oxidized in the reaction with \( \text{H}_2\text{O}_2 \), generating \( \text{HO}^• \) (Equation (5)), being regenerated (reduced) with additional \( \text{H}_2\text{O}_2 \) molecules and even with the generated hydroperoxyl radicals (\( \text{HO}_2^• \)), according to Equations (6) and (7) [52–55].

\[
\begin{align*}
\text{M}^{n+} + \text{H}_2\text{O}_2 & \rightarrow \text{M}^{(n+1)+} + \text{HO}^• + \text{HO}^- \\
\text{M}^{(n+1)+} + \text{H}_2\text{O}_2 & \rightarrow \text{M}^{n+} + \text{HO}_2^• + \text{H}^+ \\
\text{M}^{(n+1)+} + \text{HO}_2^• & \rightarrow \text{M}^{n+} + \text{O}_2 + \text{H}^+
\end{align*}
\]

The hydroxyl radical has an extremely short life-time but is very reactive as it can react with the excess of catalyst (Equation (8)) or even oxidant (Equation (9)) [50,52–55], being such reactions the undesired scavenging of the hydroxyl radicals.

\[
\begin{align*}
\text{M}^{n+} + \text{HO}^• & \rightarrow \text{M}^{(n+1)+} + \text{HO}^- \\
\text{H}_2\text{O}_2 + \text{HO}^• & \rightarrow \text{HO}_2^• + \text{H}_2\text{O}
\end{align*}
\]

The main limitations of homogeneous CWPO are the following: (i) the narrow pH range (2 to 4) in which the pollutants degradation efficiency is maximum [52,55,56], and (ii) the need to recover the catalyst after treatment, in order to comply with environmental regulations, as shown by some authors [57]. A subsequent unit is required afterwards, in which the generated sludge, containing organic compounds as well as metals, has to be further treated, becoming the overall process more complex and expensive [48,52,54–56]. In order to overcome this challenge, several studies have been reported in literature dealing with supporting metals on solid porous matrices. By doing so, the metal is deposited on the support, becoming a heterogeneous catalyst, which is present in solution in a solid form, forming a slurry (batch reactors), being easily recovered; alternatively, it can be packed in a fixed bed reactor.

The principles of the heterogeneous process are very similar to the homogeneous; however, complexity increases due to the diffusion/adsorption phenomenon. It is widely accepted that hydrogen peroxide is adsorbed on the matrix pores, but this is not completely proved [58].

The main reactions of heterogeneous CWPO (Equations (10)–(14)) are the same as the homogeneous analogue, but with the addition of the support (X):

\[
\begin{align*}
\text{X-M}^{n+} + \text{H}_2\text{O}_2 & \rightarrow \text{X-M}^{(n+1)+} + \text{HO}^• + \text{HO}^- \\
\text{X-M}^{(n+1)+} + \text{H}_2\text{O}_2 & \rightarrow \text{X-M}^{n+} + \text{HO}_2^• + \text{H}^+ \\
\text{X-M}^{(n+1)+} + \text{HO}_2^• & \rightarrow \text{X-M}^{n+} + \text{O}_2 + \text{H}^+
\end{align*}
\]
The wastewater treatment by CWPO has been extensively studied for decades, mostly using supported iron as a catalyst [29,52,58–61] (in this case, the process being called Fenton or Fenton-like). Recent studies report on catalysts where iron has been replaced by other metals, such as nickel, cobalt, copper, cerium, and manganese, as well as bimetallic particles [47,51,62–65]. However, quite often, such materials are not stable, leaching the metal to the effluent, making their reuse not possible, and their industrial application unfeasible.

In order to overcome the problem of the catalysts lack of stability, some authors report on effluents treated by CWPO using gold catalysts supported on porous matrices. Although this might seem less economically attractive, given the price of gold, these catalysts present high stability, with negligible metal leaching, and are efficient in hydrogen peroxide consumption and pollutants degradation [66–75].

In the next sections, we will discuss some methods of preparation of nano gold-based catalysts and the treatment of effluents by CWPO catalyzed by gold.

4. Nano Gold-based Catalysts

For gold to be an active catalyst, its synthesis must be carefully made in order to obtain well dispersed nanoparticles on the support. This preparation process of the gold catalysts starts by obtaining colloidal gold in suspension, by reducing Au^{3+} to Au^{0} [76,77], using different reducing agents (such as alcohols, ascorbic and citric acid, amines, citrate, hydrazines, and toluene) [77,78]. As atomic gold is formed and its concentration increases, the solution becomes saturated and the metal gradually precipitates and forms nanoparticles. Nanoparticle formation is promoted by the addition of stabilizers [76,77], like amines, quaternary alkyl ammonium ions, phosphine, carboxyl acids, and thiols [77].

Usually, gold colloids are obtained by applying the Turkevich method [79], which consists on the reaction between AuCl_{4}^{-} (using tetrachloroauric acid (HAuCl_{4}) or sodium tetrachloroauric (NaAuCl_{4})) with sodium citrate as reducing agent and capping [80], resulting in gold nanoparticles with particle diameters of 10–12 nm [76,77]. However, for catalytic purposes, it is advantageous that gold particles have smaller sizes, between 2 and 10 nm [77]. This is achieved by reducing AuCl_{4}^{-} with a strong reducing agent, such as NaBH_{4} [77,78].

Another method for generating gold colloids was developed by Brust et al. [76]. It is based in the reaction of HAuCl_{4} solution with NaBH_{4} (reducing agent), in the presence of toluene and tetraoctylammonium bromide (TOAB) which acts as a transfer cation, stabilizing agent, and anti-coagulant. First, the migration of AuCl_{4}^{-} from water to the organic phase (toluene) takes place, by ion metathesis of the counter anion on the phase transfer agent. Then, addition of sodium borohydride promotes the precursor reduction to metallic gold. This method produces gold nanoparticles with particle sizes between 2 to 6 nm [76,77].

The gold colloids can also be formed by dissolving AuCl_{4}^{-} in a solvent (like benzyl alcohol or ethylene glycol [78]), other than water. Reduction occurs by thermal treatment or addition of reducing agents [77].

The catalysts preparation is finalized by deposition of gold on a support (normally a metal oxide or a carbon material). Both processes (colloid formation and deposition on the support) can occur simultaneously and can be achieved by using several methods, namely deposition/precipitation (DP), co-precipitation (CP), impregnation, vapor-phase deposition, grafting, sol-gel, ion-exchange, among others [80–83]. These most common techniques are described below.

4.1. Deposition/Precipitation

The DP method is one of the most widely used for gold catalysts preparation. This procedure was first described by Haruta [81], who adjusted the pH of HAuCl_{4} solution in the range 6–10 with NaOH,
then added the metal oxide used as support [80], and readjusted the pH. The suspension was stirred during 1 h for gold precipitation in the form of \( \text{Au(OH)}_3 \), that was deposited on the metal oxide surface. Finally, after the deposition step, the solid in suspension was recovered, washed, dried, and submitted to a thermal treatment at 250 °C in air atmosphere [81], or at 300 °C in hydrogen atmosphere [77], for gold reduction.

Haruta [81] referred the influence of pH on gold particle size. For pH about 6, the \( \text{AuCl}_4^- \) is transformed into \( [\text{Au(OH)}_n\text{Cl}_{4-n}]^- (n = 1 \text{ to } 3) \) and the mean size of gold particles size is less than 4 nm. For pH in the range 7 to 8, the n value is close to 3, which is preferable for the preparation of the gold catalysts, depending on the support. At lower pH, the hydrolysis of the Au-Cl bond takes place in a smaller extent. Moreover, for values of pH below the oxide isoelectric point, its surface is positively charged and consequently adsorbs more negative charged gold species. This results in a higher concentration of chloride on the surface, which promotes high mobility of gold, leading to the formation of larger particles [80]. For pH values above the isoelectric point of the support, the adsorption of negatively charged gold species decreases drastically. Consequently, the gold loading is lower, and so is the chloride concentration of Au being formed [80]. Figure 2a shows a HR-TEM image and histogram of gold nanoparticle size distribution of a catalyst prepared at pH 9 by deposition/precipitation. This procedure is reproducible, very reliable, and the obtained catalysts show high catalytic activity.

![HR-TEM image and histogram of gold nanoparticle size distribution](image)

**Figure 2.** -HR-TEM image and histogram of gold nanoparticle size distribution of Au/\( \alpha \)-Fe\(_2\)O\(_3\) catalyst (a) prepared at pH 9.0 by deposition/precipitation (DP) with NaOH and heating at 70°C and -HR-TEM image of an Au/\( \alpha \)-Fe\(_2\)O\(_3\) catalyst (b) prepared by the co-precipitation method calcined at 400 °C. Nanogold particles are seen as dark spots. Adapted from Rodrigues et al. [67] and Hodge et al. [84], respectively.
4.2. Co-Precipitation

This method is based on simultaneous CP of hydroxide or carbonate and gold. For that, the gold precursor (HAuCl₄) and the soluble metal salt precursor (preferably a nitrate) are added to a Na₂CO₃ (and/or NH₄OH) solution and the suspension is mixed for a few minutes. After 1 h aging, the precipitates (Au and metal oxide) are washed and filtrated for five consecutive cycles, then dried overnight, and finally calcined in air atmosphere, to obtain a powder material [80–82,85]. This method differs from DP, in the sense that both oxide and gold are co-precipitated at the same time (in DP, Au is deposited on the already prepared support).

The preparation of catalysts by CP needs a concentration of metal salt around 0.1-0.4 M, pH range of 7–10 and temperature of precipitation and calcination between 47–87 and 227–397 °C. Within these conditions, a homogeneous dispersion of gold nanoparticles can be obtained [82].

CP is the most useful and simple method; however, its applicability is limited, as only metal hydroxides or carbonates can be co-precipitated with Au(OH)₃; moreover, reducible supports (α-Fe₂O₃, CO₃O₄, NiO, and ZnO) have to be employed in order to obtain a good dispersion of the gold nanoparticles [80,82]. Figure 2b shows a HR-TEM image of a Au catalyst prepared by this method.

4.3. Impregnation Method

This method consists in impregnating the support with a gold salt solution. This may be done by suspending the support on a large volume of metal salt, from which the solvent is removed, or by filling the pores of the support with the solution (this later procedure being called incipient wetness impregnation). Then, the precursor is dried and calcined at temperatures as high as 800 °C and reduced with hydrogen atmosphere at 120–250 °C, or aqueous oxalic acid at 40 °C, or aqueous magnesium citrate [80,83].

In the preparation of gold catalysts by this method, usually chloroauric acid (HAuCl₄·3H₂O) or auric chloride (AuCl₃ or Au₂Cl₆) are used as metal precursors. However, complex salts such as potassium aurocyanide (K₂Au(CN)₂) and the ethylenediamine complex [Au(en)₂]Cl₂ may also be employed. Regarding the supports, silica, alumina, and magnesia are often used, but titanium oxide, boehmite (Al(OH)), magnesium hydroxide, or ferric oxide (α-Fe₂O₃) have also been employed [83]. Figure 3a shows a HR-TEM image of a Au/Al₂O₃ catalyst prepared by incipient wetness impregnation.

Although impregnation is a classical procedure in the preparation of platinum group metal catalysts, it is not often applicable to gold, since the obtained catalysts show larger gold particle sizes when compared to materials prepared by CP or DP techniques. Moreover, they show low catalytic activity and it is difficult to obtain a good dispersion of the gold on the metal oxides. On one hand, gold has less affinity for these supports and lower melting point (1063 °C) than those of Pd (1550 °C) or Pt (1769 °C). On the other hand, during calcination of the precursor at low temperature (below 600 °C) the crystals of HAuCl₄ are dispersed on the surface of the support and the chloride ion markedly enhances the coagulation of gold particles [80–82,86].
4.4. Vapor-phase Deposition and Grafting Methods

The procedures for preparation of gold-based catalysts by these two methods are similar, the only difference being in the use (or not) of solvent. In the vapor-phase deposition method (also called chemical vapor deposition), a vapor of an organic gold compound (such as dimethyl-gold(III)-acetyl diketone, dimethyl-gold(III) β-diketone, or gold acetylacetonate) is transported onto a high area support by an inert gas stream and chemically reacts with the support surface to form a precursor of gold. The organic gold compound adsorbed on the support is pyrolyzed in air atmosphere, to be decomposed into small gold particles [80,81,83]. This method can be applied to a variety of metal oxides, including acidic supports, like silica oxide [80,81,83]. Figure 3b presents a HR-TEM image and histogram of particle size distribution of a gold catalyst supported on carbon nanotubes prepared by the vapor-phase deposition method.

In the grafting method, a gold complex ([Au(PPh₃)]NO₃ and/or [Au₆(PPh₃)₆][NO₃]₃) in solution is grafted onto the surface of a number of precipitated wet hydroxides (manganese and cobaltous hydroxides being particularly effective), which have many OH groups at the surface, which react with gold [80,83]. Then, drying in vacuum at room temperature and temperature-programmed calcination in air atmosphere are carried out, in order to cause a simultaneous transformation of the precursor to gold particles and oxides [80]. The deposition of gold on activated carbon is only achieved with the grafting method, however, the gold particles have too large diameters, around 10 nm [82], which leads to an inferior catalytic activity.

4.5. Sol-Gel Method

According to several authors, in the sol-gel method a sol solution of the support is obtained by mixing the support precursor (like tetra-ethyl-ortho-silicate, aluminum tri-sec-butoxide, aluminum isopropanoxide, or tetrabutoxy-titanium) with water, ethanol and methanol, and/or nitric acid. Then, the gold precursor (such as chloroauric acid, gold acetate, or hydrogen tetranitroaurate) is added to the sol solution of the support, stirring vigorously for a variable time until the gel begins to be formed. The obtained gel is dried during 12–24 h at a temperature about 100–200 °C and a calcination step follows [80].

The catalysts prepared by this method show gold nanoparticles with sizes below 6 nm, involving materials resulting from soluble precursors which form three-dimensional networks with the addition
of a base [80]. In Figure 4a it is possible to see a TEM image and gold particles size distribution of gold nanoparticles support on activated carbon prepared by the sol immobilization method.

![TEM image and particle size distribution of Au/C](image1)

**Figure 4.** (a) TEM image and particle size distribution of Au/C prepared by the sol immobilization method (dried at 60 °C until total evaporation of solution) and (b) TEM image of Au/Y prepared by ion-exchange at 25 °C and pH = 5.0. Gold nanoparticles are seen as dark spots. Adapted from Quintanilla et al. [68] and Lin et al. [89], respectively.

### 4.6. Ion-Exchange Method

The ion-exchange method consists in replacing the protons or other cations on surface, or within the structure of the support, by gold, and this leads first to atomically dispersed species and then, after calcination and reduction with hydrogen, to small gold particles [80,83].

This method is especially effective for depositing gold on zeolites, but the introduction of active species into the cavities of these supports, instead of placing gold on their external surface, presents several difficulties, for example, only limited cations or cationic complexes can be used ([Au(en)$_2$]$^{3+}$ – en = ethylenediamine - or [Au(NH$_3$)$_2$]$^+$) [80,83]. For this reason, the ion-exchange method is rarely used in the preparation of gold catalysts, although small metal particles are obtained.

Figure 4b shows a TEM image of gold supported on zeolite (Au/Y) catalyst prepared by the ion-exchange method.

In the next sections, we will present the treatment of wastewater by wet peroxidation using gold catalysts. The influence of catalyst properties and operating conditions, as well the catalyst stability, will be discussed.
5. Application of CWPO using Gold Catalysts in Wastewater Treatment

In the last century, there was an increasing interest in the use of gold catalysts by the scientific community. These materials have been used in chemical and environmental catalysis, in reactions such as CO oxidation [85,90,91], hydrogenation [92], water-gas-shift [93–95], combustion of volatile organic compounds [96–98], and organic compounds reduction [99] or oxidation [86,100–102]. Gold catalysts have also been used in wastewater treatment by catalytic wet peroxidation, which is the focus of this review, as said above. Gold has replaced catalysts that, although being efficient in the removal of pollutants, present the disadvantage of high metal leaching, like iron-based catalysts [69,70,103,104]. Gold does not leach, is stable and efficient, as mentioned above and will be further discussed ahead.

The preparation methods previously described influence the gold particles size and, consequently, the dispersion of the metal on the surface of the support. These two parameters are correlated as demonstrated by Equation (15) and have a strong effect on the catalytic activity, as well as other properties of gold-based catalysts like pore size, surface area ($S_{BET}$), mesoporosity of the support, and the oxidative state of gold.

Some authors correlate the effect of the gold amount and dispersion (and indirectly the particle size) with the catalytic performance, by evaluating the turnover frequency (TOF), which provides the number of molecules of target substrate degraded per gold atom and time unit (Equation (17)).

\[ DM \, (\%) = \frac{6 \times n_s \times MM \times 1000}{\rho \times N \times dp} \times 100 \]  

(15)

\[ n \, (\text{moles of gold}) = \frac{Y_{Au} \times W_{cat}}{MM} \]  

(16)

\[ \text{TOF} \,(\text{h}^{-1}) = \frac{C}{DM \times n \times t} \]  

(17)

where: $C$ refers to the molecules of substrate degraded, $DM$ is the gold dispersion, $n$ is the number of moles of gold used, $t$ is the time of reaction, $n_s$ is the number of atoms at the surface per unit area ($1.15 \times 10^{19} \text{ m}^{-2}$ for Au) [105], $MM$ is the molar mass of gold (196.97 g/mol), $\rho$ is the density of gold (19.5 g/cm$^3$), $N$ is the Avogadro’s number ($6.022 \times 10^{23}$ mol$^{-1}$), $dp$ is the average gold particle size (nm), $Y_{Au}$ is the amount of gold in the catalyst (wt.%), and $W_{cat}$ is the mass of catalyst (g).

The efforts to achieve active and stable gold catalysts to be used in the treatment of effluents by CWPO, in view of industrial applications, have been reported in literature. Table 2 presents an outlook of the research made, showing which pollutants were degraded, the wastewater treatment conditions used, catalysts employed, and efficiency reached.

The efficiency of the catalytic wet peroxidation process for wastewater treatment is influenced by the catalyst properties and many operating conditions, such as gold loading, pH, temperature, hydrogen peroxide dose, catalyst concentration, and also the radiation intensity source (the latter in the case of the photo-assisted wet peroxidation). The influence of the catalyst properties and the effect of such operating conditions, as well as the stability of these catalysts, will be briefly described below.
Table 2. Gold catalyst used in CWPO of model compounds degradation or wastewater treatment, operational conditions, and performances reached.

| Model Compound/ Effluent | Catalyst | Operation Conditions | Efficiency of CWPO | Ref. |
|--------------------------|----------|----------------------|--------------------|------|
| Orange II (OII) dye      | Au/Al₂O₃ (0.7 wt.%) | pH = 3.0; T = 50 °C; [H₂O₂] = 6 mM; [catalyst] = 2.0 g/L; [OII] = 0.1 mM; t = 4 h | Dye removal = 98.9%; COD removal = 49.8%; COD consumption = 95.0%; Specific Oxygen Uptake Rate = 27.8 mgO₂/(gAu.h); Inhibition of *Vibrio Fischeri* = 0.0%; Gold leaching < 0.04% | [66] |
| Acrylic Dyeing Wastewater | Au/Al₂O₃ (0.7 wt.%) | pH = 3.0; T = 50 °C; [H₂O₂] = 3.52 g/L; [catalyst] = 2.0 g/L; [OII] = 0.1 mM; t = 4 h | Color removal = 34.4%; TOC removal = 42.9%; COD removal = 50.5%; H₂O₂ consumption = 98.8%; BOD₅/COD = 0.23; Gold leaching < 0.04% | |
|                          | Au/Al₂O₃ (0.7 wt.%) | pH = 3.0; T = 30 °C; [H₂O₂] = 6 mM; [catalyst] = 2.0 g/L; [OII] = 0.1 mM; t = 16 h | Dye removal = 40.9%; TOC removal = 29.6%; Gold leaching < 0.04% | [67] |
|                          | Au/Fe₂O₃ (4.0 wt.%) from WGC | pH = 3.0; T = 50 °C; [H₂O₂] = 6 mM; [catalyst] = 2.0 g/L; [OII] = 0.1 mM; t = 4 h | Dye removal = 68.9%; TOC removal = 32.4%; H₂O₂ consumption = 91.7%; Gold leaching < 0.04% | [67] |
|                          | Au/Al₂O₃ (0.7 wt.%) | pH = 3.0; T = 30 °C; [H₂O₂] = 6 mM; [catalyst] = 2.0 g/L; [OII] = 0.1 mM; t = 16 h | Dye removal = 68.9%; TOC removal = 32.4%; H₂O₂ consumption = 91.7%; Gold leaching < 0.04% | [67] |
|                          | Au/TiO₂ (1.6 wt.%) | pH = 3.0; T = 30 °C; [H₂O₂] = 6 mM; [catalyst] = 2.0 g/L; [OII] = 0.1 mM; t = 16 h | Dye removal = 68.9%; TOC removal = 32.4%; H₂O₂ consumption = 91.7%; Gold leaching < 0.04% | [67] |
|                          | Au/ZnO (1.2 wt.%) | pH = 3.0; T = 30 °C; [H₂O₂] = 6 mM; [catalyst] = 2.0 g/L; [OII] = 0.1 mM; t = 16 h | Dye removal = 68.9%; TOC removal = 32.4%; H₂O₂ consumption = 91.7%; Gold leaching < 0.04% | [67] |
| Phenol                   | Au/TiO₂ (0.8 wt.%) | pH = 3.0; T = 50 °C; [H₂O₂] = 5 mL; [catalyst] = 2.7 g/L; V₀ = 5 mL; t = 24 h; V_solution = 45 mL | TOFphenol = 0.07*10⁶ (h⁻¹); TOFCOD = 0.07*10⁶ (h⁻¹); TOFH₂O₂ = 2.52*10⁶ (h⁻¹) | [68] |
|                          | Au(3)/C (0.5 wt.%) | pH = 3.0; T = 50 °C; [H₂O₂] = 5 mL; [catalyst] = 2.7 g/L; V₀ = 5 mL; t = 24 h; V_solution = 45 mL | TOFphenol = 0.11*10⁶ (h⁻¹); TOFCOD = 0.10*10⁶ (h⁻¹); TOFH₂O₂ = 1.84*10⁶ (h⁻¹) | [68] |
|                          | Au(5)/C (0.5 wt.%) | pH = 3.0; T = 50 °C; [H₂O₂] = 5 mL; [catalyst] = 2.7 g/L; V₀ = 5 mL; t = 24 h; V_solution = 45 mL | TOFphenol = 0.32*10⁶ (h⁻¹); TOFCOD = 0.25*10⁶ (h⁻¹); TOFH₂O₂ = 4.07*10⁶ (h⁻¹) | [68] |
|                          | Au(7)/C (0.5 wt.%) | pH = 3.0; T = 50 °C; [H₂O₂] = 5 mL; [catalyst] = 2.7 g/L; V₀ = 5 mL; t = 24 h; V_solution = 45 mL | TOFphenol = 0.25*10⁶ (h⁻¹); TOFCOD = 0.25*10⁶ (h⁻¹); TOFH₂O₂ = 2.10*10⁶ (h⁻¹) | [68] |
|                          | Au(10)/C (0.5 wt.%) | pH = 3.0; T = 50 °C; [H₂O₂] = 5 mL; [catalyst] = 2.7 g/L; V₀ = 5 mL; t = 24 h; V_solution = 45 mL | TOFphenol = 0.47*10⁶ (h⁻¹); TOFCOD = 0.43*10⁶ (h⁻¹); TOFH₂O₂ = 1.84*10⁶ (h⁻¹) | [68] |
| Phenol                   | Au/Hap (2.4 wt.% of Au) | pH = 2.0; T = 70 °C; V₀ = 5 mL; [catalyst] = 0.1 g/L; [phenol] = 100 mg/L; t = 2 h | Phenol removal = 92.5% | [70] |
Table 2. Cont.

| Model Compound/Effluent | Catalyst | Operation Conditions | Efficiency of CWPO | Ref. |
|-------------------------|----------|----------------------|--------------------|------|
| Phenol                  | Au/TiO₂ – AD (2.8 wt.%) | [phenol] = 200 mg/L; [H₂O₂] = 1520 mg/L; pH = 2.5; T = 80 °C; P = 1 atm; LHSV = 3.8 h⁻¹ | Phenol removal steady-state = 100.0%; TOC removal steady-state = ~65.0% | [71] |
| Phenol                  | Au/TiO₂ – AD (3.2 wt.%) | | Phenol removal steady-state = 100.0%; TOC removal steady-state = ~80.0% | |
| Phenol                  | Au/AC (0.8 wt.% of Au) | pH = 3.5; T = 80 °C; [H₂O₂] = 25 g/L; [catalyst] = 2.5 g/L; [phenol] = 5 g/L; t = 22 h | Phenol removal = 100%; TOC removal = 70% | [72] |
| Phenol                  | Au/DNP (1 wt.% of Au) | pH = 4.0; T = 50 °C; [H₂O₂] = 1.44 g/L; [catalyst] = 320 mg/L; [phenol] = 1 g/L; t = 7 h | Phenol removal = 100%; H₂O₂ consumption = 100%; BOD₅/COD = 0.72 | [73] |
| Phenol                  | Au/CeO₂ (1.0%) | pH = 4.0; Room temperature; [H₂O₂] = 200 mg/L; [Au] = 0.0025 mM; [phenol] = 100 mg/L; t = 24 h | Phenol removal = 7.0%; H₂O₂ consumption = 88.0%; Gold leaching = 0.8% | |
| Phenol                  | Au/Fe₂O₃ (1.5%) | | Phenol removal = 3.0%; H₂O₂ consumption = 8.0%; Gold leaching = 0.7% | |
| Phenol                  | Au/TiO₂ (1.5%) | | Phenol removal = 3.0%; H₂O₂ consumption = 19.0%; Gold leaching = 0.5% | [74] |
| Phenol                  | Au/C (0.8%) | | Phenol removal = 7.0%; H₂O₂ consumption = 14.0%; Gold leaching = 5.8% | |
| Phenol                  | Au/nDp (< 1.0%) | | Phenol removal < 1.0%; H₂O₂ consumption = 6.0%; Gold leaching = 0.5% | |
| Phenol                  | Au/HO-nDp (1.0%) | | Phenol removal = 93.0%; H₂O₂ consumption = 48.0%; Gold leaching = 0.7% | |
| Methyl Blue dye (MB)    | Au/CNT (41.0 wt.%) | [MB dye] = 50 mg/L; [catalyst] = 0.5 g/L; [H₂O₂] = 500 mM; pH = 7.08; t = 120 min | MB removal = ~100% | [88] |
| 1,1-diphenyl-2-picrylhydrazyl (DPPH) | Au/CNT placed in water/cyclohexane mixture (1/10 v/v) (41.0 wt.%) | [DPPH] = 0.2 mM; [catalyst] = 1 g/L; [H₂O₂] = 250 mM; t = 10 min; Room temperature; W/O = 1:10 v/v | DPPH removal = 100% | |
| Acid Orange 7 (AO7) dye | Au/CeO₂ (1 wt.% of Au) | [H₂O₂] = 20 mM; [catalyst] = 0.5 g/L; [dye] = 35 mg/L; t = 33 h | AO7 removal = 80% | [106] |
**Table 2. Cont.**

| Model Compound/Effluent | Catalyst | Operation Conditions | Efficiency of CWPO | Ref. |
|-------------------------|----------|----------------------|--------------------|------|
| Methyl Orange dye (MO)  | Au/TN (1.0 wt.%) | [MO] = 50 mg/L; [catalyst] = 2 g/L; [H₂O₂] = 0.15 M; pH = 3.0; T = 80 °C; t = 240 min | MO removal = 85%; TOC removal = 83% | [107] |
|                         | Au/SRAC (3.0 wt.%) | | BPA removal = 89.0%; H₂O₂ consumption = 44.1%; | |
|                         | Au/PSAC (3.0 wt.%) | | BPA removal = 23.8%; H₂O₂ consumption = 8.3%; | |
|                         | Au/CNF (3.0 wt.%) | | BPA removal = 20.4%; H₂O₂ consumption = 14.5%; | |
|                         | Au/FPDU-15 (3.0 wt.%) | [BPA] = 114 mg/L; [catalyst] = 125 mg/L; [H₂O₂] = 530 mg/L; pH = 3.0; T = 30 °C | BPA removal = 32.4%; H₂O₂ consumption = 22.8%; | [106] |
|                         | Au/X40s (10.0 wt.%) | | BPA removal = 14.5%; H₂O₂ consumption = 10.7%; | |
|                         | Au/Fe₂O₃ (5.0 wt.%) | | BPA removal = 10.1%; H₂O₂ consumption = 7.6%; | |
|                         | Au/TiO₂ (1.5 wt.%) | | BPA removal = 5.3%; H₂O₂ consumption = 10.8%; | |
|                         | Au-Fe₂O₃/Al₂O₃ (0.5 wt.%) | | BPA removal = 6.6%; H₂O₂ consumption = 15.3%; | |
| Bisphenol A (BPA)      | Au/SRAC (1.5 wt.%) | [BPA] = 89 mg/L; [catalyst] = 125 mg/L; [H₂O₂] = 530 mg/L; pH = 3.0; T = 30 °C | BPA removal = −80.0%; H₂O₂ consumption = −40.0% | |

* TOF = turnover frequency
5.1. Influence of the Catalyst Properties

As already mentioned above, the catalytic properties directly affect the efficiency of the CPWO process. Ge et al. [106] concluded that the textural properties influenced AO7 dye removal, and achieved 80% for Au/CeO$_2$ with a lower S$_{BET}$ (55 m$^2$/g) and an intermediate gold content (1 wt.%).

Alvaro et al. [109] concluded that the morphological properties of gold supported on mesoporous titania had an influence in the decontamination of Soman wastewater. The authors reached the best decontamination degree (~100%) for a catalyst with medium surface area (90 m$^2$/g) and large pore diameter (7.1 nm) associated to a highest gold loading (0.70 wt.%), among the studied samples.

Navalon et al. [74] evaluated phenol oxidation by CWPO using gold supported on CeO$_2$, TiO$_2$, carbon, Fe$_2$O$_3$, npD, and HO-npD, and observed that catalysts with smaller gold particle size (< 1 nm) and intermediate gold loading (1.0%) led to the highest performance (total phenol disappearance and 48.0% of hydrogen peroxide consumption). Moreover, a small amount of gold (0.7%) was leached from the support to the solution.

However, in the three studies reported above, the authors do not indicate any explanation why the efficiency of CPWO was the best for the catalysts selected. The main characteristics influencing the catalysts performance in CPWO are morphology and porosity (adsorption capacity), gold loading, and particle size.

The optimization of the gold loading of a catalyst is essential for economic aspects, as mentioned above, and is determinant for catalysing the reaction that generates hydroxyl radicals, influencing the efficiency of CWPO. In the work of Rodrigues et al. [67], the efficiency of CWPO decayed dramatically when the loading of gold on iron oxide increased from 0.8 to 4.0 wt.%, reducing from 99.7 to 36.6% and from 75.8 to 24.0% for OII dye and TOC removals, respectively. Moreover, a significant reduction was observed in the production of hydroxyl radicals with increasing gold content.

However, in a work using CWPO assisted with radiation to treat an OII dye solution with gold on iron oxide, the gold content had no effect on the colour removal [110]. This is due to the fact that the dye is degraded in the presence of UV/vis radiation alone. However, the authors observed a slight reduction in mineralization from 68.2 to 58.4% when the gold content increased, as well as in the production of hydroxyl radicals [110].

In both studies mentioned above, the authors pointed out an explanation for the decay of the process performance with the increase of the gold content; if in excess, gold reacts with the hydroxyl radical (HO$^·$ + Fe$_2$O$_3$-Au$^0$ → Fe$_2$O$_3$-Au$^+$ + HO$^-$), being less available to oxidize the dye, and the reaction by-products [66,110].

The degradation of methyl orange decreased from ~40 to ~10% when the gold loading increased from 1.0 to 4.0 wt.%, in a study dealing with the removal of this dye by CWPO, at 25 °C, using gold supported on modified titanium nanotubes [107]. The authors attributed this decreased of process efficiency to the fact that the catalyst with lower gold content had a smaller particle size and gold was uniformly distributed on the surface of the support [107].

On the other hand, and as shown in Figure 5a, the degradation of AO7 dye increased with gold loading until 1 wt.%; however, a further increase of catalyst content to 2 wt.% impaired the oxidation of the dye by CWPO, using gold on cerium oxide as catalyst [106], increasing the ratio between the AO7 concentration after 30 h of reaction (C) and the initial AO7 concentration ($C_0$), $C/C_0$, from 0.2 to 0.4. Furthermore, the combination of CWPO with visible radiation showed an optimum for 1 wt.% of gold loading for shorter reaction times (less than 5 h), more notorious in the period from 2 to 5 h (see Figure 5b) [106].

In contrast with the above-mentioned studies that showed the existence of an optimal gold loading, Yang et al. [108] observed that the removal of bisphenol A and consumption of hydrogen peroxide increased from 21.1 and 9% to 89.0 and 45%, respectively, after 12 h of reaction, when the gold content in an Au/AC catalyst increased from 0 to 3 wt.%. Moreover, Sempere et al. [75] observed a decrease in the turnover frequency (calculated according to Equations (15)–(17)) of phenol and hydrogen peroxide from ~100 and 320 h$^{-1}$ to ~70 and 180 h$^{-1}$, respectively, when the gold loading increased from 0.1
to 0.5 wt.% in Au/FH₂, and subsequent annealing treatment with hydrogen for sunlight assisted CWPO. The authors pointed out an explanation for the decay of the catalytic activity with increasing particle size, i.e., the catalyst having a lower gold content had a smaller particle diameter and showed more efficiency.

The studies described above allow us to conclude that the effect of the gold loading in the efficiency of CPPO depends on the type of catalyst used and the compound/wastewater to be treated.

Quintanilla et al. [68] evaluated the degradation of phenol, the mineralization and H₂O₂ consumption by wet peroxidation using Au/C, Au/Fe₂O₃, and Au/TiO₂. They verified that activated carbon is the preferable support because it has a higher adsorption capacity. Also, Au/C with less amount of gold (0.13 wt.%), lower gold size (5.1 nm), and higher fraction of Au⁶⁺ in the catalyst surface (31%), presented the highest TOF for phenol oxidation (1.19 × 10⁴ h⁻¹), TOC reduction (1.08 × 10⁴ h⁻¹), and H₂O₂ consumption (16.70 × 10⁴ h⁻¹), see Table 3. This catalyst (Au(3)/C) had gold particles with about 5 nm size, which is beneficial for the catalytic performance.

Rodrigues et al. [67] tested gold supported on titanium, zinc, aluminum, and iron oxides to treat a dye solution by CWPO. They concluded that the Au/Al₂O₃ catalyst with higher surface area (S_BET = 210 m²/g), lower amount of gold (0.7 wt.%), and an intermediate gold particle size (3.6 nm), had the best performance (removal of dye and TOC of ca. 98% and 47%, respectively, consumption of

**Figure 5.** C/C₀ ratio during AO7 degradation by CWPO (a) and assisted with visible radiation (b) processes at different gold loadings ([AO7] = 35 mg/L, [H₂O₂] = 20 mM and [Au/CeO₂] = 0.5 g/L). Adapted from Ge et al. [106].

### Table 3. Gold loading, particle size and percentage of exposed surface gold species of the catalysts, and turnover frequency (TOF) values of phenol, TOC oxidation, and hydrogen peroxide decomposition (experimental conditions: [Phenol] = 5 g/L, [Catalyst] = 2.7 g/L, V₁₂O₂ = 5 mL, t = 24 h and V_solution = 45 mL). Adapted from Quintanilla et al. [68].

| Catalyst       | [Au]_total (wt.%) | Au⁰ Fraction (%) | Au⁶⁺ Fraction (%) | Au Size (nm) | Phenol TOF x 10⁻⁴ (h⁻¹) | TOC TOF x 10⁻⁴ (h⁻¹) | H₂O₂ TOF x 10⁻⁴ (h⁻¹) |
|----------------|------------------|------------------|------------------|--------------|--------------------------|-----------------------|------------------------|
| Au/TiO₂        | 0.80             | 79               | 21               | 3.1 ± 1.8    | 0.07                     | 0.07                  | 2.52                   |
| Au(3)/AC⁺      | 0.13             | 69               | 31               | 5.1 ± 2.0    | 1.19                     | 1.08                  | 16.70                  |
| Au(5)/AC⁺      | 0.47             | 72               | 28               | 4.9 ± 1.0    | 0.32                     | 0.25                  | 4.07                   |
| Au(7)/AC⁺      | 0.48             | 71               | 29               | 6.8 ± 1.7    | 0.25                     | 0.25                  | 2.27                   |
| Au(10)/AC⁺     | 0.50             | 69               | 31               | 9.1 ± 1.1    | 0.47                     | 0.43                  | 1.87                   |

* The numbers correspond to the initial average size (nm) of gold in the colloidal solution used for the catalyst preparation.
H₂O₂ ~96%, and higher hydroxyl radicals generation) and the highest TOF value for OII dye removal (75.5 × 10⁻⁶ s⁻¹). The best efficiency of CPWO for Au/Al₂O₃ was associated to its higher adsorption capacity. The same was concluded for these same catalysts when light-assisted CWPO was used for OII dye degradation [110]. Au/Al₂O₃ permitted total discoloration and TOC removal of about 80%, combining further formation of hydroxyl radicals [110]. In the same way, Drašinac et al. [107] observed that the morphological properties of the catalysts and gold properties played an important role in methyl orange (MO) dye degradation. These authors reached the best CPWO performance (removals of 83 and 85% for TOC and MO dye, respectively) with gold supported on modified titanium nanotubes with a gold nanoparticle size of 7 nm and 1.1 wt.% loading, which had the highest total pore volume (1.31 cm³/g), pore diameter (14.8 nm), and surface area (335 m²/g). The authors reported that, in addition to the low gold content in the catalyst, the smaller particle diameter and the uniform distribution on the surface of the support benefit the catalytic process.

Yang et al. [108] reached the best performance of CWPO (89.0 and 44.1% for bisphenol A (BPA) removal and oxidant conversion, respectively) using a gold supported on styrene-based activated carbon (Au/SRAC) catalyst, which had an intermediate gold nanoparticle size (4.4 nm) and loading (3.0 wt.%). The authors pointed out the small gold size of the material as being beneficial in the degradation of the compound and oxidant conversion. On the other hand, Han et al. [70] reached the maximum removal of phenol (82%) and space-time conversion (0.53 mmol h⁻¹ L⁻¹) when using a gold supported on hydroxyapatite (Au/Hap) catalyst with higher gold particle size (4.9 nm) and intermediate loading (2.4 wt.%). The authors stated that the best catalytic activity of this sample was due to the gold particle size close to 5 nm.

These studies showed that the textural properties of the catalysts, as well as the gold particle size, play an important role in the efficiency of the catalytic process.

5.2. Effect of the Operating Conditions

The efficiency of the catalytic wet peroxidation process for wastewater treatment is also influenced by many operating conditions, such as catalyst dose, pH, temperature, hydrogen peroxide concentration, and also the radiation intensity source (the latter in the case of the photo-assisted wet peroxidation). The effect of such operating conditions will be briefly described below.

5.2.1. Catalyst Dose

The efficiency of the process increases with the catalyst concentration, since more gold will be available in the reaction medium to catalyze CWPO, generating more hydroxyl radicals. However, above a certain concentration of catalyst there is very often a negative effect, once scavenging of hydroxyl radicals by the excess of gold occurs (Equation (13)). The ideal concentration of catalyst depends on the type of effluent to treat, being necessary to optimize it. In the work developed by Domínguez et al. [72], a linear increasing dependence was obtained for initial reaction rates of phenol and oxidant disappearance for a Au/AC concentration in the range of 0–6 g/L.

As reported by Domínguez et al. [72], the work of Martín et al. [73] showed that an increase in the Au/DNP catalyst concentration from 50 to 320 mg/L proportionately increased the initial reaction rate of phenol degradation and oxidant consumption (see Figure 6). The same tendency was observed by Navalon et al. [111], as the initial phenol degradation and H₂O₂ decomposition rates increased for Au/HO-npD concentrations, in the range of 0 to 400 mg/L, for solar light assisted CWPO.

On the other hand, Rodrigues et al. [66] reached an optimum dose of Au/Al₂O₃ catalyst of 2.0 g/L that maximized the OII degradation and mineralization, as well as the formation of hydroxyl radicals (see Figure 7) when CWPO was applied to the dye solution. Moreover, the oxidant decomposition increased with catalyst dose until 2.0 g/L, but remained equal for the highest concentration tested (see Figure 7a). No gold leaching was found in any of the tests.

5.2.2. Hydrogen Peroxide Concentration

The initial concentration of \( \text{H}_2\text{O}_2 \) also plays a very important role in the oxidation of organic compounds in CWPO processes and in the operating costs of such treatment procedures; thus, it is necessary to determine the optimum dose of this reagent.
The improvement of the process by the addition of H₂O₂ is mostly due to the increased production of hydroxyl radicals, as described in Equations (10), (11), and (20). However, at high peroxide concentrations, the reaction between excess H₂O₂ and the strong oxidant HO• species becomes more relevant and, as a consequence, no subsequent improvement on the heterogeneous CWPO rate can be noticed, as the produced HO₂• radicals are less reactive than the HO• radicals (Equation (14)) [112]. Contrarily, if the concentration is low, the oxidation degree is small and there is the possibility of formation of unwanted intermediate products, which, in most cases, are more toxic and less biodegradable than the original compounds. Inherently, it is common to observe the existence of an optimum oxidant (hydrogen peroxide) dose in either wet peroxidation or radiation-assisted wet peroxidation processes.

The existence of an optimum oxidant dose was reported by several authors for CWPO catalyzed by gold on different supports [66,88,108,110,111]. In the work developed by Rodrigues et al. [66], the effect of this parameter was evaluated in the range of 3.0 to 12.0 mM. An increase in the removal of dye and TOC was observed, as well as in the formation of hydroxyl radicals with hydrogen peroxide concentration until 6.0 mM, but the efficiency of the process was reduced for higher oxidant doses (see Figure 8). The consumption of H₂O₂ also increased until 6 mM and remained constant for the higher doses. The authors reached 46.6 and 97.8% of TOC and dye removals, respectively, and ~100% for hydrogen peroxide consumption after 4 h of reaction with the optimized oxidant dose.

![Figure 8. Effect of hydrogen peroxide concentration in dye and TOC removals and hydrogen peroxide consumption after 4 h of reaction (a) and hydroxyl radicals formation during CWPO (b) for a Au/Al₂O₃ catalyst ([OII]₀ = 0.1 mM or [OII]₀ = 0.0 mM, [catalyst] = 2.0 g/L, T = 30 °C and pH_initial = 3.0). Adapted from Rodrigues et al. [66].](image)

Another study evaluated the degradation of OII dye by photo-assisted CWPO, and the reduction of color, mineralization, and oxidant consumption increased when the oxidant dose increased from 1.5 to 3.0 mM [110]; for concentrations of 6.0 and 12.0 mM, a negative effect was observed for dye and TOC removals. For the optimal oxidant concentrations, efficiencies of 85.9, 96.8, and 94.5% were reached for dye and TOC removals and H₂O₂ consumption, respectively. Moreover, authors reported that the catalyst used (gold on alumina) did not show any leaching for any of the concentrations of hydrogen peroxide evaluated.

Yang et al. [108] tested gold supported on carbon as a CWPO catalyst for the oxidation of BPA. A smaller oxidant dose (275 mg/L) allowed to remove ~50% of the model compound and ca. 35% of oxidant consumption was found after 12 h of reaction. An improvement of catalytic activity was observed when the hydrogen peroxide concentration was increased to 530 mg/L, allowing, after 12 h, to reach a bisphenol A reduction of ca. 70% and a consumption of oxidant of about 40%. However, a further increase in the oxidant dosage (835 mg/L) did not influence the efficiency of CWPO. A similar
tendency was reported by Lorençon et al. [88] for the CWPO catalyzed with Au/CNT for the degradation of a lipophilic compound (DPPH). The removal of DPPH increased with H2O2 concentration until 250 mM and remained practically constant for higher oxidant doses, as shown in Figure 9. The authors attained total degradation of DPPH for an optimal oxidant concentration of 250 mM. These two works highlighted once again the existence of an optimum oxidant dose for the gold catalysed CWPO process.

![Figure 9. Influence of initial hydrogen peroxide concentration in the degradation of a lipophilic compound (DPPH) by CWPO using Au/CNT as catalyst ([DPPH]0 = 0.2 mM, [catalyst] = 1.0 mg/mL, T = room temperature, stirring = 600 rpm). Adapted from Lorençon et al. [88].](image)

Contrarily to what was reported in the studies mentioned above, in which an optimum oxidant dose was found, in the work of Martin et al. [73] a reduction in the concentration (or even up to total absence) of phenol and intermediate compounds (catechol, quinone, and hydroquinone) was found, resulting from the oxidation of the pollutant when the H2O2 concentration was increased in the tested range (362–1447 mg/L). On the other hand, the biodegradability of the effluent improved, increasing the BOD5/COD ratio from ~ 0.05 to 0.4, when the oxidant dose increased from 362 to 1447 mg/L. However, higher doses were not tested.

The optimization of hydrogen peroxide amount is not only important for economic reasons, as the reagent is relatively expensive, but also to guarantee that H2O2 solution is not in excess. On one hand, too much H2O2 is detrimental to the subsequent biological treatment, if required, having deleterious effects on the microorganisms and leading to a decrease in the efficiency of the biological process. On the other hand, an excess of H2O2 contributes to the COD of the treated effluent, a commonly legislated parameter, and can give an erroneous indication of the possibility of the effluent discharge into water bodies.

5.2.3. Initial pH

The efficiency of wet peroxidation is also strongly dependent on the pH of the medium. A pH < 2.5 allows the scavenging reaction between the hydroxyl radical and H+ to take place (Equation (18)) [113]. Furthermore, at neutral or alkaline conditions, hydrogen peroxide self-decomposition into water and oxygen (Equation (19)) is promoted, decreasing the amount of available oxidant to yield hydroxyl radicals to promote organics degradation.

\[
\text{H}^+ + \text{HO}^* + e^- \rightarrow \text{H}_2\text{O} \tag{18}
\]

\[
2\text{H}_2\text{O}_2 \rightarrow 2\text{H}_2\text{O} + \text{O}_2 \tag{19}
\]
In the homogeneous process, for very acidic pH values, Au\(^+\) is present in lesser amounts, while higher pHs lead to precipitation of gold in insoluble form (AuHO), resulting in reduction of the amount of Au available, which leads to small formation of radicals (Equation (5)). These drawbacks are overcome when gold is supported on a porous support (heterogeneous process), once the metal is inside the pores and confined within the structure of the solid matrix \([52,114]\). This reduces the precipitation of gold that occurs in the homogeneous process; thus, the catalyst is available to decompose the hydrogen peroxide and generate the hydroxyl radical.

However, Domínguez et al. \([72]\) observed that the CWPO of phenol using gold supported on activated carbon was efficient in acid and neutral pH range (3.5–7.5) and decreased significantly for pH = 10.5. The authors achieved, after 24 h of reaction, removals of phenol and TOC of ~100 and ~60\%, respectively, with an efficiency of hydrogen peroxide use (\(\eta\), evaluated by the ratio between the amount of TOC removal and oxidant consumption) of 0.8 for pH values between 3.5 to 7.5.

A similar behavior was found by Martin et al. \([73]\). These authors showed that gold supported on diamond nanoparticles was catalytically active in the pH range between 4 and 7, with almost all phenol being degraded. They also observed a strong decay in its removal for the pH of 8 and 9 and total consumption of hydrogen peroxide for all values of pH tested (Figure 10). These authors reported that the biodegradability (evaluated by the BOD\(_5\)/COD ratio) was higher (~0.7) for the lower pH value tested (4.0), which was associated with no phenol detection and the lowest concentration of catechol and hydroquinone (intermediate compounds resulting from the oxidation of phenol) after CWPO treatment at this pH value, and decreased to values near 0.4 for higher pHs (5.0–7.0) that present phenol and higher concentrations of catechol and hydroquinone in solution after treatment.

**Figure 10.** Effect of pH in phenol (a) and hydrogen peroxide (b) concentration during CWPO reaction with gold supported on diamond as catalyst at different pH values. Lines were added merely to better illustrate the data trends ([phenol]\(_{\text{initial}}\) = 1.0 g/L, [H\(_2\)O\(_2\)]\(_{\text{initial}}\) = 1.44 g/L, [catalyst] = 320 mg/L and T = 50 °C). Adapted from Martin et al. \([73]\).

Studies of OII degradation by CWPO, without \([66]\) and with radiation \([110]\), using gold supported on alumina showed that the initial pH has an effect on the efficiency of the process. The authors observed optimum activity for initial pH = 3, which maximized the removal of OII (>97\%) and TOC (85.9\%) (see Figure 11), as well as the generation of hydroxyl radicals. The formation of radicals was evaluated in runs carried out without OII dye and when hydroxyl radicals were formed, and thus became in contact with 1,5-diphenyl carbazide, 1,5-diphenyl carbazone was formed, which presented a brown color that was measured at 563 nm. However, the oxidant consumption increased with pH (see Figure 11). As the support used in this study was alkaline, it increased the pH of the medium and, for the optimum initial pH found, the pH after 2 h of reaction was 4, which is in agreement with the
The best pH value reported by Domínguez et al. [72] and Martin et al. [73]. In these studies, all oxidant was consumed and the gold did not leach into solution during the oxidation for the initial pH range tested.

Concerning the industrial application perspective, the gold catalysts that allow the use of oxidative processes at neutral pH are more advantageous, because they reduce the costs associated with the acid consumption needed to decrease the pH to the acid range, and the base necessary to neutralize the effluent after the treatment, before it is discharged into the water bodies or subsequent treatment processes, as biological degradation.

5.2.4. Temperature

The temperature has a large influence on the efficiency of the CWPO process. The possibility to increase the operating temperature, as a way of improving the efficiency of the process, has been scarcely investigated, because the idea of thermal decomposition of H$_2$O$_2$ into O$_2$ and H$_2$O seems
to be widely accepted as a serious drawback [11]. However, according to the Arrhenius law, higher temperatures (often up to ca. 50–70 °C) can lead to a more efficient use of \( \text{H}_2\text{O}_2 \) upon enhanced generation of \( \text{HO}^\bullet \) radicals, at low metal concentrations. A decrease of the metal dose is important, since it improves the efficiency of \( \text{H}_2\text{O}_2 \) use, by minimizing competitive scavenging reactions [116]. Moreover, increasing the temperature accelerates oxidation of the organic compounds by the radicals.

Therefore, an increase in the temperature can be considered as a way to intensify the treatment process. Domínguez et al. [72] observed a positive effect of temperature, in the range of 50–80 °C, in the removal of phenol by CWPO using gold supported on activated carbon. The same tendency was reported by Drašinac et al. [107], as the methyl orange dye removal increased from ~30% at 25 °C to 85% at 80 °C, after 250 min of reaction, when the process was catalyzed by gold supported on modified titanium nanotubes.

Martin et al. [73] found that the reaction rate of phenol degradation and hydrogen peroxide consumption, when using an Au/DNP catalyst, increased with temperature in the range of 40–100 °C, reducing the reaction time from ~30 h to ~2.5 h, respectively, always reaching 100% compound removal and total consumption of oxidant. The same tendency was observed in a study that evaluated the methyl orange dye degradation, as its removal and TOC reduction increased with temperature in the range of 25 to 80 °C (see Figure 12), reaching 85 and 83% for MO and TOC reduction, respectively, for the optimal temperature (80 °C) [107].

![Figure 12](image-url)  
**Figure 12.** Methyl Orange (MO) decolorization during CWPO using gold supported on titanium nanotubes as catalyst, at different temperatures. Lines were added merely to better illustrate the data trends ([MO] \_initial = 50 mg/L, \[\text{H}_2\text{O}_2\] \_initial = 0.15 M, [catalyst] = 2.0 g/L and pH \_initial = 3.0). Adapted from Drašinac et al. [107].

In contrast, the effect of decomposition of hydrogen peroxide in water and oxygen was observed in the works of Yang et al. [108] and Rodrigues et al. [66,110]. Yang et al. [108] observed a significant increase in bisphenol A degradation and consumption of the hydrogen peroxide during CWPO, using Au/SRAC as catalyst, when the temperature increased from 30 to 40 °C (see Figure 13). Also when the temperature was raised to 50 °C, a little increase in the process efficiency was obtained. However, increasing the temperature to 60 °C, showed no improvement in the degradation of bisphenol A.
Table 4 shows an outline of these studies, reporting the operating conditions used and the efficiency achieved. The use of radiation increases the rate of oxidation since there are additional mechanisms for the formation of free radicals, according to the following three processes: i) the catalytic decomposition of hydrogen peroxide in the presence of radiation (Equation (20)) [42,117], ii) the decomposition of hydrogen peroxide by incidence of radiation (Equation (21)), and iii) photolysis of gold hydroxide (Equation (22)).

\[
\text{X-Au}^{n+} + \text{H}_2\text{O}_2 + \nu \rightarrow \text{X-Au}^{(n+1)+} + \text{HO}^* + \text{HO}^- \tag{20}
\]

\[
\text{H}_2\text{O}_2 + \nu \rightarrow 2\text{HO}^* \tag{21}
\]

\[
\text{X-Au(OH)}^{n+} + \nu \rightarrow \text{X-Au}^{n+} + \text{HO}^* \tag{22}
\]

On the other hand, an improvement of the performance in radiation-assisted CWPO can also occur due to direct photolysis of the organic compounds to degrade.

Some authors report that an increased radiation intensity has a positive effect on the performance of the treatment process. Navalon et al. [74] showed that an increase of laser power from 0 to 70 mJ/pulse improved the phenol degradation when Au/OH-npD catalyst was used. Rodrigues et al. [110] observed an increase of color and TOC removal, as well as of hydrogen consumption, with the radiation intensity of a TQ150 mercury lamp when Au/Al$_2$O$_3$ was used as photo-catalyst.
Table 4. Gold catalysts used in CWPO assisted with radiation for model compounds degradation or wastewater treatment, operational conditions, and performances reached.

| Model Compound/Effluent | Catalyst | Operation Conditions | Efficiency of CWPO assisted with Radiation | Ref. |
|-------------------------|----------|----------------------|------------------------------------------|-----|
| Phenol                  | Au/FH₂ (0.1%) | pH = 4.0; Room temperature; [phenol] = 100 mg/L; t = 3.5 h; Radiation: Sunlight | Phenol removal = 100%; H₂O₂ consumption = ~60% |    |
|                         | Au/FN₂ (0.5%) | pH = 4.0; Room temperature; [H₂O₂] = 200 mg/L; [phenol] = 100 mg/L; t = 8 h; Radiation: Sunlight | Phenol removal = -20%; H₂O₂ consumption = -20% | [75]|
|                         | Au/F (0.5%)   | pH = 4.0; Room temperature; [H₂O₂] = 200 mg/L; [phenol] = 100 mg/L; t = 8 h; Radiation: Sunlight | Phenol removal = -10%; H₂O₂ consumption = -15% |    |
| Acid Orange 7 dye (AO7) | Au/CeO₂ (1.0 at.%) | pH = 3.0; T = 30 °C; [H₂O₂] = 20 mM; [catalyst] = 0.5 g/L; [AO7] = 35 mg/L; t = 6 h; Radiation: Visible light | Dye removal = 100% | | [106]|
| Orange II (OII) dye     | Au/Al₂O₃ (0.7 wt.%) | pH = 3.0; T = 30 °C; [H₂O₂] = 6 mM; [catalyst] = 2.0 g/L; [OII] = 0.1 mM; t = 2 h; Radiation: UV/visible light (500 W/m²) | Dye removal = 97.8%; TOC removal = 68.2% |    |
|                         | Au/Fe₂O₃ (0.8 wt.%) | pH = 3.0; T = 30 °C; [H₂O₂] = 6 mM; [catalyst] = 2.0 g/L; [OII] = 0.1 mM; t = 2 h; Radiation: UV/visible light (500 W/m²) | Dye removal = 96.9%; TOC removal = 58.4% |    |
|                         | Au/Fe₂O₃ (4.0 wt.%) from WGC | pH = 3.0; T = 30 °C; [H₂O₂] = 6 mM; [catalyst] = 2.0 g/L; [OII] = 0.1 mM; t = 2 h; Radiation: UV/visible light (500 W/m²) | Dye removal = 98.5%; TOC removal = 73.5% |    |
|                         | Au/TiO₂ (1.6 wt.%) | pH = 3.0; T = 30 °C; [H₂O₂] = 6 mM; [catalyst] = 2.0 g/L; [OII] = 0.1 mM; t = 2 h; Radiation: UV/visible light (500 W/m²) | Dye removal = 99.8%; TOC removal = 73.4% |    |
|                         | Au/ZnO (1.2 wt.%) | pH = 3.0; T = 50 °C; [H₂O₂] = 3 mM; [catalyst] = 2.0 g/L; [Al₂O₃] = 0.1 mM; t = 2 h; Radiation: UV/visible light (500 W/m²) | Dye removal = 99.9%; TOC removal = 90.9%; H₂O₂ consumption = 98.6%; Gold leaching < 0.5 mg/L | [110]|
| Acrylic dyeing wastewater | Au/Al₂O₃ (0.7 wt.%) | pH = 3.0; T = 50 °C; [H₂O₂] = 6 mM; [catalyst] = 2.0 g/L; [Al₂O₃] = 0.1 mM; t = 2 h; Radiation: UV/visible light (500 W/m²) | Color removal = 100%; TOC removal = 72.4%; COD removal = 70.0%; BOD₅/COD = 0.5; Specific Oxygen Uptake Rate = 17.9 mgO₂/(gVSs h); Inhibition of Vibrio Fischeri = 0.0% |    |
| Phenol                  | Au/HO-npD (1.0 wt%) | pH = 4.0; T = 30 °C; [H₂O₂] = 2.5 g/L; [catalyst] = 400 mg/L; [phenol] = 100 mg/L; t = 2 h; Radiation: Sunlight | Phenol removal = 100%; H₂O₂ consumption = 100%; COD removal = 69.7%; BOD₅/COD = 0.4 | [111]|
| Phenol                  | Au/HO-npD (1.0 wt%) | pH = 4.0; [H₂O₂] = 200 mg/L; [catalyst] = 160 mg/L; [phenol] = 100 mg/L; t = 2 h; Radiation: Laser Flash (70 mJ/pulse) | Phenol removal = 100%; H₂O₂ consumption = ~90% |    |
| Phenol                  | Au/CeO₂ (1.0 wt%) | pH = 4.0; [H₂O₂] = 200 mg/L; [catalyst] = 160 mg/L; [phenol] = 100 mg/L; t = 3 h; Radiation: Laser Flash (70 mJ/pulse) | Phenol removal = -15%; H₂O₂ consumption = -100% | [115]|
| Phenol                  | Au/TiO₂ (1.0 wt%) | pH = 4.0; [H₂O₂] = 200 mg/L; [catalyst] = 160 mg/L; [phenol] = 100 mg/L; t = 3 h; Radiation: Laser Flash (70 mJ/pulse) | Phenol removal = -10%; H₂O₂ consumption = -80% |    |

5.3. Catalyst Stability

The deactivation of catalysts in CWPO is mostly associated with the loss of metal by leaching from the solid support to the effluent during the treatment process, but also with possible gold nanoparticle sintering and/or pore blockage. For application of catalytic wet peroxidation in the treatment of real effluents, a crucial aspect to be taken into account is the reutilization of the catalysts, without reduction of their efficiency and stability in consecutive cycles of use, making it imperative to evaluate how stable and durable they are.

Ferentz et al. [71] evaluated the long-term stability of Au/TiO₂, with 2.8 and 3.2 Au wt.%, in a fixed bed reactor. The first catalyst was stable during 50 h, achieving constant removals of TOC and phenol >90 and >99%, respectively. In the period of 50 to 75 h, the efficiency of CWPO dropped to ~65% of
TOC removal and remained constant for higher reaction times, corresponding to the performance reached with pristine TiO$_2$. The authors associated the deactivation of the catalyst to the gold particles growth (as the size increased from 3–4 nm in the fresh catalyst to 20–30 nm after CWPO). For the second catalyst (3.2 wt.% Au/TiO$_2$), the efficiency of the process, in terms of TOC removal, decreased from ~95% to ~80% during the first 50 h of reaction and remained practically constant for higher reaction times. This loss of catalytic activity was attributed to: i) an increase of the gold size from 7–8 nm to 13–15 nm for fresh and used catalysts after 300 h, respectively, and ii) adsorption of dicarboxylic acids in gold, with the equilibrium established at 50 h.

In the work developed by Domínguez et al. [72], the activated carbon supported gold deactivated in the first cycle of CWPO of phenol. The removal of the model compound observed by the authors in next three cycles was attributed to the activity of activated carbon alone (see Figure 14). Since the authors did not observe any gold leaching for the solution, the catalyst deactivation was attributed to the presence of dicarboxylic acids (by-products formed in the oxidation of phenol), which adsorb on gold nanoparticles. In order to recover the catalytic activity, the catalyst was regenerated after the first cycle by: i) alkaline washing to pH 14 with Na$_2$CO$_3$, in order to dissolve the absorbed species, and subsequent washing with distilled water until neutralization, and ii) oxidative thermal treatment at 200 °C, during 14 h, in air atmosphere, which allowed to burn-off the carboxylic acids adsorbed on the gold nanoparticles. The first regeneration process led to an activity recovery of about 60% and the catalyst loss was 2 wt.% of gold by leaching. For the second process, a more significant loss of gold (10 wt.%) was observed, however, the catalyst activity was fully restored (see Figure 14).

In contrast to Ferentz et al. [71] and Domínguez et al. [72] that reported a loss of the catalytic activity of gold catalysts during CWPO, several studies in the literature refer to the stability of gold on different solid supports [66,67,70,73,108,110] when used in subsequent reutilization cycles. In the investigations performed by Rodrigues et al. [66,67,110], the gold supported on alumina, zinc oxide, titanium oxide, and iron oxide was stable during 3–5 consecutive cycles in acid medium (pH = 3.0), with OII and TOC removals and H$_2$O$_2$ consumption remaining unchanged during the cycles. The authors observed no catalyst loss of gold by leaching during the reactions.

Gold supported on activated carbon also did not deactivate in acidic pH (3.0) for degradation of bisphenol A by CWPO, with the removal of BPA and the consumption of oxidant being more or less constant, in ~80 and ~40%, respectively, during four consecutive cycles. This demonstrates that the catalyst can be reused several times [108]. The same tendency was observed by Han et al. [70], who evaluated the stability of gold supported on hydroxyapatite, showing that the conversion of phenol was constant after five cycles, either at pH 2.0 (>90%) or 5.0 (~80%) (see Figure 15). Similar results were
obtained by Sempere et al. [75] that reused diamond supported gold (submitted to a thermal treatment at 420 °C and subsequent annealing treatment with hydrogen) three times in sunlight assisted CWPO in the oxidation of phenol. This catalyst did not lose the catalytic activity and the leaching of gold was negligible (< 1% of the initial gold in the first cycle of utilization) or was not observed (in the second and third cycles).

Figure 15. Conversion of phenol by CWPO using gold supported on hydroxyapatite as catalyst in consecutive five cycles of reutilization at pH = 2.0 and 5.0 ([phenol] = 100 mg/L, T = 70 °C, [catalyst] = 0.1 g/L and V_{H2O2} with 30 wt.% = 1 mL). Adapted from Han et al. [70].

Martin et al. [73] evaluated the reuse of a diamond supported gold (Au/DNP) catalyst in phenol oxidation by CWPO during four cycles. The authors exhaustively washed the material with water at pH = 10 and, finally, with distilled water, in order to eliminate the deactivation of the catalyst by adsorption of carboxylic acids (intermediate products generated by phenol oxidation) on gold, as also pointed out by Domínguez et al. [72] as the main reason for deactivation. The Au/DNP, after a simple treatment by washing, can be reused during four times, reaching, in all cycles, not only total conversion of phenol and consumption of H2O2 in the end of reaction, but the same temporal profiles [73] (see Figure 16). However, the washing of the catalyst led to a loss of gold to the solution, but the leaching decreased with an increase of the cycles (from 3 to <0.1 wt.% after 1st use and 4th use, respectively), so that the performance of the catalyst was not affected by the small leaching and no deactivation was found [73].

Figure 16. Evolution of phenol (a) and hydrogen peroxide (b) concentration during consecutive reuse cycles ([phenol] = 1 g/L, T = 50 °C, pH = 4.0, [Au/DNP] = 320 mg/L and [H2O2] = 1.44 g/L). Adapted from Martín et al. [73].
Navalon et al. [74] reused Au/HO-npD catalysts during three consecutive cycles, washing with water at pH 10 after each cycle. The authors did not observe any significant change in phenol and H$_2$O$_2$ disappearance profiles. Additionally, a run was carried out with a large excess of pollutant (40 g/L) and 0.5 mg/L of catalyst, in order to have an equivalent of 400 consecutive reuse cycles and use 5.5 equivalent of oxidant with respect to phenol. This run allowed to conclude that Au/HO-nDP had the ability to remove 36% of the pollutant before being deactivated, however, an alkaline washing before the second and third cycles allowed the catalyst to recover the catalytic activity.

6. Conclusions

This review showed that gold catalysts can be efficiently used in CWPO processes and that their catalytic activity depends on several operational variables of the process (such as pH, temperature, oxidant and catalyst concentrations, and gold content), as well as on the properties of the catalysts. We believe that such gathered information will provide useful insights that might lead to a more rapid and effective optimization of catalytic wet peroxidation processes using these materials.

Despite the recent progresses in this area, it is still crucial that work continues to be done to better apply these catalysts in the treatment of real effluents by CWPO and to deepen the knowledge coming from the laboratory studies for the scale-up of the process. The main limiting step in the application of this technology might be the costs of the treatment, since gold has a high cost (compared to other more common materials); however, gold also shows advantages, namely, high stability, high efficiency, and absence of leaching into the solution, which might justify the investment.

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Abbreviations

- AO7: Acid Orange dye
- AOPs: Advanced Oxidation Processes
- Au/AC: Gold on activated carbon
- Au/C: Gold on carbon
- Au/CNF: Gold on carbon nanofibers
- Au/CNT: Gold on carbon nanotubes
- Au/X40s: Gold on coconut shell carbon
- Au/DNP: Gold on diamond nanoparticles
- Au/F: Gold on diamond after thermal treatment at 420 °C in air atmosphere
- Au/FH$_2$: Gold on diamond after thermal treatment at 420 °C in air atmosphere and at 500 °C in hydrogen atmosphere
- Au/FN$_2$: Gold on diamond after thermal treatment at 420 °C in air atmosphere and at 500 °C in nitrogen atmosphere
- Au/Hap: Gold on hydroxyapatite
- Au/npD: Gold on nano power diamond
- Au/HO-npD: Gold on nano power diamond previously treated with Fenton reagent
- Au/FDU-15: Gold on ordered mesoporous carbon
- Au/PSAC: Gold on pitch-based spherical activated carbon
- Au/SRAC: Gold on styrene-based activated carbon
- Au/TN: Gold on titanium nanotubes functionalization with hydrogen peroxide
- Au/TiO2-AD: Gold on titanium oxide prepared by adsorption method
- BOD$_5$: Biological oxygen demand after 5 days
- BPA: Bisphenol A
- CWPO: Catalytic Wet Peroxidation
COD  Chemical oxygen demand  
DM  Gold metal dispersion  
DPPH  1,1-diphenyl-2-picrylhydrazyl  
EU-WFD  European Union Water Framework Directive  
HR-TEM  High-resolution transmission electron microscopy  
Iν  Ultraviolet radiation  
M  Transmission of metallic cations  
MB  Methyl Blue dye  
MO  Methyl Orange dye  
NHE  Normal hydrogen electrode  
OII  Orange II dye  
RH  Organic matter  
TOC  Total organic carbon  
TOF  Turn off frequency  
WGC  World Gold Council  
X  Support

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