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A revised photocatalytic transformation mechanism for chlorinated VOCs: experimental evidence from C$_2$Cl$_4$ in the gas phase

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Abstract

The photodegradation of gaseous perchloroethylene (PCE) was investigated on titanium dioxide under UV light at 365 nm at the gas/solid interface in a CSTR photo-reactor (Continuous Stirred Tank Reactor). The rate and products are strongly affected by oxygen presence. Gaseous products of PCE degradation agree with literature. The production of active chlorine (sum of Cl$_2$, HClO, ClO and ClO$_2$) was investigated both in the presence of oxygen and in anoxic conditions. At low O$_2$ concentration no gaseous active chlorine was determined, while a significant amount was measured in the presence of oxygen. By considering that in the absence of O$_2$ the only possible form of active chlorine is Cl$_2$, this highlights that Cl$^-$ is not produced, and that chain reactions promoted by the chlorine radical do not occur on the TiO$_2$ surface.

The photocatalytic transformation of C$_2$Cl$_4$ was investigated at different concentrations. The rate follows a first order kinetic that is rationalized with a photocatalytic kinetic model in which the substrate is able to react simultaneously with both photogenerated holes and electrons. In anoxic conditions adsorbed halogenated organic compounds with molecular weights higher than that of PCE were produced and chloride ions accumulated at the surface. Reductive pathways have a key role in PCE degradation. The water vapor has a detrimental role on the PCE transformation rate due to the competition with PCE adsorption on reactive sites with 2:1 stoichiometry.

The addition of chloride ions on TiO$_2$ surface slows down the PCE degradation rate and the production of gaseous CCl$_4$ but increments that of C$_2$Cl$_6$ in anoxic conditions. This is rationalized by a mechanism in which direct hole transfer to substrate occurs followed by chloride anion addition to the carbocation.

Keywords

Photocatalysis, Titanium Dioxide, Perchloroethylene, Chlorine radical, Gas/solid interface, Active chlorine
1 Introduction

The photocatalytic process is a well-known method for abatement of air pollutants. Early studies showed that alkanes, aromatic hydrocarbons, alcohols, aldehydes, ketones, and halogenated solvents can be degraded although in a different extent. The studies on halogenated solvents, namely trichloroethylene (TCE), tetrachloroethylene (PCE) and chloroform date at late 80’s.[1] The main products observed were DCAC (dichloroacetyl chloride), COCl₂ (phosgene), CO₂, CO and HCl. Nimlos et al. [2] were also able to identify Cl₂ using a molecular beam mass spectrometer. In the degradation of TCE and PCE, CHCl₃ and CCl₄ [3,4] were formed as the O₂ mole fraction in the feed gas decreases. The formation of more halogenated products was ascribed to the initial formation of Cl• radicals, and the subsequent preferential attack at the -CHCl side.[3] Comparison of literature data [5] under similar conditions and reactor configuration using the Langmuir Hinshelwood model showed that toluene, m-xylene, acetone and TCE have similar order of magnitude rate constant and apparent binding constant.[6] A photocatalytic rate enhancement for the conversion of many pollutants was reported when co-fed with chloro-olefins.[7] Quantum yield greater than 1 has been cited several times [1,8], although the original papers are almost not available. A detailed mechanism was also proposed through ab initio molecular orbital calculations, showing that addition of 'OH and Cl• radicals to TCE is possible, being the first more exothermic than the second one.[3,8] Nowadays, PCE and TCE are still used as model pollutants for reactor configuration studies.[9]

The positive synergic effect, the formation of more halogenated intermediates, and the cited quantum yield greater than 1 for chlorinated compounds were ascribed to chain reactions promoted by the chlorine radical that is formed at the TiO₂ surface.[3,10] The chlorine radical can be formed a) from chloride oxidation by 'OH or holes, or b) directly released from oxidized intermediates. The supposed Cl• formation by hydroxyl radicals produced at the surface of TiO₂ exposed to near-UV irradiation in the presence of adsorbed water vapor [11] was able to justify the observed intermediates.[8] The role of chlorine radical was also supported by experiments in which chloride ions were added in the gas phase or at the surface. After addition of HCl in catalytic quantity in the air flow or by PCE overheating, a sensible increase of the reaction rate was obtained.[12] The addition of surface Cl• through prechlorination with HCl increased the rate for toluene and hexane at the very beginning of the reaction when compared to the co-feeding with TCE, for which the enhancement was observed after some reaction time needed to accumulate a useful surface Cl• level.[13]

The chain reaction mechanism assumes that chorine radical is an initiator which adds to TCE, forming a more halogenated radical, which in some further steps releases Cl• or HCl after reaction with oxygen.[3] The chain is terminated when Cl• reacts with water to form 'OH (a reaction possible only for pH > 2-2.5 as suggested by the standard redox potential for the couples E°(Cl•/Cl−)= 2.60 V and E°('OH,H⁺/H₂O)=2.73 V [14]), or in dry conditions to form Cl₂.[15] Chlorine radical could also be depleted by photoelectrons in the conduction bands (reducing to chloride ion that is adsorbed on the surface) or by reaction with O₂, forming ClO₂.[3]

The aforementioned framework based on the key role of Cl• seems to be consistent. However, parallel to earlier studies, alternative explanations could be suggested. For example, literature-reported values of photon efficiency are consistent with a photocatalytic mechanism
without chain reactions mediated by Cl• radical. Using a fluorescence UV source and potassium ferrioxalate actinometry, the TCE photon efficiency was calculated as 13% over conditioned catalysts.[16] Jacoby et al.[1,2] measured quantum yields from 0.5 to 0.8 under high TCE concentration. In addition, the inhibition of the rate in the presence of halide ions in aqueous solution was suggested to be due to the competition with organic molecules for the oxidative species (i.e. photogenerated holes).[17] The chlorine radical, if formed, is subjected to rapid recombination with conduction band electrons, creating a short-circuit cycle that depresses the conversion. An analogous inhibition was observed in the gas-phase photocatalytic degradation of CHCl₃ due to the increase of surface Cl• with the space time.[18] The photoactivity was recovered washing the catalyst. Recently, EPR measurements were conducted to detect the presence of chlorine radicals by using the PBN spin-trap in chloroform solution.[19] The PBN–Cl• radical was not identified, suggesting that the existence of chlorine radical can be largely disregarded, although not completely excluded. Then the rate decrease was ascribed to the surface sites occupancy without a redox involvement of chloride ion. If so, also the formation of •OH radicals, more endoergonic, could hardly be suggested.

The existence of •OH radicals in photocatalysis was debated for years.[20] Although free •OH is still often invoked in the recent literature, the surface trapped hole is most probably the reactive species, often indistinguishable as reactivity from •OH radical. In the photocatalyzed oxidation of TCE in the presence of ¹⁸O₂, oxygen-18 is incorporated into DCAC, phosgene, and CO, indicating that only the oxygen from the gas phase is involved in the oxidation of TCE, whilst oxygen-16 from H₂¹⁶O is not incorporated in the photooxidation products.[21] This proved that •OH driven oxidation mechanism is not operative in the photooxidation of TCE. The direct electron transfer from organic substrate to the photogenerated holes was recently demonstrated as a unique electron transfer mechanism in the case of photocatalytic transformation of melamine, which is unreactive toward •OH radical.[22] This indicates that direct electron transfer from the surface trapped hole could be in general the sole process responsible for oxidation.[23] Consistently, the product analysis obtained with O₂-sensitised photo-oxidation of indane and some of its hetero-analogs in deaerated CH₃CN and in the presence of Ag₂SO₄ was explained with the intervention of a carbocation in the first reaction event resulting from an electron-transfer mechanism (from the substrate to the photogenerated hole).[24]

To clarify the possible role of •Cl radical some new data on intermediate formation in the photocatalytic degradation of PCE are here reported, both in the presence and absence of O₂, together with the analysis of kinetic data. The formation of gaseous Cl₂ is assessed by a simple trapping experiment.

2 Materials and Methods

Details on used materials, preparation of the catalyst, and experimental setup are reported in Supplementary Material (hereafter SM). The experiments of PCE transformation were carried out in a homemade flow-through photo-reactor working in a regime of perfect mixing (Continuous Stirred Tank Reactor, CSTR, see SM under 1.4).[25,26] Under steady state the output concentration of gaseous C₂Cl₄ does not change with the time, if there is no change in the photoactivity of the catalyst, e.g. no poisoning or photo-activation as a consequence of the irradiation. Consequently, defining the conversion at the time t as \( \eta(t) = \frac{(C_o - C_{out}(t))}{C_o} \), where
$C_o$ is the feed concentration and $C_{out}$ the concentration inside the reactor and measured at its outlet, the rate under stationary conditions is directly obtained from the experiment (eq. 1).[25]

\[
Rate(C_{out}) = \frac{C_o \eta F}{S} \quad \text{(eq. 1)}
\]

The monitoring of C₂Cl₄ flowing out of the reactor was carried out with a Photo Ionization Detector. The volatile VOCs were analysed through both GC-MS after cryofocusing and directly through an FTIR spectrophotometer equipped with a cell for gases with 8 m long optical path length. The catalyst surface modification was assessed through UV-Vis reflectance spectra and FT-IR. The surface-deposited compounds have been extracted and analysed with GC/MS and IC. All details are reported in SM.

The determination of the active chlorine formed during the photocatalytic test was carried out according to a standard spectrophotometric methods,[27] based on the reaction of active chlorine (Cl₂/HClO/ClO⁻/ClO₃⁻) with N,N-diethyl-p-phenylenediamine (DPD). This reaction produces a pink coloured species that can be quantified spectrophotometrically at 510-515 nm. The gas flowing out of the reactor was bubbled through a fritted glass in a slightly basic solution (NaOH 2 mM). This solution favoured the trapping of volatile active chlorine species through: 1) deprotonation of the volatile hypochlorous acid into the non-volatile hypochlorite anion; 2) dismutation of Cl₂ into Cl⁻ and reactive ClO⁻ (Cl₂ + 2 OH⁻ → H₂O + Cl⁻ + ClO⁻); 3) reaction of ClO with the excess of hydroxyl ions to form ClO⁻ (ClO⁻ + OH⁻ → ClO⁻ + CH⁺OH); 4) dismutation of ClO₂ in chlorite and chlorate species (2 ClO₂ + H₂O ⇌ HClO₂ + HClO₃, $K=1.2 \times 10^{-7}$). At different irradiation times a fixed volume of the solution was withdrawn and mixed with DPD solution buffered at pH 6.2 (phosphate buffer, 0.17 mol·L⁻¹ $\text{HPO}_4^{2-}$ e 0.33 mol·L⁻¹ $\text{H}_2\text{PO}_4$). The DPD produced pink colored species was quantified at 510 nm with a Cary 100 UV-Vis spectrophotometer.

3 Results and Discussion

3.1 Rate as a function of the inlet C₂Cl₄ and water vapor concentration

The kinetic of the photocatalytic transformation of C₂Cl₄ was evaluated i) at different concentrations of the inlet flow ($C_o$) and ii) at different concentration of water vapor concentration. In both cases under stationary state the conversion was obtained and from this the transformation rate was computed (eq. 1). It is worth mentioning that the rate calculated from the experimental conversion refers to the concentration inside the reactor, and to $C_{out}=C_o(1-\eta)$.

Fig. 1A shows the photonic efficiency (PE) at different $C_o$ and plotted vs the actual concentration inside the reactor ($C_{out}$). The photocatalytic efficiency PE is given by the photocatalytic rate [mol m⁻² s⁻¹] divided by the incoming light irradiance $I_o$ [mol of photons m⁻² s⁻¹]. The inset in Fig. 1A shows the stationary state (SS) conversion as a function of the inlet concentration of perchloroethylene. As the conversion is almost independent of the inlet concentration, a quasi-first order kinetic was observed ($rate=m\times C_{out}^n$ where $n=1$ for first order, observed $n=0.90$).

A basic kinetic approach to the photocatalytic process previously reported [28] and widely discussed,[23] based on 5 primary steps (light absorption with production of an exciton in the bulk, charge carriers migration to the surface, recombination of the charge carriers at the
surface, interface charge transfer of holes to a reduced substrate and conduction band
electrons transfer to an oxidant species), gives the rate=$\varphi \times QY$, where the quantum yield ($QY$)
of the photocatalytic process is:

$$QY = \left( -y + \sqrt{y(y+2)} \right)$$  \hspace{1cm} (eq. 2)

The master variable $y$ in eq. 2 is formally the ratio of two rates, the chemical rate at the
catalyst surface and $\varphi$, which is the rate of absorption of photons in units consistent with those
of the chemical rate (e.g. mol m$^{-2}$ s$^{-1}$):

$$y = \frac{k_0 \{C_{red}\} \{C_{ox}\}}{\varphi}$$  \hspace{1cm} (eq. 3)

where $k_0$ is a combination of microscopic kinetic constants, and $\{C_{red}\}$ and $\{C_{ox}\}$ are the
concentrations of the reduced and oxidized species adsorbed at the surface according to a
proper adsorption isotherm (e.g. in the most simplified case a Langmuir isotherm). The above
model, where the oxidant and reductant are different species, foresees a reaction order $0 \leq n \leq 0.5$,
often experimentally verified [22,28] and recently deeply tested for the oxidation reaction
of formic acid in the presence of oxygen.[29]

The first order observed in the case of PCE (see Fig. 1A) can be rationalized in the
framework of the basic kinetic model briefly summarized above considering the possibility
that C$_2$Cl$_4$ reacts both with $h_s^+$ and $e_s^-$. This double reactivity of the substrate has been hardly
ever considered in the gas-phase halocarbon degradation because i) the most of the
experiments are carried out in the presence of O$_2$, ii) oxygen is believed to be an efficient
electron scavenger and iii) the main goal of the photocatalytic processes is the full oxidation
(abatement) of the substrate. An effective competition between O$_2$ and PCE for the
photoproduced electrons can be operating also in the presence of excess oxygen as few PCE
ppm were able to increase toluene degradation in the presence of air (200000 ppm).[7]

The importance of reductive pathways was proved in aqueous slurry in the study of photo-
induced hydrolysis of chlorinated methanes under anaerobic conditions [30], in which the
interconversion of CCl$_4$, CHCl$_3$ and CH$_2$Cl$_2$ was supposed to proceed by reductive release of
chloride ions. An initial reductive dominance of the reductive pathway was observed for CCl$_4$
even in the presence of oxygen. Reductants such as alcohols (methanol, propan-2-ol, tert-
butanol) remarkably enhance CCl$_4$ degradation as they scavenge holes.[31] Accordingly, it is
worth noting that the early literature reported that compounds exhibiting TCE rate promotion
were toluene, ethylbenzene, m-xylene,[7] methyl ethyl ketone (MEK), acetaldehyde,
butyraldehyde, methyl tert-butyl ether (MTBE), methyl acrylate, 1,4-dioxane,[32] and
methanol.[33] All these compounds are degraded mainly by an oxidative pathway. Then e$_{CB}$
scavenging by TCE would increase the rate. Rate inhibition by TCE was exhibited for
acetone, methylene chloride, chloroform, and 1,1,1-trichloroethane.[32] These compounds are
more likely degraded by a first reduction step like TCE, which then acts as a competitor
reducing their transformation rate.

On the hypothesis that C$_2$Cl$_4$ reacts both with $h_s^+$ and $e_s^-$, it is possible to predict that the
presence of oxygen is not mandatory for an effective transformation of PCE (vide infra),
because both surface holes and electrons are able to be scavenged by the substrate itself (i.e
$\{C_{red}\} = \{C_{ox}\}$). In this case

$$QY = 2 \left( -y + \sqrt{y(y+2)} \right)$$  \hspace{1cm} (eq. 4)
where the master variable \( y = k_0[PCE]^2/\varphi \). Eq. 4 shows that when the substrate can be simultaneously oxidized and reduced the reaction order is \( 0 \leq n \leq 1 \). The first order at low \( \{PCE\} \) (low value of \( y \) and then low value of \( QY \) or \( PE \), see Fig. 1) is evident when \( y \) value is significantly lower than 2. For this case

\[
QY \approx 2[PCE]^{\frac{2k_0}{\varphi}}
\]  

(eq. 5)

As the absorbed light is proportional to the incident one \( (\varphi = \alpha I_0) \), and the photocatalytic rate is given by \( rate = \varphi QY \), the photonic efficiency \( PE = \frac{rate}{I_0} \) is

\[
PE = 2[PCE]^{\frac{1}{2\alpha k_0 I_0}} = k'[PCE]
\]  

(eq. 6)

where \( k' = \sqrt{8\alpha k_0 I_0} \). This equation justifies the first order rate observed for the PCE photocatalytic transformation (Fig. 1A), and supports the role of the reductive pathway, that will be deeply investigated under anoxic conditions (see below).

The transformation rate of PCE was also investigated at increasing relative humidity. Fig. 1B shows that the rate normalized for the incoming light intensity \( I_0 \) decreases with increasing water vapor content, as previously reported when the mole fraction of \( H_2O \) is 50–60 times higher than that of PCE.[34] The inhibition of the PCE photocatalytic rate with the increment of the water vapor concentration can be rationalized by considering the competitive adsorption of PCE with water \( W \) for the same adsorption sites \( s \), and can be modeled using a competitive Langmuir isotherm

\[
\{PCE\} = \frac{K_{ads}^{PCE}\{s\}[PCE]}{1 + K_{ads}^{PCE}\{PCE\} + K_{ads}^{W}[W]^n}
\]  

(eq. 7)

where \( \{ \} \) and \( [\] \) are concentrations at the surface and in the gas phase, respectively, and \( \{s\} \) is the concentration of adsorption sites. The above isotherm can be easily derived using the balance of surface sites and the equilibrium constants of reactions \( s + PCE \rightleftharpoons s-PCE \) and \( s + nW \rightleftharpoons s-W_n \), which are \( K_{ads}^{PCE} \) and \( K_{ads}^{W} \) respectively.

Combining equations (6) and (7) the experimentally determined \( PE \) is related to the concentration of gaseous water in the reactor by

\[
\frac{1}{PE} = a + b[W]^n
\]  

(eq. 8)

where \( a = \frac{1 + K_{ads}^{PCE}\{PCE\}}{k K_{ads}^{PCE}\{s\}[PCE]} \) and \( b = \frac{K_{ads}^{W}}{k K_{ads}^{PCE}\{s\}[PCE]} \). In the used experimental conditions \( a \) and \( b \) are constants ([PCE] = 3 ppm, constant surface area, constant light intensity).

By numerical fit of experimental data \( n = 2.2 \pm 0.1 \) was obtained, suggesting a displacement of 2 molecules of water for molecule of adsorbed PCE. The dashed curve in Fig. 1B is obtained with \( n=2.2, a=18.3, b=920 \) and shows the goodness of the model. Also linearization with eq. 8 has \( R^2 = 0.992 \). The model also predicts that \( PE \) is not dependent of the humidity at low water content, supplementing literature data [34] and confirming that traces of water are uninfluential for experiments carried out under dry condition. From the ratio \( a/b \) and under the reasonable hypothesis that \( K_{ads}^{PCE}\{PCE\} >> 1 \) one obtains \( K_{ads}^{PCE} = (1.6 \pm 0.2) \times 10^2 K_{ads}^{W} \). Because the dependence of the photocatalytic rate from the water content is highly dependent on the catalyst preparation [35], the reported value refers to the catalyst specimen used in this work.
As H$_2$O is needed to compensate surface unoccupied atomic orbitals,[36] the calculated value indicates that PCE is strongly adsorbed.

The competitive adsorption of H$_2$O and C$_2$Cl$_4$ on the same reactive sites is a strong proof that the photocatalytic transformation of PCE takes place at the TiO$_2$ surface, where the direct electron transfer occurs.

As a consequence of the detrimental role of water in the PCE photocatalytic transformation, the below reported experiments were carried out in dry gas. Although under this condition, traces of water are unavoidable, but such traces are uninfluential (see above).

### 3.2 Basic mechanism

The reactivity (transformation kinetics and by-products) was compared in the presence of synthetic air (N$_2$ 80% + O$_2$ 20%) and in anoxic conditions (N$_2$ 100%) with the aim to give insights on the role of oxidative and reductive pathways. A complete anoxia cannot be reached as a consequence of the ubiquitous atmospheric oxygen which unavoidably diffuses into the reactor and tubes, even though a continuous flow through the reactor with N$_2$ is fed. This hinders to reach a complete anoxia even at very long irradiation time. On the contrary in an ideal batch reactor in which diffusion of O$_2$ from the atmosphere is absent, initial traces of oxygen would be easily removed during the photocatalytic process and authentic anoxic condition would be reached. In actual anoxic experimental conditions a residual O$_2$ concentration ≤ 10 ppm was attained.

Fig. 2A shows the residual concentration of PCE going out of the reactor under irradiation. A significant photocatalytic transformation of PCE was observed both in the presence and in the absence of oxygen with a slightly higher conversion in oxygenated atmosphere. Despite of the most common photocatalytic transformations of organic substrates are completely blocked in the absence of electron scavengers (when {C$_{ox}$}=0, in eq. 3 $y$ = 0, and consequently $QY$ and the rate are null), the rate of PCE degradation is only partially depressed with oxygen shortage. This supports the previous hypothesis that C$_2$Cl$_4$ is able to be concurrently oxidized by holes and reduced by the photoelectrons, as depicted in the following reactions:

$$C_2Cl_4(ad) + h^+_u \rightarrow C^+Cl_2$$
reaction (r2) and that, in oxic conditions, it reacts in later steps of degradation. Then reductive electron transfer, which occurs only at the catalyst surface, is always operating.

From the comparison between the amount of cumulated chloride and the total amount of PCE photocatalytically transformed in the same time, the molar fraction $\chi$ of $C_2Cl_4$ converted in chloride was computed (ratio between the $\mu$mol of $C_2Cl_4$ degraded and the $\mu$mol of deposited chloride, both normalized per unit area). The values of $\chi$ are reported as inset in Fig. 2B. In the presence of $O_2$ larger values of $\chi$ were observed, which indicate that PCE is oxidized according to reaction (r1) without losing chloride anion, and later reactions with $O_2$ increment its degradation rate. Then organic chlorine must be released in some other forms as well as chloride.

The transformation of PCE was investigated on TiO$_2$ films on which different amount of chloride ion was deposited through evaporation of an aerosol of sodium chloride solution. To prevent that the amount of surface chloride changed significantly during the photocatalyzed reaction, the experiments were carried out at low conversion and shortly terminated (after 10 minutes of irradiation). Fig. 3A shows the PE with and without oxygen as a function of the chloride amount added at the surface. In both oxic and anoxic condition the addition of chloride ion had a detrimental effect on the PCE degradation rate for the reasons already discussed (a redox null cycle or physical occupation of sites, preventing PCE adsorption).

The former hypothesis was that organic chlorine could be released as Cl$^\bullet$, which also favors chain reactions useful for further degradation, and that also chloride ion can be oxidized to Cl$^\bullet$ radical as mentioned in the Introduction. Then, the production of active chlorine gives insights not only on the role of the surface chloride in the photocatalytic process, but also on some other peculiar aspects of the photocatalytic transformation of PCE.

The production of active chlorine (AC, i.e. $HClO/ClO^-$, $Cl_2$, $ClO$ and $ClO_2$) in the gaseous phase is reported in Fig. 3B as a function of the irradiation time under the four investigated conditions. Only in the presence of oxygen the production of a significant amount of AC was observed (after a delay time of roughly 30 minutes due to the exponential dilution in the CST reactor) while in anoxic conditions the concentration of AC is lower than the limit of detection. Because in the absence of $O_2$ the only active chlorine species that can be produced is $Cl_2$, the transformation of PCE does not involve pathways in which the production of $Cl_2$ is operational. The absence of $Cl_2$ excludes that Cl$^\bullet$ radicals are produced through oxidation by holes of chloride ion self-released or added as NaCl, or through release from intermediates, and agrees with the absence of Cl$^\bullet$ evidenced by EPR spectroscopy.[19] The holes at the surface ($h^\bullet$) have a sufficient redox potential for the oxidation of $Cl^-$ to $Cl^\bullet$.[17] A possible competition of chloride ion for $h^\bullet_{VB}$ is possible as a decrease of the rate was observed (Fig. 3A). However, this radical is evidently not formed because surface mediated recombination processes with $e^-_{CB}$ could be operational. Then, the suggested production of a significant amount of chlorine radical at the irradiated surface [2,4,39] is not in agreement with the absence of AC observed here without oxygen.

Conversely, in the presence of oxygen there is production of AC. Even if the eventually formed chlorine radical reacts with oxygen to give $ClO_2$, under irradiation it can be immediately transformed into $ClO$ through its photolysis via the reaction $ClO_2$ ($^3B_2$) $\rightarrow$ $ClO(2\Pi) + O(^3P)$ [38], and then the detected AC is composed of $HClO$ and/or $ClO$. However, because $Cl_2$ is not formed under anoxic conditions as above reported, there is good reason to assume that only oxygenated species such as $HClO/ClO^-$, and $ClO$ are produced in the
presence of oxygen. Furthermore, as no significant difference between the production of AC 337 in the presence of oxygen was observed for the pristine and surface chlorinated catalysts (see Fig. 3B), these species must derive from intermediates, and are produced through their dehalogenation, that is organic chlorine is directly released as HClO or ClO. Then, in the presence of oxygen the operational mechanism for PCE transformation would imply the production of oxygenated chlorinated species.

3.3 Main intermediates

The main by-products of the PCE photocatalytic transformation are those previously reported [2,4,39], namely phosgene, trichloroacetyl chloride (TCAC), tetrachloromethane and esachloroethane. The gaseous species were identified through FT-IR spectrophotometry equipped with a cell with long optical path (8 m) and GC-MS. From FT-IR spectra (see Fig. 3-SM) in the 2500-500 cm$^{-1}$ CO$_2$, COCl$_2$, traces of TCAC and other unknown chlorinated compounds were detected (see SM under 3.). The identification of the main products was carried out by comparing the recorded spectra with the spectra of pure gaseous standard.[40] From GC-MS with cryofocusing the residual C$_2$Cl$_4$, the produced CCl$_4$ and C$_2$Cl$_6$ were detected (see Fig. 4-SM).

Fig. 4A shows the concentration of photocatalytic produced carbon tetrachloride in the presence and absence of oxygen. The presence of chlorides has a detrimental effect on the production of CCl$_4$ both with and without O$_2$. The effect is larger under anoxic conditions. The production of CCl$_4$ is strongly inhibited in the presence of oxygen. The concentration of this molecule increased monotonically during the experiments in both cases, but the concentrations reached in anoxic conditions are roughly 1-2 order of magnitude higher than in the presence of oxygen (see Fig. 5-SM). Then the increase of CCl$_4$ with irradiation time is NOT correlated with the concomitant increase of released chloride observed during the photocatalytic degradation of PCE (see Fig. 2B). CCl$_4$ must be released from intermediates formed upon further chlorination, favored under anoxic conditions, and the chlorination is not due to chlorine radical, as discussed before.

During the same experiments the evolution of hexachloroethane was also monitored. As for CCl$_4$, the production of C$_2$Cl$_6$ is totally impeded by an excess of oxygen and favored by anoxic conditions. Conversely, chloride ions at the surface favor its production (Fig. 4B). Given that *Cl radical is not present and that the primary reducing step decrements by one the number of organic chlorine, there is the sole possibility that the carbocation C$^+Cl_2$
carbocation was demonstrated as necessary to justify observed reaction products.[24] The reaction of an anion with a carbocation, actually formed for direct hole transfer to PCE (reaction r1) and with CCl$_3$. 
The mechanism of formation of the detected compounds and of higher MW compounds with conjugated double bonds could be sketched as: 1) the addition of C₂Cl₄ to neutral radical formed upon reduction (r2); 2) the arrangement of the radical by chlorine transposition; 3) loss of CCl₄ (as suggested before). Amongst these steps the oxidation of intermediates by holes forms a carbocation that adds one chloride anion, giving molecules with less double bonds and more chlorine (toward chloroalkanes, as depicted in r4 and r5). Alternatively, amongst these steps, the reduction of intermediates by e⁻ and loss of chloride ion forms a neutral molecule or radical that can undergo the previous reactions. The formation of the detected 1,1,2,3,4,4-hexachlorobuta-1,3-diene is here proposed as an example (see also Fig. 6).

\[ \text{CCl}_2=\text{C}^\ast\text{Cl}_{(ad)} + \text{C}_2\text{Cl}_4 \rightarrow \text{CCl}_2=\text{C}({\text{Cl}}) \]
FIGURE CAPTIONS

**Fig. 1.** (A) Photon efficiency PE=rate/I₀ of PCE transformation as a function of actual concentration in the reactor. Inset: conversion as a function of inlet concentration C₀. Conditions: C₀ = 2-9 ppm, total flow = 3.0 L min⁻¹ (N₂ 80%, O₂ 20%), S = 15 cm², I₀=10 W m⁻². (B) Photon efficiency PE of PCE transformation as a function of actual water vapor concentration. The dashed line shows the data interpolation with eq. 8 (see text). Conditions: C₀ = 3 ppm, total flow = 1.6 L min⁻¹ (N₂ 80%, O₂ 20%), S = 15 cm², I₀=10 W m⁻².

**Fig. 2.** (A) residual concentration of PCE going out of the reactor in the presence of oxygen (N₂ 80% + O₂ 20%) and in quasi-anoxic conditions (N₂ 100 %) as a function of the irradiation time. Experimental conditions: C₀ = 1 ppm, total flow = 1.6 L min⁻¹ (N₂ 80% + O₂ 20% or N₂ 100%), S = 100 cm², I₀=10 W m⁻², volume sampled in the cryogenic trap = 50 mL. (B) Evolution of the surface concentration of Cl⁻ as a function of the irradiation time. Inset: molar fraction χ of C₂Cl₄ converted in chloride ion. Experimental conditions: C₀ = 3 ppm, total flow = 1.6 L min⁻¹ (N₂ 80% + O₂ 20% or N₂ 100%), S = 15 cm², I₀=10 W m⁻².
Fig. 3. (A) Photon efficiency PE=Rate/Io for C₂Cl₄ degradation (after 10 minutes of irradiation) in the presence or absence of oxygen as a function of the amount of chloride ions added (as NaCl) at the TiO₂ film surface. Experimental conditions: C₀ = 3 ppm, total flow = 1.6 L min⁻¹ (N₂ 80% + O₂ 20% or N₂ 100%), S = 100 cm², 10 W m⁻². (B) Concentration of active chlorine flowing out of the reactor as a function of the irradiation time in the presence or absence of added chloride (1000 µmol NaCl) at the TiO₂ film surface. Experimental conditions: C₀ = 10 ppm, total flow = 2.4 L min⁻¹ (N₂ 80% + O₂ 20% or N₂ 100%), S = 100 cm², 10 W m⁻².

Fig. 4. Concentration of formed CCl₄ (A) and Chromatographic peak area for C₂Cl₆ (B) after 10 minutes of irradiation during the photocatalytic transformation of C₂Cl₄ in the presence or absence of oxygen as a function of the amount of chloride ions added (as NaCl) at the TiO₂ film surface. Experimental conditions: C₀ = 3 ppm, total flow = 1.6 L min⁻¹ (N₂ 80% + O₂ 20% or N₂ 100%), S = 100 cm², Io=10 W m⁻².
**Fig. 5.** Mechanism of reaction for the production of \( \text{C}_2\text{Cl}_6 \) during the photocatalytic transformation of gaseous PCE under irradiated TiO\(_2\) in anoxic condition.

**Fig. 6.** Suggested mechanism proposed for the production of polyene compounds at the surface of the photocatalyst in anoxic conditions. The structural analogy between the first and the last radicals of the reaction chain allows the perpetuation of the reaction with the production of compounds with higher molecular weight.
References

A revised photocatalytic transformation mechanism for chlorinated VOCs: experimental evidence from C$_2$Cl$_4$ in the gas phase

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SUPPLEMENTARY MATERIAL

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1 Materials and Methods

1.1 Materials

Titanium dioxide (Hombikat N100: 100% anatase, BET specific surface area 100 m² g⁻¹, average crystal size 20 nm) was purchased from Sachtleben Chemie GmbH (Duisburg, Germany). Perchloroethylene (PCE, 99.9%), Titanium(IV) isopropoxide (97%), N,N-Diethyl-p-phenylenediamine (97%), NaOH (≥98%), Na₂HPO₄ (≥98%) and NaH₂PO₄ (99%) were purchased from Sigma Aldrich and absolute ethanol from Fluka (≥98%). Nitrogen was obtained from the evaporation of pure liquid nitrogen (99.99%, SIAD, Italy) while gaseous oxygen (research grade 99.9%) and chromatographic grade Helium (5.5 grade) were supplied by SIAD (Italy). Gaseous standards of PCE were prepared in 15 L SilcoCanTM (Restek) canisters diluting PCE in nitrogen at the desired concentration. All the compounds were used as received without any further purification step. Water was purified with a MilliQ plus apparatus (TOC = 2 ppb, conductivity 18.2 MΩ cm).

1.2 Preparation of the TiO₂ films

TiO₂ films were prepared as follow. 12.5 g of TiO₂ Hombikat N100 were suspended in 50 mL of a Titanium(IV) isopropoxide (TIP, 0.33 mol L⁻¹) in absolute ethanol. 2 mL of the suspension were deposited on the surface of 10×10 cm² Pyrex glass substrates, in this way 0.55 g of TiO₂ were deposited on each Pyrex glass substrate as sum of the nanocrystalline catalyst (0.50 g) and the amorphous TiO₂ phase obtained from the hydrolysis of TIP (0.05 g). The samples were dyed at 100 °C for 150 min and UV irradiated (18 W m⁻² in the 300-400 nm range) for 18 hours with the aim to photocatalytically remove the organic residues from the catalyst surface. The hydrolysis of TIP is essential to obtain a compact and well anchored films. The TiO₂ film samples were storage in the dark and in a closed box to avoid possible poisoning of the surface before use.

1.3 Methods and experimental set-up

The monitoring of C₂Cl₄ flowing out the reactor was carried out with a Photo Ionization Detector PID-AH (from α-Sense with Kr as filling gas for the glow discharge lamp) which signal was amplified (model AD327 amplifier, Analog device), digitalized and recorded by using an analog-to-digital converter (model NI USB-6221, National Instruments). The volatile VOCs flowing out the reactor were analysed by both GC-MS after cryofocusing and directly by FTIR spectrophotometry. The cryofocusing system was a Model 7000 Entech system able to focus at the temperature of liquid nitrogen the VOCs, eliminate both CO₂ and water vapour, and inject into the chromatographic system the preconcentrated samples. The GC-MS system used for the VOCs determination was a 6890 GC hyphenated with a 5973 Mass Selective Detector (Agilent Technologies). The column was an Agilent CP-Sil 5 CB column (length 60 m, i.d. 0.32 mm, film thickness 0.1 µm). The analyses were carried out starting from 35°C, increasing the temperature at 5 °C/min up to 140°C and from 140°C to 240°C at 15 °C/min.

Volatile by-products of the PCE photocatalytic transformation were also detected through FT-IR (Cary 670 spectrophotometer, Agilent Technologies) equipped with a cell for gases with 8 m long optical path length. Both the cell and the transfer line were thermostated at 150 °C. The FT-IR spectrophotometer was equipped with a KBr beam splitter and a MCT detector.

The organic compounds cumulated at the catalyst surface in the anoxic conditions were determined through extraction with dichloromethane after irradiation of the film of TiO₂ and
analysed through an Agilent 6890 GC equipped with a GERSTEL CIS4 PTV injector (injection volume 2 μL in splitless mode). The column used was an Agilent HP-5MS (length 30 m, i.d. 0.25 mm, film thickness 0.25 μm). The starting oven temperature was 40 °C for 3 min with a ramp of 5°C/min up to 320 °C (maintained for 9 minutes). The mass spectra obtained were interpreted for comparison with the standard spectra reported in the Wiley 7n library (Agilent part No. G1035B). For both the GC analysis the carrier gas was He of chromatographic grade (Sapi, Italy).

The deposition of water soluble anions at the TiO₂ film surface during the photocatalytic transformation of PCE with and without oxygen was evaluated. After different irradiation times in condition of constant flow of PCE, the films were removed from their Pyrex glass supports, extracted with ultrapure water and the aqueous samples analyzed. The determination of water extractable anions cumulated at the catalyst surface was done through ion chromatography (Dionex 500, with GP40 pump, Rheodyne 9126, 50 μL loop, LC30 oven, ASRS-II Ultra conductivity suppressor and ED40 detector) with the analytical column AS9-HC and column guard AG9-HC. The eluent was K₂CO₃ 9 mM, and the flow rate adopted was 1 mL/min keeping the column temperature oven at 30 °C.

Some TiO₂ samples obtained at different irradiation times and conditions were analysed by FT-IR Spectrophotometry to investigate the changes in the surface properties. The samples were pressed in self-supporting pellets (“optical thickness” of ≈ 10 mg cm⁻²) and placed in an IR cell with KBr windows (Aabspec), permanently attached to a vacuum line (residual pressure = 1.0x10⁻⁶ Torr), allowing desorption and thermal treatments to be carried out in situ. We used a Perkin-Elmer System 2000 FT-IR spectrophotometer equipped with a MCT cryodetector able to record spectra in the 7200-580 cm⁻¹ range.

Finally, the UV-Vis reflectance spectra of the film obtained at different irradiation time in the presence of C₂Cl₄ both in oxygen rich and in anoxic conditions were recorded by a Cary 5000 (Varian) spectrophotometer with an integration sphere in polytetrafluoroethylene (PTFE) with a 150 mm diameter. The elaboration of the %R spectra was carried out by the Kubelka-Munk equation using a PTFE standard sample as reference blank.

1.4 Photochemical reactor and related kinetic treatment

The experiments of PCE transformation were carried out through the experimental set up schematized in Fig. 1-SM. The homemade flow-through photo-reactor worked in a regime of perfect mixing (Continuous Stirred Tank Reactor, CSTR). An internal fan allows the absence of any concentration gradient inside the reactor. The reaction chamber (340x180x90 mm) was in poly-methyl-methacrylate (PMMA) sealed with epoxy glue to avoid air leakage. The cover of the reactor was a Pyrex glass 5 mm thick with a full optical transparency in the emission wavelength range of the adopted light source. A set of two Philips PL-S 9W/2P BLB lamps was used as irradiation source with an emission spectra in the 350-400 nm range with a maximum of emission at 365 nm. To modify the UV irradiance reaching the sample the relative distance between the light source and the sample was properly modified. The light irradiance was measured through a photometer purchased by the Italian vendor CO.FO.MEGRA. The radiative power [W m⁻²] was converted in the related photon flux [mol of photons m⁻² s⁻¹] on the basis of the lamp spectrum. A homogeneous irradiation of the whole catalyst surface was always assured. The reactor was built according to the CEN/TS 16980-1:2016 Technical Specification proposed for the Determination of the degradation of nitric oxide (NO) in the air by photocatalytic material.[1] A set of valves allowed the regulation of
the inlet/outlet gas flow and the by-pass of the reactor chamber. The gas flow flowing out the
reactors is properly analysed as reported above. For a detailed description of the testing
procedure see ref. [2].

The mass balance for C\textsubscript{2}Cl\textsubscript{4} in the reactor as a function of the reaction time (for
photochemical transformation the irradiation time \( t \)) is

\[
V_R \frac{d}{dt} C_{out}(t) = F[C_0 - C_{out}(t)] - S \text{Rate}(t)
\]

(eq. 1-SM)

where \( V_R \) is the reactor volume, \( C_{out}(t) \) is the concentration of C\textsubscript{2}Cl\textsubscript{4} going out the reactor at
the irradiation time \( t \), \( F \) is the total flow, \( S \) is the geometric area of the TiO\textsubscript{2} film, \( C_0 \) is the
concentration of the reactant entering the reactor and \( \text{Rate}(t) \) is the transformation rate
normalized for unit area of C\textsubscript{2}Cl\textsubscript{4} in the reactor at the irradiation time \( t \).

In condition of steady state (SS, \( t \rightarrow \infty \) and no change in the photoactivity of the catalyst, e.g. no
poisoning or photo-activation as a consequence of the irradiation) the concentration of C\textsubscript{2}Cl\textsubscript{4}
does not change with the time (\( V_R \frac{d}{dt} C_{out}(t) = 0 \)) and consequently

\[
F[C_0 - C_{out}(t)] - S \text{Rate}^{SS}(t) = 0
\]

(eq. 2-SM)

Defining the conversion at the time \( t \) as

\[
\eta(t) = \frac{C_0 - C_{out}(t)}{C_0}
\]

(eq. 3-SM)

The transformation rate can be expressed as a function of the conversion and so directly
measured from the experimental profiles in steady state conditions (eq. 4-SM) [2].

\[
\text{Rate}^{SS} = \frac{C_0 \eta_s F}{S}
\]

(eq. 4-SM)

The inlet gas flows and their concentrations were set with an Entech-4600A Mass Flow
Controller (MFC) mixing pure N\textsubscript{2}, O\textsubscript{2}, and concentrated C\textsubscript{2}Cl\textsubscript{4} in N\textsubscript{2}. Relative humidity was
set by bubbling dry N\textsubscript{2} in thermostated pure water under a fixed pressure.

A thermohygrometric sensor placed in the reaction chamber allowed the determination of
both the relative humidity and the temperature. The absolute humidity was computed on the
basis of eq. 5-SM [3]

\[
p_{H_2O} = 6.11 \frac{RH\%}{100} \frac{10^{7.5T}}{237.7+T}
\]

(eq.5-SM)

in which \( p_{H_2O} \) is the absolute humidity (atm), \( RH\% \) is the relative humidity % and \( T \) is the
temperature (°C). In order to express the absolute humidity in ppm the value of \( p_{H_2O} \) (atm)
was multiplied for \( 10^6 \) (the atmospheric pressure was considered stable and equal to 1 atm).
Fig. 1-SM Scheme of the experimental apparatus adopted for the investigation of the photocatalytic transformation of C$_2$Cl$_4$. 

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2 Main Intermediates

Evolution of the amount of oxalate (normalized for the film surface area) deposited on the TiO$_2$ surface in the presence or absence of oxygen.

Fig. 2-SM Evolution of the amount of oxalate (normalized for the film surface area) deposited on the TiO$_2$ surface in the presence or absence of oxygen as a function of the irradiation time during the photocatalytic transformation of gaseous C$_2$Cl$_4$. Experimental conditions: C$_0$ = 3 ppm, total flow = 1.6 L min$^{-1}$ (N$_2$ 80% + O$_2$ 20% or N$_2$ 100%), S = 15 cm$^2$, 10 W m$^{-2}$.
3 Detection of gas species

The gaseous species were identified through FT-IR spectrophotometry with long optical path cell (8 m) and GC-MS.

In FT-IR spectra (see Fig. 3-SM) there is the absence of the two main IR signals of the PCE at 914 and 780 cm\(^{-1}\) due to the abatement of C\(_2\)Cl\(_4\) at concentration lower than the limit of detection of the technique. The main products observed both in the presence and absence of oxygen are carbon dioxide (see the doublet with maxima at 2330 and 2360 cm\(^{-1}\) related to its asymmetric stretching) and phosgene (main peaks at 794, 846, 856, 1016, 1680, 1818 and 1834 cm\(^{-1}\)). Furthermore, traces of both TCAC and other chlorinated compounds can be hypothesized due to the presence of other minor peaks (e.g. 794, 668 and 569 cm\(^{-1}\)) not ascribable to the main transformation products.

The analysis of the by-products formed during the photocatalytic transformation of PCE were carried out also by GC-MS through cryofocusing at the temperature of the liquid nitrogen. The chromatograms obtained (an example is reported in Fig. 4-SM) are dominated by 3 main peaks related to the presence of the residual C\(_2\)Cl\(_4\) (\(t_r = 14.6\) min), the photocatalytically produced CCl\(_4\) (\(t_r = 12.8\) min) and C\(_2\)Cl\(_6\) (\(t_r = 28.7\) min). The identification of the nature of these compounds were carried out on the basis of the mass spectra (matching between the recorded spectra and the spectra reported in the NIST library \(\geq 95\%\)). Note that the absence of phosgene in the chromatogram can be easily explained considering that this compound reacts in the focusing system with the traces of water producing CO\(_2\) and HCl and consequently cannot be determine by the adopted GC-MS system.

Fig. 3-SM FT-IR spectra in the 2500-500 cm\(^{-1}\) range of the gaseous species flowing out of the reactor chamber during the photocatalytic transformation of C\(_2\)Cl\(_4\) in the presence of oxygen (red line) and in quasi-anoxic conditions (blue line). Experimental conditions: C\(_0\) = 20 ppm, total flow = 2.0 L min\(^{-1}\) (N\(_2\) 80% + O\(_2\) 20% or N\(_2\) 100%), S = 100 cm\(^2\), 10 W m\(^{-2}\), resolution 4 cm\(^{-1}\).
Fig. 4-SM GC analysis of the gaseous species flowing out of the reactor during the C$_2$Cl$_4$ photocatalytic transformation under UV-irradiated TiO$_2$ film. Example of chromatogram (Selective Ion Monitoring (SIM) mode, m/z 117, 166 and 201) and Mass Spectra of the main compounds. For the three compounds identified the related spectra present in the MS library Wiley 7n library were reported together with the goodness of fit (≥95%). Experimental conditions: C$_0$ = 1 ppm, total flow = 1.6 L min$^{-1}$ (N$_2$ 80% + O$_2$ 20%), S = 100 cm$^2$, 10 W m$^{-2}$, volume sampled in the cryogenic trap = 50 mL.
Fig. 5-SM Concentration of the photocatalytically generated CCl$_4$ (B) in the presence of oxygen (N$_2$ 80% + O$_2$ 20%) and in anoxic conditions (N$_2$ 100 %) as a function of the irradiation time. Experimental conditions: C$_0$ = 1 ppm, total flow = 1.6 L min$^{-1}$ (N$_2$ 80% + O$_2$ 20% or N$_2$ 100%), S = 100 cm$^2$, 10 W m$^{-2}$, volume sampled in the cryogenic trap = 50 mL.

4 Surface species

4.1 UV-Vis reflectance measurements and poisoning

Fig. 6-SM Evolution of the TiO$_2$ film surface during the photocatalytic transformation of C$_2$Cl$_4$ in quasi anoxic-condition under prolonged UV-irradiation (from 0 to 24 hours of continuous irradiation). A) UV-Vis reflectance spectra of the TiO$_2$ films; B) Pictures of the films obtained after different irradiation times (note the progressive evolution of the surface color toward pale yellow). Experimental conditions: C$_0$ = 10 ppm, total flow = 2.1 L min$^{-1}$ (N$_2$ 100%), S = 100 cm$^2$, 10 W m$^{-2}$.
4.2 Conversion in CSTR in anoxic conditions

During irradiation the experimental conversion in CSTR (after the first transient) decreases, due to poisoning. This effect can’t be diagnosed in batch experiments.

**Fig. 7-SM** Residual concentration of C$_2$Cl$_4$ under anoxic-condition as a function of the irradiation time. Experimental conditions: C$_0$ = 10 ppm, total flow = 2.1 L min$^{-1}$ (N$_2$ 100%), S = 100 cm$^2$, 10 W m$^{-2}$.

**Fig. 8-SM** Effect of the irradiation in the presence of oxygen on the TiO$_2$ film obtained at the end of a long time irradiation in the presence of C$_2$Cl$_4$ in quasi anoxic-condition. Note that the yellow-pale compounds deposited during in the anoxic phase disappeared during the irradiation in the presence of oxygen. Experimental conditions: C$_0$ = 10 ppm, total flow = 2.1 L min$^{-1}$ (N$_2$ 100%), S = 15 cm$^2$, 10 W m$^{-2}$. 

hv, O$_2$
4.3 FT-IR spectroscopy of surface species

Film samples at different irradiation times were removed from their Pyrex glass substrate and pressed as self-supported pellets. The spectra of the samples in the 1000-4000 cm\(^{-1}\) range were recorded after out-gassing at room temperature and at 50 °C for 30 minutes. The spectra of samples before and after 6 hour of irradiation in the presence of oxygen almost overlap (Fig. 9A-SM), indicating a negligible accumulation. The comparison among the spectra obtained after 0, 6, 15 and 24 hours of irradiation in anoxic conditions (Fig. 9B-SM) show a progressive increment of the intensity of: 1) the peak at 1620 cm\(^{-1}\) is related to the bending of the water molecules adsorbed at the TiO\(_2\) surface; 2) a new peak formed at 1370 cm\(^{-1}\); 3) the broad absorption band spread over the 3550-2500 cm\(^{-1}\) interval and the partly resolved components between 3750 and 3550 cm\(^{-1}\).

As already stated, water traces are always present in the experimental system. The peaks in 3550-2500 cm\(^{-1}\) interval are attributed to O–H stretching bond of hydroxyl surface groups (linear or bridged) and adsorbed water. The pattern at 3750-3550 cm\(^{-1}\) is commonly attributed to free \(\nu OH\) oscillators while the broad band at 3550-2500 cm\(^{-1}\) to H-bonded \(\nu OH\) oscillators.[4,5] The increment of the intensity of these signals, together with the concomitant increment of the 1620 cm\(^{-1}\) peak can be related to the increase of the hydration state of the TiO\(_2\) surface. The higher amount of surface water can be related to the presence at the surface of species able to increase the Lewis acidity, and consequently to increase the ability to adsorb water molecules [5], and to the production of water soluble species (e.g. Cl\(^-\)), which are able to decrease the vapor tension of the water multilayers located at the TiO\(_2\) surface, and consequently hinder their desorption.

The 1370 cm\(^{-1}\) peak can be attributed to the vibration modes of C–Cl bond closed to carbon/carbon double bonds. As an example, 3-chloropropene shows an intense absorption band in the same region with a maximum located at 1417 cm\(^{-1}\).[6] Then compounds of similar structure are cumulated, most probably with increased conjugated double bonds as Vis absorption is developed with irradiation time.

**Fig. 9-SM** Evolution of the FT-IR spectra of the TiO\(_2\) films removed from the support and analyzed as self-supported pellets during the photocatalytic transformation of C\(_2\)Cl\(_4\). (A) Comparison between the samples before and after 6 hour of irradiation in the presence of oxygen; (B) Comparison among the spectra before and after 6, 15 and 24 hours of irradiation in quasi-anoxic conditions. Experimental conditions: C\(_0\) = 10 ppm, total flow = 2.1 L min\(^{-1}\) (N\(_2\) 80% + O\(_2\) 20% or N\(_2\) 100%), S = 100 cm\(^2\), 10 W m\(^{-2}\).
4.4 GC-MS analysis of surface species

Fig. 10-SM Gas chromatogram (TIC mode) of the sample obtained extracting with CH₂Cl₂ the TiO₂ film after the photocatalytic transformation of C₂Cl₄ in quasi-anoxic condition (C₀ = 10 ppm ppm, flow = 2100 cm² (N₂ 100%), S = 100 cm³). The peaks of the five main compounds formed are highlighted in red.
Table 1-SM Retention time, molecular weight and molecular structure of the 5 compounds formed and deposited on the TiO$_2$ film during the photocatalytic transformation of C$_2$Cl$_4$ under irradiated TiO$_2$ film in quasi-anoxic conditions ($C_0 = 1$ ppm, flow = 1600 cm$^2$ (N$_2$ 100%), $S = 100$ cm$^2$).

<table>
<thead>
<tr>
<th>$t_r$ (min)</th>
<th>Compounds</th>
<th>Molecular weight</th>
</tr>
</thead>
<tbody>
<tr>
<td>6.81</td>
<td><img src="cl8.png" alt="" /> $1,1,2,3,3$-hexachloropropene</td>
<td>249</td>
</tr>
<tr>
<td>6.85</td>
<td><img src="cl9.png" alt="" /> $1,1,2,3,4,4$-hexachlorobuta-1,3-diene</td>
<td>261</td>
</tr>
<tr>
<td>8.24</td>
<td><img src="cl10.png" alt="" /> perchloropropane</td>
<td>320</td>
</tr>
<tr>
<td>9.24 - 9.84</td>
<td><img src="cl11.png" alt="" /> Trans-1,1,2,3,4,4-ottachlorobut-2-ene</td>
<td>332</td>
</tr>
<tr>
<td></td>
<td><img src="cl12.png" alt="" /> Cis-1,1,2,3,4,4-ottachlorobut-2-ene</td>
<td></td>
</tr>
</tbody>
</table>
Fig. 11-SM Mass spectra of the molecules formed during the photocatalytic transformation of C₂Cl₄ under irradiated TiO₂ film in quasi-anoxic conditions (C₀ = 1 ppm ppm, flow = 1600 cm² (N₂ 100%), S = 100 cm²). For the compounds with τᵣ = 6.81 and 6.85 min the spectra present in the MS library Wiley 7n library for the related compounds were reported for comparison with the experimental data.
Fig. 12-SM. Area of the peaks of species formed during the photocatalytic transformation of C₂Cl₄ and deposited on the TiO₂ films under anoxic condition as a function of the irradiation time. Experimental conditions: C₀ = 10 ppm, total flow = 2.1 L min⁻¹ (N₂ 100%), S = 100 cm², I₀=10 W m⁻².
5 References