Photocatalytic degradation of atrazine by heteropolyoxotungstates

Loubna Youssef, Ghassan Younes and Rami Al-Oweini

Department of Chemistry, Faculty of Science, Beirut Arab University, Beirut, Lebanon

ABSTRACT
An elegant and eco-friendly photocatalytic degradation approach for the most commonly used herbicide, Atrazine (ATZ); allowed a selective and partial degradation into atrazine-2-hydroxy (Hydroxyatrazine, HAT), which was the sole obtained metabolite. The effect of tungstosilicates and tungstophosphates was investigated as photocatalyst under UV irradiation at two different wave-lengths; 254 and 366 nm; was investigated by measuring the absorbance at 10-min intervals, using 720-UV spectrophotometry. ATZ conversion rate was the highest when irradiating with the higher-energy 254 nm UV light. This resulted into 54% degradation in the presence of the silicon-based α-Keggin dodecatungstosilicate (α-SiW12O40)4−, whereas 17% was decomposed in its absence. The nature of the heteroatom as well as the structural type of the studied polyoxometalates had a significant effect on the degradation percentage. Accordingly, 31% of ATZ was only decomposed in presence of the phosphorus-analogue (α-PW12O40)4− whereas the cyclic super-lacunary octatetracontatungstooctaphosphate ([P8W48O184]40) increased the degradation to 41%.

1. Introduction
The contamination of our water supplies as well as our food chain by various persistent organic pollutants has emphasized the attention of many researchers since several decades [1–8]. Herbicides are phototoxic chemicals that have been identified as organic pollutants due to their ubiquitous influence on the environment and their ability to remain nondegradable for long periods of time [9–12]. Atrazine, 6-chloro-N-ethyl-N’-(1-methylethyl)-triazine-2,4-diamine (ATZ), was ranked as the second mostly used agricultural herbicide [13–16]. Its usage has heavily intensified in many countries especially in the Middle East region [17]. Moreover, this pesticide was found to be an endocrine disruptor that can affect reproduction and development [18]. It is highly persistent in the environment, lasting from days up to several years, due to the stability of the s-triazine ring that doesn’t undergo any natural degradation. Therefore, the use of this herbicide is restricted in many countries nowadays [19]. The US Environmental Protection Agency announced that the legal limit for ATZ in drinking water is just 3 ppb [20]. In spite of these regulations, ATZ have been reported to exceeding the recommended value in many countries [21–23].

UV irradiation has been an alternative technology for water purification and treatment of various hazardous contaminants in the presence of different catalysts. Many studies have reported the use of ozone gas, hydrogen peroxide, zinc oxide and titanium oxide as photocatalysts for the degradation of Atrazine by both oxidative and reductive pathways [24–27]. Accordingly, the degradation of ATZ can produce 11 different metabolites depending on the used degradation process [28]. Table 1 summarizes the obtained metabolites from different degradation processes. Our study adds yet another approach, however unique, for the degradation of ATZ using for the very first time a highly-selective and discrete metal–oxo cluster facilitating the degradation into only one major metabolite, atrazine-2-hydroxy (HAT), which can be easily monitored.

Hence, metal oxide clusters, known as polyoxometalates (POMs), have drawn the attention of many researchers as photocatalysts in various reactions of both organic and inorganic pollutants [35–37]. It has been found that these POMs offer many advantages as homogeneous photocatalysts, mainly due to being highly redox active [38–41]. Additionally, POMs experience strong light absorption in the UV region as well as in the visible region for those containing molybdenum and vanadium [42]. The irradiation of POM with UV light below 400 nm activates their oxidizing ability by the formation of OH radicals that are able to decompose a certain organic substrate, denoted here by “S”, according to the following scheme. Furthermore, the structural
Table 1. Atrazine degradation products of some reported degradation methods in the literature.

| Degradation method                                    | Degradation product                                                                 | Reference |
|-------------------------------------------------------|--------------------------------------------------------------------------------------|-----------|
| Photolytic, Sonolytic and Sonophotolytic degradation   | HIE, CVT, OVT, OIE, CDIT, COET, OOT, ODEIT, CIAT, OODIT, ODE...
Figure 2. (a,b) The degradation percentage of ATZ in the absence and presence of photocatalyst under UV irradiation at 366 and 254 nm, respectively.

(Figure 2(b)) since the energy of a UVC light is much higher than that of UVA causing faster degradation and more efficient processes [5].

The effect of nonatungstophosphate (PW₉) and nonatungstosilicate (SiW₉) on the decomposition of the ATZ herbicide is illustrated in Figure 3(a,b), respectively, under 254 nm UV irradiation. These two metal oxide clusters belong to the same Keggin structural family, both of which are trilacunary. The behaviour of both catalysts is the same till 30 min of irradiation. As time proceeds the tungstosilicate succeeded to degrade 14% of ATZ whereas the presence of tungstophosphate resulted in 11% degradation.

The plenary and complete Keggin-structured polyoxometalates (PW₁₂) and (SiW₁₂) have been investigated as photocatalysts for the treatment of Atrazine. Their effect is shown in Figure 4(a,b), respectively. The concentration of ATZ drops directly after 10 min of irradiation in the presence of dodecatungstosilicate to reach 0.93 × 10⁻⁵ M at 60 min. The degradation percentage reaches 54% in the presence of this photocatalyst. However, the dodecatungstophosphate catalyst reaches a modest 31% of ATZ degradation.

On the other hand, the effect of the cyclic superlacunary octatetracontatungstooctaphosphate salt (P₈W₄₈) was also examined in this study. This polyanion was able to degrade 41% of ATZ under the studied experimental conditions (Figure 5). The behaviour of this catalyst in the reaction mixture is somehow different than the other photocatalysts in our work. The concentration of ATZ gradually dropped in a linear manner after 20 min of UV irradiation.

As seen from the above results, the Si-based POMs gave better results than P-based ones. This can be attributed to the important influence of the heteroatom nature on the photocatalytic activity of the
The bond length of X-O affects the electron density on the W-Oaxial that in turn highly affects the redox properties of these metal oxide clusters. The long Si-O bond lengths compared to that of P-O increases the electron density at the tungsten atoms in the Keggin-structured [SiW12] [52].

The degradation product resulting from the photodegradation of ATZ in the presence of [SiW12] as the best photocatalyst in our study was determined. As shown in Figure 6, the only peak appearing at t0 is related to ATZ whose maximum wavelength was found to be 222 nm. However, at the end of our experiment the absorbance at 222 nm drops indicating a loss in ATZ concentration and a new peak forms at 244 nm. This new peak is due to the formation of HAT [27] as the only metabolite for the ATZ decomposition under our experimental conditions.

4. Conclusion

Polyoxometalates, being discrete, environmentally benign, and thermally stable, are promising photocatalysts for the selective degradation of Atrazine to Atrazine-2-hydroxy, as the only produced metabolite. The use of Keggin-type dodecatungstooctaphosphate polyanion resulted in 54% decomposition under simple experimental conditions, which could be easily mimicked under natural circumstances. The catalytic effect of octatetracontatungstooctaphosphate polyanion [P8W48] on the degradation of atrazine was firstly reported here. This complex succeeded to degrade 41% of the studied pollutant. UVC irradiation showed better effect than UVA on the catalytic activity of the studied polyoxometalates. Finally, the kinetic behaviour of these polyanions as well as other affecting parameters on the photodegradation of Atrazine is going to be reported in our next paper.

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Disclosure statement

No potential conflict of interest was reported by the authors.

ORCID

Loubna Youssef http://orcid.org/0000-0002-5503-5708
Ghassan Younes http://orcid.org/0000-0001-6927-3523
Rami Al-Oweini http://orcid.org/0000-0003-1891-3716

References

1. Readman JW, Albanis TA, Barcelo D, et al. Herbicide contamination of Mediterranean estuarine waters: results from a MED POL pilot survey. Mar Pollut Bull. 1993;26:613–619. doi:10.1016/0025-326X(93)90500-J
2. Gray N. Drinking water quality. problems and solutions. Chichester: John Wiley & Sons; 1994. p. 132–148.
3. Encyclopedia of surface and colloid science. New York: Marcel Dekker; 2002.
4. Ali I, Aboul-Enein HY. Instrumental methods in metal ions speciation: chromatography, capillary electrophoresis and electrochemistry. New York: Taylor & Francis; 2006.
5. Chen C, Yang SG, Guo YP, et al. Photolytic destruction of endocrine disruptor atrazine in aqueous solution under UV irradiation: products and pathways. J Hazard Mater. 2009;172:675–684.
6. Ali I, Aboul-Enein HY, Gupta VK. Nanochromatography and nanocapillary electrophoresis: pharmaceutical and environmental analyses. Hoboken (NJ): Wiley & Sons; 2009.
7. Gupta VK, Ali I. Environmental water: advances in treatment, remediation and recycling. Amsterdam: Elsevier; 2012.
8. Basheer AA. Chemical chiral pollution: impact on the society and science and need of the regulations in the 21st century. Chirality. 2018;30(4):402–406. doi:10.1002/chir.22808
9. Ali I, Jain CK. Groundwater contamination and health hazards by some of the most commonly used pesticides. Curr Sci. 1998;75(10):1011–1014.
10. Ali I, Aboul-Enein HY. Determination of chiral ratio of o,p-DDT and o,p-DDD pesticides on polysaccharides chiral stationary phases by HPLC under reversed-phase mode.
[10] Gupta PK. Reproductive and developmental toxicology. Academic Press: Elsevier; 2011. p. 503–521.

[11] Ali I, ALOthman ZA, Al-Warthan A. Sorption, kinetics and thermodynamics studies of atrazine herbicide removal from water using iron nano-composite material. Int J Environ Sci Technol. 2016;13(2):733–742. doi:10.1007/s13762-015-0199-6

[12] Nesnovik NK, Elezović I, Karan V, et al. Acute and subacute toxicity of atrazine to carp (Cyprinus carpio L.). Ecotoxicol Environ Safety. 1993;25:173–182.

[13] Jain CK, Ali I. Determination of pesticides in water, sediments and soils by gas chromatography. Int J Environ Anal Chem. 1997;68(1):83–101. doi:10.1080/03067319708030482

[14] Pathak RK, Dikshit AK. Atrazine and its use. Int J Res Chem Environ. 2012;2:1–6.

[15] Hofman RS, Capel PD., Larson SJ. Comparison of pesticides in eight U.S. urban streams. Environ Toxicol Chem. 2000;19:2249–2258.

[16] Ali I, Al-Othman ZA, Al-Warthan A. Removal of secbumeton herbicide from water on composite nanoadsorbent. Desalin Water Treat. 2016;57(22):10409–10421. doi:10.1080/19443994.2015.1041164

[17] Canny J, Teze A, Thouvenot R, et al. Disubstituted tungstosilicates. 1. Synthesis, stability, and structure of the lacunary precursor polyanion of a tungstosilicate. J. Cluster Sci. 2009;919(1-3):140–145. doi:10.1016/j.molstruc.2008.08.025

[18] Kortz U. Polyoxometalates. Eur J Inorg Chem. 2009;2009(214):336–342.

[19] Kortz U, Müller A. Introduction: a special issue dedicated to Michael T. Pope. J Cluster Sci. 2006;17(2):139–141. doi:10.1007/s10876-006-0065-x

[20] Kortz U, Müller A, van Slageren J, et al. Polyoxometalates: fascinating structures, unique magnetic properties. Coord Chem Rev. 2009;253(19-20):2315–2327. doi:10.1016/j.ccr.2009.01.014

[21] Kortz U, Liu T. The best of polyoxometalates. Eur J Inorg Chem. 2013;2013(10-11):1559–1560. doi:10.1002/ejic.201300230

[22] Al-Oweini R, Aghyar S, El-Rassy H. Immobilized polyoxometalates onto mesoporous organically-modified silica aerogels as selective heterogeneous catalysts of anthracene oxidation. J Sol-Gel Sci Technol. 2012;61(3):541–550. doi:10.1007/s10971-011-2657-7

[23] Al-Oweini R, Sartorel B, Bassil BS, et al. Photocatalytic water Oxidation by a mixed-valent MnIII3MnIVO3 manganese core that mimics the Natural Oxygen-Evolving Center. Angewandte Chem - International Edition. 2014;53(42):11182–11185. doi:10.1002/anie.201404664

[24] Natai M, Bazzan I, Gobena-Ferron S, et al. Photo-assisted water oxidation by high-nuclearity cobalt-oxo cores: tracing the catalyst fate during oxygen evolution turnover. Green Chem. 2017;19(10):2416–2426. doi:10.1039/c7gc00052a

[25] Canny J, Teze A, Thouvenot R, et al. Disubstituted tungstosilicates: 1. Synthesis, stability, and structure of the lacunary precursor polyolavan of a tungstosalicylate. gamma-SiW10O36–6H2O. Inorg Chem. 1986;25:2114–2119.

[26] Ginsberg AP. Inorganic syntheses. Berkeley Heights (NJ): Wiley-Interscience; 1990; Vol. 27.

[27] Al-Oweini R, El-Rassy H. Synthesis and characterization by FTIR spectroscopy of silica aerogels prepared using several Si(OR)4 and R′Si(OR)3 precursors. J Mol Struct. 2009;919(1-3):140–145. doi:10.1016/j.molstruc.2008.08.025
[48] Al-Oweini R, Bassil BS, Palden T, et al. The manganese(III)-
containing tungstophosphate [MnIII3(\(\text{H}_2\text{O}\))5(\(\text{A-}\alpha-\text{PW}_9\text{O}_{34}\))2]9. Polyhedron. 2013;52:461–466. doi:10.1016/j.poly.2012.08.050

[49] Al-Oweini R, Bassil BS, Friedl J, et al. Synthesis and
characterization of multinuclear manganese-containing

tungstosilicates. Inorg Chem. 2014;53(11):5663–5673. doi:10.1021/ic500425c

[50] Al-Oweini R, Bassil BS, Itani M, et al. The mixed-valent 10-
manganese(III/IV)-containing 36-tungsto-4-arsenate(V),
[MnII6MnIV4(OH)12(\(\text{H}_2\text{O}\))12(\(\text{A-}\beta-\text{AsW}_{9}\text{O}_{34}\))4]22.

[51] Poblet JM, López X, Bo C. Ab initio and DFT modelling of
complex materials: towards the understanding of elec-
tronic and magnetic properties of polyoxometalates.
Chem Soc Rev. 2003;32:297–308.

[52] Allmen KV, Moré R, Miller R, et al. Nickel-containing
Keggin-type polyoxometalates as hydrogen evolution
catalysts: Photochemical structure-activity relationships.
ChemPlusChem. 2015;80:1389–1398.