We are IntechOpen, the world's leading publisher of Open Access books Built by scientists, for scientists

5,900 Open access books available 145,000

180M



Our authors are among the

TOP 1% most cited scientists





WEB OF SCIENCE

Selection of our books indexed in the Book Citation Index in Web of Science™ Core Collection (BKCI)

Interested in publishing with us? Contact book.department@intechopen.com

Numbers displayed above are based on latest data collected. For more information visit www.intechopen.com



Competition Kinetics: An Experimental Approach

Murtaza Sayed, Luqman Ali Shah, Javed Ali Khan, Noor S. Shah, Rozina Khattak and Hasan M. Khan

Additional information is available at the end of the chapter

http://dx.doi.org/10.5772/intechopen.70483

Abstract

In this chapter, free radical kinetics with the help of competition kinetics and some experimental results calculated by competition kinetics to find out the rate constant of reactive species ($^{\circ}OH$, e_{aq}^{-} , $^{\circ}H$) with target compound, which is used by radiation chemists is briefly discussed. The competition kinetics method is well validated by taking ciprofloxacin, norfloxacin and bezafibrate as example compounds. The bimolecular rate constants of hydroxyl radical, hydrate electron and hydrogen atom has been calculated for example solute species (ciprofloxacin, norfloxacin and bezafibrate).

Keywords: competition kinetics, rate constants, norfloxacin, ciprofloxacin, bezafibrate

1. Introduction

Radiation chemistry involves extensive study of competition between fast reactions of transient species, reactive and intermediates. Such knowledge is useful to investigate the mechanism of a radiolytic reaction and to propose which process is taking place and which experimental condition is governing a reaction and to know the chemical kinetics of a radiolytic reaction under study.

Generally, in a chemical process, the reactant is converted to products in an individual step. However, in a radiation induced chemical reaction, all steps are taken into consideration including deposition of energy by a charged particle in the system and then formation of a final stable chemical product, and certainly will be a rather complex set of reactions [1]. In the following sections, we will briefly discuss the fast kinetics, i.e. competition kinetics to find the unknown rate constants of a compound with reactive species like hydroxyl radical ($^{\circ}$ OH) or hydrated electron (e_{aq}^{-}), by considering a reference compound whose rate constant with these reactive species is already known.



© 2018 The Author(s). Licensee InTech. This chapter is distributed under the terms of the Creative Commons Attribution License (http://creativecommons.org/licenses/by/3.0), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited. [cc] BY

2. Competition kinetics

For detailed investigation of competing reactions, it is necessary to have a good knowledge about the rate constant data that will used to propose which reaction is predominant. For instance, the Fricke dosimeter contains three main active species (350 mol m⁻³ H⁺, 1 mol m⁻³ Fe²⁺ and 0.25 mol m⁻³ O₂) that have very high rate constants with e_{aq}^{-} and to find out the reaction mechanism involved in the dosimetry, it is necessary to investigate which solute(s) will mainly react with e_{aq}^{-} .

The reactions and their corresponding rate constants are given as [1–3]:

$$e_{aq}^{-} + H^{+} \to H$$
 $k_{1} = 2.3 \times 10^{10} \text{ M}^{-1} \text{s}^{-1}$ (1)

$$e_{aq}^{-} + Fe^{2+} \rightarrow Fe^{+}$$
 $k_2 = 1.6 \times 10^8 \text{ M}^{-1} \text{s}^{-1}$ (2)

$$e_{aq}^{-} + O_2 \rightarrow O_2^{-}$$
 $k_3 = 1.9 \times 10^{10} \,\mathrm{M}^{-1} \mathrm{s}^{-1}$ (3)

The extent of reaction is proportional to the product *k*[solute] for each of the three solutes, that is,

$$e_{aq}^{-} + H^{+}: e_{aq}^{-} + Fe^{2+}: e_{aq}^{-} + O_{2} = k_{1}[H^{+}]: k_{2}[Fe^{2+}]: k_{3}[O_{2}] = 5 \times 10^{4}: 1:30$$
(4)

Alternatively, it can also be concluded that 99.94% of the hydrated electrons reacting with the three solutes will react with H^+ , so under such conditions the reaction of e_{aq}^- with Fe^{2+} and O_2 will be ignored. Therefore, it is compulsory to have an wide collection of rate constant data to apply kinetics for a radiation induced chemical reaction.

In Fricke dosimeter, hydrogen ions are considered as strong scavengers of hydrated electrons [4–6]. The effectiveness of a chemical scavenger depends upon the product *k*[scavenger] that must have a higher value than *k*[substrate]. For example, *tert*-butanol is used to scavenge hydroxyl radicals and by using concentration of 1 mol m⁻³ *tert*-butanol it has *k*[tert-butanol] = $6.0 \times 10^5 \text{ s}^{-1}$. Therefore, *tert*-butanol would be an efficient scavenger for hydroxyl radical (can scavenger over 99% of the hydroxyl radical), if *k*[solute] for the reaction of hydroxyl radical with the solute is less than $6.0 \times 10^5 \text{ s}^{-1}$ [7]. Similarly, oxygen is used to scavenge hydrated electrons and hydrogen atoms from aqueous media [5, 8, 9] and for both the radical (e_{aq}^{-} and $^{\circ}$ H) in air-saturated media, *k*[O₂] = $5 \times 10^6 \text{ s}^{-1}$, so that oxygen can be expected to interfere in the radiolysis of aqueous media if *k*[solute] for the reactions of hydrogen atoms and hydrated electrons with the solute are of the same order as, or less than, $5 \times 10^6 \text{ s}^{-1}$.

In case of radiolysis of organic species, their products also itself often act as scavengers and it is commonly found that the product yield is not in direct relation with the absorbed dose. To estimate the possible reasons of such effects, competition kinetics can be employed in an effective way if the radiolysis mechanism is known and the necessary rate constants are available. For example, cyclohexene is produced when cyclohexane is irradiated and both cyclohexene and cyclohexane have appreciable rate constants with hydrogen atoms, one of the radical specie produced during gamma radiolysis of aqueous media. The reactions are summarized below as:

$${}^{\bullet}\text{H} + \text{C}_{6}\text{H}_{12} \to \text{H}_{2} + {}^{\bullet}\text{C}_{6}\text{H}_{11} \qquad k_{4} = 3.0 \times 10^{7} \text{ M}^{-1}\text{s}^{-1}$$
 (5)

$${}^{\bullet}H + C_6 H_{10} \rightarrow {}^{\bullet}C_6 H_{11} \qquad \qquad k_5 = 3.0 \times 10^9 \text{ M}^{-1} \text{s}^{-1} \tag{6}$$

Under such conditions, the hydrogen atoms will be reacting equally with cyclohexane and cyclohexene when k_4 [cyclohexane] = k_5 [cyclohexene] or in other words we can say that only when 1% of the cyclohexane has been converted to cyclohexene.

Furthermore, competition kinetics is also employed when measuring rate constants by pulse radiolysis using a reference compound.

2.1. Computation of bimolecular rate constant of 'OH with ciprofloxacin

Ciprofloxacin (CIP) belongs to a class of fluoroquinolone family and is used globally as a human and veterinary medication [10]. It has been very much concentrated that the event of these wide range antibiotics in the water bodies may position genuine dangers to the environment and human wellbeing by producing expansion of bacterial medication inactivation. The natural event of these fluoroquinolones anti-infection agents in numerous nations, similar to Switzerland, Australia and China have been affirmed in recent literature [10–13]. It has likewise been watched that most quinolone antibiotics are not completely utilized in the human body and accordingly are discharged and acquainted with the amphibian condition through wastewater sewages because of poor execution of ordinary water treatment plants [12, 14–16] bringing about adversative impacts to sea-going microorganisms and fish [17, 18]. Thus, it becomes necessary to advice alternative physiochemical techniques for effective removal of these contaminants and diminish their ecological effects [12, 19]. For this reason, the deterioration of ciprofloxacin (CIP) in water utilizing ionizing radiations was evaluated to examine the rate constant of [•]OH with CIP. The degradation curves of CIP by gamma irradiation at various absorbed doses has been shown by **Figure 1**.

For computation of bimolecular rate constant of ${}^{\bullet}$ OH with CIP by competition kinetics, phenol was selected as reference compound that has second order rate constant of $6.6 \times 10^9 \text{ M}^{-1} \text{ s}^{-1}$ with ${}^{\bullet}$ OH [20]. The sample solution of total 150 mL having CIP and reference compound phenol together in equivalent quantity was immersed with oxygen gas to change over e_{aq}^{-1} and ${}^{\bullet}$ H to superoxide radical anions quickly [21], which are less responsive compared to ${}^{\bullet}$ OH.

$$CIP + \bullet OH \xrightarrow{k_{CIP}} Products$$
 (7)

$$Phenol + \bullet OH \xrightarrow{R_{phenol}} Products \tag{8}$$

While, the notations, k_{CIP} and k_{phenol} denotes the bimolecular rate constants of [•]OH with CIP and phenol, respectively. Keeping the condition that the dose rate (DR) was kept constant, the rate of decay of CIP is directly related to the rate constant and the concentration of [•]OH [21]:

So,

$$-\frac{d[CIP]}{dD} = -\frac{1}{DR}\frac{d[CIP]}{dt}$$
(9)



Figure 1. The UV spectra of CIP solution observed by gamma irradiation at various absorbed doses ranging from 0 to 870 Gy. Inset shows the influence of gamma-irradiation on degradation of 4.6 mg L^{-1} of CIP solution [8].

Or,

$$-\frac{d}{dD}[CIP] = -\frac{1}{DR}k_{CIP}[CIP][^{\bullet}OH]$$
(10)

Similarly,

$$-\frac{d}{dD}[Phenol] = -\frac{1}{DR}k_{phenol}[Phenol][^{\bullet}OH]$$
(11)

While the absorbed ionizing dose and the total time for which irradiation was performed, are represented by "D" and "t," respectively. Subsequently, the original concentrations of both CIP and phenol are same. Therefore, the rate of decay of CIP to phenol would be equal to the ratio of their individual rate constants as follows [8]:

$$\frac{-\frac{d}{dD}[CIP]}{-\frac{d}{dD}[phenol]} = \frac{k_{CIP}}{k_{phenol}}$$
(12)

Or,

$$\ln \frac{[CIP]_D}{[CIP]_0} = \frac{k_{CIP}}{k_{phenol}} \ln \frac{[phenol]_D}{[phenol]_0}$$
(13)

In Eq. (13), at time 0, the concentration of CIP and phenol are represented by $[CIP]_0$ and $[Phenol]_0$, respectively; while after absorbed dose "D" of gamma irradiation, the corresponding concentration of CIP and phenol are represented by $[CIP]_D$ and $[Phenol]_D$, respectively.



Henceforth, the bimolecular rate constant of ${}^{\bullet}$ OH with CIP was calculated to be $2.75 \times 10^9 \text{ M}^{-1} \text{ s}^{-1}$. Dodd et al. [22] likewise ascertained apparent second order rate constant of with ${}^{\bullet}$ OH-radical with CIP to be $4.1 \ (\pm 0.3) \times 10^9 \text{ M}^{-1} \text{ s}^{-1}$, which is somewhat higher than the esteem calculated in the current report [8].

2.2. Computation of $^{\bullet}OH$, e_{aq}^{-} and $^{\bullet}H$ with norfloxacin

Norfloxacin (NORO) may be called likewise chemotherapeutic antibacterial agent, furthermore is regularly utilized to medicine for urinary tract infections [23]. Its occurrence in surface water and wastewater overflows has been accounted for at follow ppb levels [12, 14, 24–27]. Even though, the detected concentration of NORO is very low and normally ranges from ng L⁻¹ to μ g L⁻¹ in water bodies and μ g kg⁻¹ to mg kg⁻¹ in soils and sediments, still these fluoroquinolone family are categorized as "pseudopersistant" contaminants because of their continuous and regular discharge into the water bodies [28, 29]. González-Pleiter et al. [30] concentrated on those unique united toxicities from claiming norfloxacin, amoxicillin, erythromycin, levofloxacin, furthermore anti-microbial prescription toward two maritime organisms, i.e. Cyanobacterium Anabaena CPB 4337. Similarly as a goal existing being and the green alga *Pseudokirchneriella subcapitata* as a non-target existing continuously. They assigned norfloxacin on a chance to be a greater amount dangerous on cyanobacterium over green alga. Furthermore, norfloxacin alone and additionally its mixture for different antibiotics might stance genuine idle danger to oceanic environment.

The presence of NORO in the fresh water bodies indicate that traditional wastewater or water treatment techniques are not efficient to remove NORO from aquatic environment due to its aromatic nature and its occurrence cause thoughtful health associated problems by using contaminated drinking water [31–33]. Therefore, it becomes an issue of interest to remove NORO from the aquatic environment.

In a typical experiment for gamma radiolysis of NORO, the apparent bimolecular rate constant of ${}^{\bullet}OH$, $e_{aq}{}^{-}$ and ${}^{\bullet}H$ with NORO was assessed, using competition kinetics.

The following Eq. (14) was employed to measure the bimolecular rate constant of $^{\circ}OH$, e_{aq}^{-} and $^{\circ}H$, which are the main species produced during gamma radiolysis of aqueous media [21].

$$k_{\bullet \text{OH/NORO}} = \frac{\ln \left([\text{NORO}]_0 / [\text{NORO}]_D \right)}{\ln \left([2 - \text{CP}]_0 / [2 - \text{CP}]_D \right)} k_{\bullet \text{OH/2-CP}}$$
(16)

2-Chlorophenol (2-CP) was selected as reference compound which have recognized rate constants with [•]OH, e_{aq}^{-} and [•]H ($k_{\bullet OH/2-CP} = 1.2 \times 10^{10} \text{ M}^{-1} \text{ s}^{-1}$, $k_{e_{aq}^{-}/2-CP} = 1.3 \times 10^{9} \text{ M}^{-1} \text{ s}^{-1}$, $k_{\bullet H/2-CP} = 1.5 \times 10^{9} \text{ M}^{-1} \text{ s}^{-1}$) [20]. To permit only [•]OH to react with NORO, and scavenge e_{aq}^{-} and [•]H, O_2 saturated sample (O_2 changes e_{aq}^{-} and [•]H to superoxide radical anions, which are less responsive opposite to [•]OH) [20] was applied for computing the bimolecular rate of [•]OH with NORO. In the same way, the bimolecular rate constant of e_{aq}^{-} with NORO (k_{eaq} -/NORO) was calculated by N₂-puging the sample solution added with 0.1 M *iso*-propanol (*iso*-propanol is used to scavenges both [•]OH and [•]H) [20]. Similarly, the computation of bimolecular constant of [•]H with NORO ($k_{\bullet H/NORO}$) was made by N₂ saturating the solution of 0.1 M *tert*-butanol (*tert*-butanol is used to scavenge [•]OH) [20] at pH 2.2. Low pH was maintained to get high yield of [•]H through reaction of e_{aq}^{-} with ⁺H [34].

A linear plot with slope equal to $k_{\bullet OH/NORO}/k_{\bullet OH/2-CP}$ was observed by plotting ln([NORO]₀/ [NORO]_D) vs ln ([2-CP]₀/[2-CP]_D) at several absorbed ionizing doses. The same calculation was implemented for measurement of bimolecular rate constant of e_{aq}^{-} and $^{\bullet}H$ with NORO, respectively. Applying the obtained slope values, the second order rate constants of $^{\bullet}OH$, e_{aq}^{-} and $^{\bullet}H$ with NORO were computed to be $(8.81\pm0.03) \times 10^9 \text{ M}^{-1} \text{ s}^{-1}$, $(9.54\pm0.16) \times 10^8 \text{ M}^{-1} \text{ s}^{-1}$ and $(1.10\pm0.20) \times 10^9 \text{ M}^{-1} \text{ s}^{-1}$, respectively [5], which also indicates that $k_{e_{aq}^{-}/NORO}$ is lesser to $k_{\bullet H/NORO}$, or in other words the reactivity of e_{aq}^{-} to NORO is less than the reactivity of $^{\bullet}H$ with NORO. Thus, in the removal of NORO by ionizing irradiation $^{\bullet}H$ is of immense importance. The bimolecular rate constant of $^{\bullet}OH$ with NORO in the current report is analogous with the study of Santoke et al. [35], in which they calculated the bimolecular rate constants of $^{\bullet}OH$ with six common fluoroquinolones (orbifloxacin, flumequine, marbofloxacin, danofloxacin, enrofloxacin and model compound, 6-fluoro-4-oxo-1,4-dihydro-3-quinolone carboxylic acids) and was found to be in the range of $6.4 - 9.03 \times 10^9 \text{ M}^{-1} \text{ s}^{-1}$.

2.3. Measurement of bimolecular rate constant of 'OH with bezafibrate

Bezafibrate (BZF) is also the most commonly detected pollutant among various pharmaceuticals excreted into the sewage system and is categorized as persistent organic pollutants [36]. In drinking water its concentration has been noticed at the levels of 27 ng L⁻¹ [37] in rivers at the concentrations level of $0.1-0.15 \ \mu g \ L^{-1}$ [37], in small streams in the range of $0.5-1.9 \ \mu g \ L^{-1}$ [37], in surface waters in the range of $3.1 \ \mu g \ L^{-1}$ [38], and up to $4.6 \ \mu g \ L^{-1}$ level in sewage treatment plant effluents. Owing to its high use and persistence nature, the elimination of BZF from aqueous media has emerged as a hot research topic. The qualitative and quantitative analysis of its degradation products besides its degradation kinetics is also of great concern. Keeping in view all these problems, the degradation of BZF was investigated by photo catalysis using hydrothermally synthesized TiO₂/Ti films with exposed {001} facets. Besides photo catalysis, there are other many advanced treatment options for efficient removal of BZF from aqueous media, such as nanofiltration techniques, ultraviolet (UV) radiation and advanced oxidation processes (AOPs) [39] and these have been thoroughly studied. In AOPs (the most reliable and efficient technique), as compared to other treatment techniques the pollutant of interest is converted in to more stable, harmless inorganic species such as carbon dioxide, water and mineral salts. AOPs are categorized as ozonation (O₃), H₂O₂, O₃/H₂O₂/photocatalysis, and O₃/H₂O₂/UV photocatalysis [39, 40]. TiO₂ photocatalysis is considered as more auspicious and efficient technique among semiconductor photocatalysis [41, 42]. TiO₂ photo active material has shown a great potential in many applications, including water splitting to generate O₂ and H₂ [43, 44] water and wastewater treatment [45, 46], gas phase treatment [47, 48], as well as in solar cells [49]. So, in this case the degradation of BZF was performed by VUV photo active material with exposed {001} faceted TiO₂/Ti material.

For measurement of absolute bimolecular rate constant of \bullet OH with BZF, *para*-chlorobenzoic acid (*p*-CBA) was used as probe molecule was calculated using *para*-chlorobenzoic acid (*p*-CBA) as probe molecule and by employing competition kinetics technique established by Pereira et al. [50] and given in Eq. (15).

$$k_{\bullet OH(s)} = \frac{k_{s(UV/H_2O_2)} - k_{s(UV)}}{k_{ref(UV/H_2O_2)} - k_{ref}} \times k_{\bullet OH(ref)}$$
(17)

Where, $k_{\bullet OH}$, $k_{(UV/H_2O_2)}$ and $k_{(UV)}$ represent the second order rate constant of hydroxyl radical, UV/H₂O₂, fluence based rate constant of UV/hydrogen peroxide process and UV, fluence based rate constant of direct photolysis, respectively. The notations "s" and "ref" represents the substrate and reference compounds, which in our case is BZF and *p*-CBA, respectively. For the determination of [•]OH rate constant with BZF, two sets of experiments were performed. In one set of experiments, the solution containing 27.63 µM of BZF, 27.63 µM of *p*-CBA and 1 mM



Figure 2. Determination of bimolecular rate constant of BZF with $^{\circ}$ OH; inset shows the degradation kinetics of BZF alone, p-CBA alone, BZF+ H₂O₂, pCBA + H₂O₂ exposed to UV-irradiation.

of H_2O_2 was exposed to UV irradiation, while another set of experiments was free of H_2O_2 to calculate k_{UV} . The concentration of H_2O_2 was kept higher for ensuring production of efficient [•]OH with UV-photocatalysis.

Figure 2 shows degradation curves for BZF and *p*-CBA, both BZF and *p*-CBA were found to follow pseudo-first order degradation kinetics. The second order rate constant of [•]OH with p-CBA is $5.0 \times 10^9 \text{ M}^{-1} \text{ s}^{-1}$ [20]. By substituting pseudo-first order degradation constants ($k_{\text{UV/H}_2O_2}$ and k_{UV}) values in Eq. (15), the bimolecular rate constant of [•]OH with BZF was calculated and found out to be $5.66 \times 10^9 \text{ M}^{-1} \text{ s}^{-1}$.

3. Conclusions

The overall conclusion of this chapter is that, in radiation chemistry to have a good knowledge about the mechanism of a reaction mechanism it is necessary that one must have sufficient understanding about the free radical kinetics. In addition, competition kinetics model can be successful applied for the determination of unknown rate constants of reactive species with solute molecule. The competition kinetics can not only be applied for $^{\circ}OH$ rate constants with the solute but also for measurement of e_{aq}^{-} and $^{\circ}H$ with the target species. The competition kinetics method is validated by taking ciprofloxacin, norfloxacin and bezafibrate as example compounds. However, it should be make sure that competing reactions do not disobey the kinetics rules.

Author details

Murtaza Sayed¹*, Luqman Ali Shah¹, Javed Ali Khan¹, Noor S. Shah^{1,2}, Rozina Khattak³ and Hasan M. Khan¹

*Address all correspondence to: murtazasayed_407@yahoo.com

1 Radiation and Environmental Chemistry Laboratory, National Centre of Excellence in Physical Chemistry, University of Peshawar, Pakistan

2 Department of Environmental Sciences, COMSATS Institute of Information Technology, Vehari, Pakistan

3 Department of Chemistry, Shaheed Benazir Bhutto Women University, Peshawar, Pakistan

References

- [1] Spinks JWT, Woods RJ. An Introduction to Radiation Chemistry. John-Wiley and Sons, Inc., New York; 1990.
- [2] Fricke H, Hart EJ. Chemical dosimetry. In: Radiation Dosimetry 2. Academic Press, New York; 1966. pp. 167-239

- [3] Buxton GV, Stuart CR. Re-evaluation of the thiocyanate dosimeter for pulse radiolysis. Journal of the Chemical Society, Faraday Transactions 1995;**91**:279-281
- [4] Sayed M, Fu P, Shah LA, Khan HM, Nisar J, Ismail M, Zhang P. VUV-photocatalytic degradation of bezafibrate by hydrothermally synthesized enhanced {001} facets TiO2/Ti film. The Journal of Physical Chemistry A 2015;120:118-127
- [5] Sayed M, Khan JA, Shah LA, Shah NS, Khan HM, Rehman F, Khan AR, Khan AM. Degradation of quinolone antibiotic, norfloxacin, in aqueous solution using gamma-ray irradiation. Environmental Science and Pollution Research 2016;23:13155-13168
- [6] Sayed M, Shah LA, Khan JA, Shah NS, Khan HM, Khan RA, Khan AR, Khan AM. Hydroxyl radical based degradation of ciprofloxacin in aqueous solution. Journal of the Chilean Chemical Society 2016;61:2949-2953
- [7] Rehman F, Murtaza S, Ali Khan J, Khan HM. Removal of crystal violet dye from aqueous solution by gamma irradiation. Journal of the Chilean Chemical Society 2017;62:3359-3364
- [8] Sayed M, Ismail M, Khan S, Tabassum S, Khan HM. Degradation of ciprofloxacin in water by advanced oxidation process: Kinetics study, influencing parameters and degradation pathways. Environmental Technology 2016;37:590-602
- [9] Nisar J, Sayed M, Khan FU, Khan HM, Iqbal M, Khan RA, Anas M. Gamma-irradiation induced degradation of diclofenac in aqueous solution: Kinetics, role of reactive species and influence of natural water parameters. Journal of Environmental Chemical Engineering 2016;4:2573-2584
- [10] Zhang H, Huang C-H. Oxidative transformation of fluoroquinolone antibacterial agents and structurally related amines by manganese oxide. Environmental Science & Technology 2005;39:4474-4483
- [11] Ikehata K, Gamal El-Din M, Snyder SA. Ozonation and advanced oxidation treatment of emerging organic pollutants in water and wastewater. Ozone: Science & Engineering 2008;30:21-26
- [12] Watkinson AJ, Murby EJ, Costanzo SD. Removal of antibiotics in conventional and advanced wastewater treatment: Implications for environmental discharge and wastewater recycling. Water Research 2007;41:4164-4176
- [13] Xu W, Zhang G, Li X, Zou S, Li P, Hu Z, Li J. Occurrence and elimination of antibiotics at four sewage treatment plants in the Pearl River Delta (PRD), South China. Water Research 2007;41:4526-4534
- [14] Kümmerer K, Al-Ahmad A, Mersch-Sundermann V. Biodegradability of some antibiotics, elimination of the genotoxicity and affection of wastewater bacteria in a simple test. Chemosphere 2000;40:701-710
- [15] Mascarelli AL. New mode of action found for pharmaceuticals in the environment. Environmental Science & Technology 2010;44:1159-1160

- [16] Kümmerer K. Antibiotics in the aquatic environment—a review—part I. Chemosphere 2009;75:417-434
- [17] Christensen AM, Ingerslev F, Baun A. Ecotoxicity of mixtures of antibiotics used in aquacultures. Environmental Toxicology and Chemistry 2006;25:2208-2215
- [18] Carlsson G, Orn S, Larsson D. Effluent from bulk drug production is toxic to aquatic vertebrates. Environmental Toxicology and Chemistry 2009;28:2656-2662
- [19] Andreozzi R, Campanella L, Fraysse B, Garric J, Gonnella A, Giudice RL, Marotta R, Pinto G, Pollio A. Effects of advanced oxidation processes (AOPs) on the toxicity of a mixture of pharmaceuticals. Water Science and Technology 2004;50:23-28
- [20] Buxton G, Greenstock C, Helman W, Ross A. Critical review of rate constants for reactions of hydrated electrons, hydrogen-atoms and hydroxyl radicals OH/O⁻ in aqueous solution. Journal of Physical and Chemical Reference Data 1988;17:513-886
- [21] Kimura A, Osawa M, Taguchi M. Decomposition of persistent pharmaceuticals in wastewater by ionizing radiation. Radiation Physics and Chemistry 2012;81:1508-1512
- [22] Dodd MC, Buffle M-O, Von Gunten U. Oxidation of antibacterial molecules by aqueous ozone: Moiety-specific reaction kinetics and application to ozone-based wastewater treatment. Environmental Science & Technology 2006;40:1969-1977
- [23] Prieto A, Moder M, Rodil R, Adrian L, Marco-Urrea E. Degradation of the antibiotics norfloxacin and ciprofloxacin by a white-rot fungus and identification of degradation products. Bioresource Technology 2011;102:10987-10995
- [24] Kolpin DW, Furlong ET, Meyer MT, Thurman EM, Zaugg SD, Barber LB, Buxton HT. Pharmaceuticals, hormones, and other organic wastewater contaminants in US streams, 1999-2000: A national reconnaissance. Environmental Science & Technology 2002;36:1202-1211
- [25] Leung HW, Minh TB, Murphy MB, Lam JC, So MK, Martin M, Lam PK, Richardson BJ. Distribution, fate and risk assessment of antibiotics in sewage treatment plants in Hong Kong, South China. Environment International 2012;42:1-9
- [26] Gracia-Lor E, Sancho JV, Serrano R, Hernandez F. Occurrence and removal of pharmaceuticals in wastewater treatment plants at the Spanish Mediterranean area of Valencia. Chemosphere 2012;87:453-462
- [27] Oberlé K, Capdeville M-J, Berthe T, Budzinski Hln, Petit F. Evidence for a complex relationship between antibiotics and antibiotic-resistant *Escherichia coli*: From medical center patients to a receiving environment. Environmental Science & Technology. 2012; 46:1859-1868
- [28] Lindberg RH, Olofsson U, Rendahl P, Johansson MI, Tysklind M, Andersson BAV. Behavior of fluoroquinolones and trimethoprim during mechanical, chemical, and active sludge treatment of sewage water and digestion of sludge. Environmental Science & Technology 2006;40:1042-1048

- [29] Hernando MD, Mezcua M, Fernandez-Alba AR, Barcelo D. Environmental risk assessment of pharmaceutical residues in wastewater effluents, surface waters and sediments. Talanta 2006;**69**:334-342
- [30] González-Pleiter M, Gonzalo S, Rodea-Palomares I, Leganés F, Rosal R, Boltes K, Marco E, Fernández-Piñas F. Toxicity of five antibiotics and their mixtures towards photosynthetic aquatic organisms: Implications for environmental risk assessment. Water Research 2013;47:2050-2064
- [31] Liu WF, Zhang J, Zhang CL, Ren L. Sorption of norfloxacin by lotus stalk-based activated carbon and iron-doped activated alumina: Mechanisms, isotherms and kinetics. Chemical Engineering Journal 2011;**171**:431-438
- [32] Rosal R, Rodriguez A, Perdigon-Melon JA, Petre A, Garcia-Calvo E, Gomez MJ, Aguera A, Fernandez-Alba AR. Occurrence of emerging pollutants in urban wastewater and their removal through biological treatment followed by ozonation. Water Research 2010;44: 578-588
- [33] Xu WH, Zhang G, Zou SC, Li XD, Liu YC. Determination of selected antibiotics in the Victoria Harbour and the Pearl River, South China using high-performance liquid chromatography-electrospray ionization tandem mass spectrometry. Environmental Pollution 2007;145:672-679
- [34] Sánchez-Polo M, López-Peñalver J, Prados-Joya G, Ferro-García MA, Rivera-Utrilla J. Gamma irradiation of pharmaceutical compounds, nitroimidazoles, as a new alternative for water treatment. Water Research 2009;43:4028-4036
- [35] Santoke H, Song W, Cooper WJ, Greaves J, Miller GE. Free-radical-induced oxidative and reductive degradation of fluoroquinolone pharmaceuticals: Kinetic studies and degradation mechanism. The Journal of Physical Chemistry A 2009;113:7846-7851
- [36] Sui Q, Gebhardt W, Schröder HF, Zhao W, Lu S, Yu G. Identification of new oxidation products of bezafibrate for better understanding of its toxicity evolution and oxidation mechanisms during ozonation. Environmental Science & Technology 2017;51:2262-2270
- [37] Ternes T. Pharmaceuticals and Metabolites as Contaminants of the Aquatic Environment. ACS Publications; 2001.
- [38] Weston A, Caminada D, Galicia H, Fent K. Effects of lipid-lowering pharmaceuticals bezafibrate and clofibric acid on lipid metabolism in fathead minnow (Pimephales promelas). Environmental Toxicology and Chemistry. American Chemical Society. 2009;28: 2648-2655
- [39] Klavarioti M, Mantzavinos D, Kassinos D. Removal of residual pharmaceuticals from aqueous systems by advanced oxidation processes. Environment International 2009;35: 402-417
- [40] Parsons S. Advanced Oxidation Processes for Water and Wastewater Treatment. IWA publishing; 2004

- [41] Madhavan J, Grieser F, Ashokkumar M. Combined advanced oxidation processes for the synergistic degradation of ibuprofen in aqueous environments. Journal of Hazardous Materials. American Chemical Society. 2010;178:202-208
- [42] Grabowska E, Diak M, Marchelek M., Zaleska A. Decahedral TiO₂ with exposed facets: Synthesis, properties, photoactivity and applications. Applied Catalysis B: Environmental 2014;156:213-235
- [43] Kim HS, Kim D, Kwak BS, Han GB, Um M-H, Kang M. Synthesis of magnetically separable core@ shell structured NiFe2O4@TiO2 nanomaterial and its use for photocatalytic hydrogen production by methanol/water splitting. Chemical Engineering Journal 2014;243:272-279
- [44] Wang C, Hu Q-Q, Huang J-Q, Deng Z-H, Shi H-L, Wu L, Liu Z-G, Cao Y-G. Effective water splitting using N-doped TiO2 films: Role of preferred orientation on hydrogen production. International Journal of Hydrogen Energy 2014;39:1967-1971
- [45] Choi J, Lee H, Choi Y, Kim S, Lee S, Lee S, Choi W, Lee J. Heterogeneous photocatalytic treatment of pharmaceutical micropollutants: Effects of wastewater effluent matrix and catalyst modifications. Applied Catalysis B: Environmental 2014;147:8-16
- [46] Lee HU, Lee G, Park JC, Lee Y-C, Lee SM, Son B, Park SY, Kim C, Lee S, Lee SC. Efficient visible-light responsive TiO₂ nanoparticles incorporated magnetic carbon photocatalysts. Chemical Engineering Journal 2014;240:91-98
- [47] Mrowetz M, Balcerski W, Colussi A, Hoffmann MR. Oxidative power of nitrogen-doped TiO₂ photocatalysts under visible illumination. The Journal of Physical Chemistry B 2004; 108:17269-17273
- [48] Nischk M, Mazierski P, Gazda M, Zaleska A. Ordered TiO₂ nanotubes: The effect of preparation parameters on the photocatalytic activity in air purification process. Applied Catalysis B: Environmental 2014;144:674-685
- [49] Dwivedi C, Dutta V, Chandiran AK, Nazeeruddin MK, Grätzel M. Anatase TiO₂ hollow microspheres fabricated by continuous spray pyrolysis as a scattering layer in dyesensitised solar cells. Energy Procedia 2013;33:223-227
- [50] Pereira VJ, Weinberg HS, Linden KG, Singer PC. UV degradation kinetics and modeling of pharmaceutical compounds in laboratory grade and surface water via direct and indirect photolysis at 254 nm. Environmental Science & Technology 2007;41:1682-1688