The Metal Catalyst Influences the Kinetics and Mechanisms of MS2 Inactivation in Fenton-like Systems

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Abstract: Human enteric viruses are frequent microbial contaminants of surface water and groundwater. Waterborne viruses can be effectively inactivated by oxidants, such as those generated in Fenton-like systems. However, the mechanisms by which this inactivation occurs are not understood. Here we investigated how two Fenton-like systems, Cu/H₂O₂ and Fe/H₂O₂/light, affect the infectivity and structural integrity of MS2 coliphage, a frequently used surrogate for human enteric viruses. The extent of MS2 genome and capsid protein degradation was evaluated by quantitative PCR and protein mass spectrometry, and was related to the observed level of inactivation. Even though inactivation in both systems occurred via the same oxidant, hydroxyl radical, the contributions of genome and capsid protein degradation to inactivation differed. Inactivation in the Cu/H₂O₂ system was rapid and involved both genome and protein damage. In contrast, inactivation in Fe/H₂O₂/light proceeded at a slower rate and encompassed solely genome damage. Our findings demonstrate that not only the oxidant, but also its source, the metal catalyst, determines the inactivation kinetics and mechanism in Fenton-like systems. This work provides the first evidence of the impact of the metal catalyst on virus inactivation in Fenton-like systems.

Keywords: Disinfection · Fenton · Inactivation mechanisms · Protein mass spectrometry · Waterborne virus

Introduction

The disinfection of drinking water, and in some countries of wastewater, is an important step in preventing the transmission of waterborne diseases such as poliomyelitis or hepatitis. Water disinfection is typically performed by treatment with an oxidizing chemical, such as free chlorine (HOCl) or ozone, or by physical methods such as the irradiation by UVC light. These methods, while effective, each have disadvantages, such as a high energy consumption, or the formation of hazardous disinfection by-products. Therefore, much research has been conducted in recent years to devise alternative disinfection strategies that rely on more energy-efficient and environmentally sustainable disinfection methods.

Over the past decade, Fenton-like processes have been explored as alternative methods for water disinfection. The most attention has been dedicated to the photo-Fenton system, in which ferric iron is photo-reduced to ferrous iron, which subsequently reacts with hydrogen peroxide (H₂O₂) to generate oxidants. Fenton-like systems using other transition metals or iron (hydr) oxide minerals as catalyst to generate oxidants have also been explored. Fenton-like systems using other transition metals or iron (hydr) oxide minerals as catalyst to generate oxidants have also been explored. Fenton-like systems can successfully inactivate various pathogens, including viruses, through the Fenton process, have been implicated as inactivating agents in these processes, though their mechanism of action is not understood. Specifically, it is unclear which virus constituents (genome and proteins) are most efficiently targeted.

HO• radicals are known to be reactive toward the different viral building blocks. HO• radicals can oxidize all amino acids, though tryptophan, tyrosine, histidine and cysteine are the most suscep-
HO• is also reactive toward all nucleotides that make up the viral genome. Previous studies conducted on viruses have shown that HO• generated by TiO$_2$ photocatalytic disinfection induced genome damage in RNA and DNA coliphages, though protein degradation was not considered. To our knowledge, damage to virus components in Fenton-like systems has not been investigated.

Our previous work has shown that Fenton-like systems (Fe/\(\text{H}_2\text{O}_2/\text{light}\), and Cu/\(\text{H}_2\text{O}_2\) systems) efficiently inactivate bacteriophage MS2\(^9\) (Fig. 1). The aim of this work was to investigate chemical degradation of the components of MS2 during treatment in these Fenton-like systems, and to link the observed degradation to inactivation. To this end, we combined Fenton chemistry with protein mass spectrometry and traditional tools of environmental virology (virus measuring to culture infectivity; quantitative polymerase chain reaction (qPCR) to measure genome integrity). MS2 is a frequently used model organism for human enteric viruses. It is a non-enveloped coliphage that consists of an RNA genome (single-stranded RNA, 3569 nucleotides) surrounded by 180 copies of a capsid protein (CP; 13.7 kDa, 129 amino acids). In addition, the virion contains a single copy of an assembly protein (A protein, 43.9 kDa, 339 amino acids). Inactivation and damage to the MS2 genome and capsid proteins was studied in both Fenton-like systems, and the patterns of damage were compared to elucidate the role of the oxidant and the metal catalysts in inactivation.

Materials and Methods

MS2 phages were inactivated in two Fenton-like systems: one containing Cu/\(\text{H}_2\text{O}_2\) and one composed of Fe/\(\text{H}_2\text{O}_2\) irradiated by simulated sunlight. Samples were periodically taken and analyzed for remaining MS2 infectivity, as well as genome and protein integrity. Damage to different segments of the viral genome was quantified by qPCR. Individual loss of the capsid protein fragments were determined by protein mass spectrometry (Matrix-assisted laser desorption ionization time-of-flight mass spectrometry; MALDI-TOF-MS).

Reagents and Organisms

Reagents

Copper sulfate (CuSO$_4$·5H$_2$O), hydrogen peroxide (H$_2$O$_2$, 30%), EDTA, catalase (2000–5000 units mg$^{-1}$ protein), and \(\alpha\)-cyano-4-hydroxycinnamic acid were purchased from Sigma-Aldrich (St. Louis, Missouri). Ferric chloride (FeCl$_3$·6H$_2$O), sodium bicarbonate (NaHCO$_3$, 99+%), NaCl (99+%), CaCl$_2$ (99+%), NaH$_2$PO$_4$ (99+%), trifluoroacetic acid (TFA), acetonitrile (ACN, HPLC grade) were obtained from Acros Organics (Geel, Belgium). Trypsin and chymotrypsin were purchased from Worthington Biochemical Corporation (Lakewood, NJ).

Organisms

MS2 coliphage (DSMZ 13767) and its bacterial host E. coli (DSMZ 5695) were purchased from the German collection of microorganisms and cell cultures (DSMZ, Braunschweig, Germany). The propagation and purification of MS2 was performed as previously described.\(^1\) The infective MS2 phage concentrations were enumerated by the double-layer agar method and measured in plaque-forming units per ml (PFU ml$^{-1}$). The MS2 stock solution had a concentration of 10$^{14}$ PFU ml$^{-1}$ and was stored in dilution buffer (DB; 5 mmol l$^{-1}$ NaH$_2$PO$_4$, 10 mmol l$^{-1}$ NaCl, pH 7.4) at 4 °C. A 15N-labeled stock of MS2 was used as the internal standard for quantitative protein mass spectrometric analyses. 15N-labeled MS2 was propagated and purified as described previously,\(^1\) using E. coli (ATCC 15597; LGC Standards, Molsheim, France) as the host. The 15N-labeled MS2 stock solution had a concentration of 10$^{10}$ PFU ml$^{-1}$ and was stored in DB at 4 °C.

Inactivation Experiments

Experiments were carried out in sacrificial glass reactors (50 ml), with an initial infective MS2 concentration of 10$^{10}$ PFU ml$^{-1}$ (detection limit for the protein mass measurement) in 20 ml of carbonate-buffered saline (CBS; 0.1 mmol l$^{-1}$ NaHCO$_3$, 15 mmol l$^{-1}$ NaCl) at pH 6.9. Cu(II) or Fe(III) were added to the reactors from stock solutions (1 mmol l$^{-1}$ CuSO$_4$, or FeCl$_3$) to obtain a final concentration of 10 µmol l$^{-1}$. H$_2$O$_2$ was added from a 25 mmol l$^{-1}$ stock solution to a final concentration of 50 µmol l$^{-1}$. EDTA (2 µmol l$^{-1}$) was added to complex any trace metals, which have shown to catalyze Fenton-like processes.\(^9\) This EDTA concentration did not cause virus inactivation. The solutions were stirred at 400 rpm on a multi-position stir plate. To obtain different levels of inactivation, 500 µL of catalase (500 units ml$^{-1}$) were added to sacrificial reactors at different times to scavenge the remaining H$_2$O$_2$ and halt the Fenton-like reactions. H$_2$O$_2$-free control samples were also evaluated. After catalase addition, sample aliquots were withdrawn from reactors for virus quantification by culturing. The remainder of the solution was used for genome and capsid protein analysis as described below. All experiments were performed in triplicate.

For photo-Fenton experiments, Fe/H$_2$O$_2$ solutions were irradiated using a solar simulator (ABET Technologies, Sun 2000) with a 1000 W Xe lamp, an air mass 1.5 filter and UVB cut-off filter. The UVB cut-off filter was used to avoid confounding effects arising from direct genome damage induced by UVB light. The resulting light spectrum has been previously reported.\(^8\) The total irradiance in the visible spectrum was 320 W m$^{-2}$. The sample temperature was controlled by placing the reactors in a cooled water bath at 20 ± 2 °C. Dark control experiments were conducted in reactors covered with aluminium foil.

Virus Aggregation

Virus aggregation was measured by dynamic light scattering (Malvern Instruments, Nano ZS) as previously described.\(^1\) Aggregation measurements were conducted for the same solution conditions as used in the inactivation experiments.

Measurement of Genome Damage by qPCR

Quantitative polymerase chain reaction (qPCR) was used to quantify and localize the genome damage of MS2 upon inactivation. 200 µL of samples were withdrawn from the reactors for the immediate extraction of virus genomes as previously described.\(^1\) Damage to six different genome segments of approximately 300 nucleotides was quantified using a RotorGene 3000 quantitative PCR platform (Corbett Life Science, Sidney, Australia). Jointly, the six segments covered approximately 50% of the entire ge-
nomo. The exact location of the segments within the genome, as well as the primers used, are described in Table 1. Each segment was first reverse transcribed and then amplified using the one step SYBR PrimeScript RT-PCR kit according the manufacturer’s procedure (Takara Bio Inc., Shiga, Japan). The qPCR protocols are described in detail elsewhere.[17] SYBR green was used as the fluorescent probe for all segments.

Damage to the entire genome was extrapolated from the measured damage to the six segments as described previously.[20] Briefly, the proportion of intact whole genomes (N/N_0) in the sample at any level of inactivation can be calculated as:

\[
\frac{N}{N_0} = \prod_{i=1}^{6} \left( \frac{n_i}{n_i,0} \right) \tag{1}
\]

where \( \prod_{i=1}^{6} \left( \frac{n_i}{n_i,0} \right) \) corresponds to the product of the intact proportions of each of the six individual genome segments (\( n_i \)) considered. The total genome length corresponds to the MS2 genome size (3569 nucleotides) and the length of six genome segments is the sum of all genome segment sizes shown in Table 1.

The extent of damage to each genome segment was assessed before and after inactivation. The inactivation levels studied were between 2 log (99%) and 7 log (99.99999%) for Cu/H_2O_2 system and between 1 log (90%) and 7 log for the Fe/H_2O_2/ light system.

Measurement of Capsid Protein Damage by MALDI-TOF-MS

Protein analysis was conducted on the same samples as genome analysis. Protein mass spectrometry was used to identify the regions of the MS2 capsid protein that were most susceptible to damage. After inactivation, 20 \( \mu \)l of 15N-labeled MS2 was added to each reactor as an internal standard for the quantification of protein damage. The ratio of initial native MS2 to labeled MS2 was 1:1. The capsid protein was denatured, alkylated and digested by two protease enzymes (trypsin or chymotrypsin) into small peptides as previously described.[18] The MS2 capsid protein sequence and the peptides generated by the digestion are shown in Tables 2 and 3, respectively. Jointly, the peptides resulting from digestion by the two protease enzymes yielded a 98% coverage of the capsid protein.

Calibration curves for each peptide were obtained by analyzing six different ratios of native to labeled MS2 (ratio range 0.2–1.2). The standards were denatured, alkylated and digested in the same way as the samples. The intact fraction of each peptide upon inactivation was determined by comparing the peak ratio of the native to labeled peptide in a sample to the ratios measured in the calibration curve.

Peptide analyses were carried out in positive ion linear mode MALDI-TOF-MS (ABI 4800; Applied Biosystems, Rotkreuz, Switzerland). The MALDI matrix used was 7 mg ml\(^{-1}\) of \( \alpha \)-cyano-4-hydroxy-cinnamic acid solution diluted in 50% ACN and 50% TFA (0.1%) as the solvent. The ratio between matrix and sample was 10 to 1. These mixtures of each sample were placed on a MALDI plate by quadruped and were air-dried. MALDI spectra were analyzed with Applied Biosystems Data Explorer Software.

Results

Inactivation Kinetics

MS2 was readily inactivated in both the Cu/H_2O_2 and the Fe/ H_2O_2/ light systems (Fig. 2). Virus inactivation followed (pseudo-) first-order kinetics for both disinfection systems. In the H_2O_2-free control samples, a decrease in infectious MS2 of approximately 90% (1 log loss) in 15 min and 99% (2 log loss) in 30 min was observed for Cu and Fe systems, respectively. Metal-free H_2O_2 samples have previously been shown to not cause inactivation at the concentrations studied herein.[9]

Table 1. Primer sets, segment position and segment sizes used for the quantification of MS2 genome damage. nt = nucleotide.

| Segment number\(^a\) | Primer direction | Primer sequence (5’ to 3’) | Genome position | Segment size (nt) |
|----------------------|------------------|----------------------------|-----------------|------------------|
| 2                    | Forward          | AAGGTGCCCTACAAGCGAAGT      | 344 to 678      | 335              |
|                      | Reverse          | TTCGTTTAGGGCAAGGTGC        |                 |                  |
| 3                    | Forward          | CCGCTACCTTGCCCTAAAC        | 657 to 959      | 303              |
|                      | Reverse          | GACGACAACCATGCGCAAC        |                 |                  |
| 6                    | Forward          | CCTAAAGTGGAACCCAGAC        | 1530 to 1818    | 289              |
|                      | Reverse          | AAAGATCGCGGAGGAAGATCA      |                 |                  |
| 7                    | Forward          | CGCGATCTTTTCTCTGAAT        | 1809 to 2125    | 317              |
|                      | Reverse          | GACGATCGGTAGCCAGAGG        |                 |                  |
| 10                   | Forward          | ATAGTCAAGCGACAAAACT        | 2724 to 3033    | 310              |
|                      | Reverse          | GGCCTGGATCTCGACATCA        |                 |                  |
| 12                   | Forward          | GAATACCCAGACGCATGAA        | 3285 to 3528    | 240              |
|                      | Reverse          | AATCCCCGGGTCCTCTCTTTA      |                 |                  |

\(^a\) Numbering corresponds to that in Pecson et al.[17]

Table 2. Amino acid sequence of the MS2 capsid protein (PDB file 2MS2)

| Amino acid sequence of the MS2 capsid protein (PDB file 2MS2) |
|-------------------------------------------------------------|
| ASNFTQFVLY\(^{20}\)DNNGTGTDVTV\(^{20}\)APSNFANGVA\(^{90}\)EWISSNSRSQ\(^{60}\)AYKVTCSRQ\(^{30}\)SSAQNRKYTI\(^{10}\)KVEVPKVATQ\(^{70}\)TVGGVELPVA\(^{90}\)AWRSYNMEL\(^{150}\)TIPFATNSD\(^{100}\)CELIKAMQG\(^{150}\)LLKDGNPIPS\(^{20}\)AIAANSGY |
Genome Damage

Two aspects of genome damage were assessed: the distribution of damage across the genome, and the extent of genome damage as a function of inactivation. Damage was assessed before treatment and at two levels of inactivation, as well in the relevant H$_2$O$_2$-free control samples.

In the control samples without H$_2$O$_2$, all genome segments remained intact in both systems investigated. However, a decrease in the qPCR signal indicative of genome damage was measured at each level of inactivation (Fig. 3). At the lower levels of inactivation (1 or 2 log), a similar, heterogeneous pattern of damage was observed in both systems. The most affected genome segments were segments 2 and 3, whereas little or no damage was observed in the other four segments. At higher levels of inactivation (7 log), the differences in degradation between the segments was less apparent. Furthermore, at 7 log inactivation, genome damage in the Fe/H$_2$O$_2$/light exceeded that in the Cu/H$_2$O$_2$.

To evaluate the extent of damage across the whole genome as a function of inactivation, the proportions of intact, whole genome (N/N$_0$) and remaining infectivity (C/C$_0$) were compared (Fig. 4). Hereby, C/C$_0$ was assessed by culturing, whereas N/N$_0$ was determined by extrapolating qPCR data, using Eqn. (1). For the Cu/H$_2$O$_2$ system, the extent of genome integrity loss was approximately equal to that of infectivity loss (Fig. 4a). In contrast, in the Fe/H$_2$O$_2$/light system, the extent of genome damage exceeded that of inactivation, in particular at high levels of inactivation (Fig. 4b).

Capsid Protein Damage

As for genome damage, the extent of capsid protein damage incurred at two levels of inactivation were evaluated for the Cu/H$_2$O$_2$ and Fe/H$_2$O$_2$/light systems, respectively. We assessed whether the inactivation by these treatments leads to protein degradation, and if so, which protein segments (peptides) are most susceptible.

The pattern of damage significantly differed between the two systems studied (Fig. 5): in the Cu system, the H$_2$O$_2$-free control sample showed a homogeneous loss in intact peptide concentration of roughly 20% (Fig. 5a). Upon addition of H$_2$O$_2$, an increasing loss in intact peptide concentration was observed for all peptides at 2 log and 7 log of inactivation. Notably, some peptides were affected to a greater extent, namely those found at m/z 1761 (position 8-25), 1754 (position 67-83) and 2671 (position 84-106). In the Fe-based system, a heterogeneous decrease in peptide signal was observed in the H$_2$O$_2$-free control (Fig. 5b). In contrast to the Cu system, most peptides did not degrade further upon addition of H$_2$O$_2$. Only selected peptides (e.g. m/z 184, position 1-7) exhibited a measurable decay after 7 log inactivation compared to the H$_2$O$_2$-free control.

Discussion

To assess the utility of Fenton-like systems for virus disinfection, it is important to understand the inactivation mechanisms involved. Here, virus inactivation was evaluated by means of investigating the inactivation kinetics, along with the accompanying
genome and protein degradation. The results demonstrate that, while both systems tested efficiently inactivated MS2, important differences exist in both kinetics and inactivation mechanisms.

**Influence of Metal Catalyst on MS2 Inactivation**

At the same metal and peroxide concentration, MS2 was inactivated more rapidly in the Cu/H₂O₂ system (Fig. 2). This is particularly interesting given the fact that the production rate of hydroxyl radicals, the main oxidant in both treatment systems, was greater in the Fe/H₂O₂/light system. This finding provides an indication that the metal catalyst involved in oxidant production plays a crucial role. We have previously demonstrated that under our experimental conditions, HO• is mainly produced from dissolved Cu(ii) species in the Cu/H₂O₂ system, and from colloidal particles in the Fe/H₂O₂/light system. We have furthermore shown that the metal catalyst, and hence the site of oxidant production, must be in close vicinity to the virion to effectively cause inactivation. The small size of the dissolved Cu(ii) likely allows a denser coverage of the virion with metal catalyst compared to the larger Fe colloids. This spatially distinct arrangement of the site of oxidant production in the two systems investigated (illustrated in Fig. 1) may influence the extent and pattern of genome and protein degradation induced by the oxidants produced, and may ultimately account for the more efficient inactivation in the Cu/H₂O₂ system. The effect of the metal catalyst on genome and protein degradation is discussed below.

**Contribution of Genome Damage to Inactivation**

By quantifying the degradation of different genome segments, as well as of the whole genome, as a function of inactivation, several aspects regarding the mechanism of virus inactivation upon treatment by Fenton-like processes could be addressed. Specifically, we could assess 1) if inactivation by these Fenton-like leads to genome damage; 2) if damage is homogeneously distributed across the entire genome or selectively degrades particular genome regions; and 3) if genome damage is sufficiently extensive to account for the entire observed inactivation.

Inactivation was accompanied by significant genome damage (Fig. 3). The most affected genome segment in both systems was the 5'-end of the genome (segment 2), indicating the 5'-end of the genome was more susceptible to attack by the oxidants than segments in the center and at the 3'-end. The distribution of genome damage is thus not a random process, and it is influenced by the genome structure but not the metal catalyst. While the distribution of damage was similar in the two systems tested, the extent of genome damage differed, with the Fe/H₂O₂/light system inducing more extensive genome degradation per unit inactivation than the Cu/H₂O₂ system (Fig. 4). The 1:1
the ability of iron (hydr)oxides to denature proteins.\[22\] It is thus
ity to the genome. A possible explanation for this finding lies in
the inactivation of MS2 in the Cu/H\textsubscript{2}O\textsubscript{2} system. Instead, protein
sume that genome damage alone was not sufficient to account for
in was quantified by qPCR, whereas it is not clear if each lesion
related between genome damage and inactivation observed in
the Cu/H\textsubscript{2}O\textsubscript{2} system is consistent with MS2 being inactivated by
single-hit lesions to the genome. However, genome damage here-
in was quantified by qPCR, whereas it is not clear if each lesion detected by qPCR also causes MS2 inactivation. In fact, several
studies have demonstrated that tandem and clustered lesions are
are prone to degradation by Fenton-like processes.\[29,30\] Segment
84-106 contains two sulfur-bearing amino acids, namely a me-
thionine (Met88) and a cysteine (Cys101), which may account
for its propensity to degrade in the presence of oxudants. Other
sulfur-containing residues (e.g. Cys46 in segment 44-49), which
were less solvent-accessible, were less susceptible to degradation.
Overall, it can be concluded that oxidants generated in the Cu/
H\textsubscript{2}O\textsubscript{2} system can attack the entire capsid protein, though protein
regions that are solvent-exposed and contain easily oxidizable
residues are the most affected. It is not known, however, if the
degradation of these regions directly contributes to loss of virus
infectivity.
In summary, our results demonstrate virus inactivation in
different Fenton-like systems can differ with respect to both in-
activation kinetics and mechanisms, and is closely linked to the
aqueous chemistry of the transition metal catalyst. Specifically,
the solubility and resulting spatial arrangement of the catalyst
around the virion, and hence the site oxidant production, likely
determine the disinfection outcome. In the Cu/H\textsubscript{2}O\textsubscript{2} system, HO\textsuperscript{+}
was produced by dissolved Cu ions, which can readily access the
virion. The resulting inactivation was rapid and involved the deg-
radation of both viral genome and proteins. In the Fe H\textsubscript{2}O\textsubscript{2}/light system, in contrast, HO\textsuperscript{+} was produced by iron colloids, which
are likely positioned at a greater distance to the virion surface and
are thus less effective at oxidizing the viral proteins. Instead,
the iron colloids may have denatured the protein and exposed the RNA. The resulting inactivation occurred at a slower rate, and involved solely genome damage. The dissolved or colloidal state of the catalyst should be considered when applying Fenton-like solutions. The dissolved or colloidal state promotes both genome and protein damage should be considered.

Acknowledgements
This research was funded by Swiss National Science Foundation (Projects No. 200021_118077 and 200020_131918). The authors thank Thérèse Sigstam and Loic Decrey for performing the MALDI-TOF-MS measurements.

Conflict of Interest
No conflict of interest declared.

Received: October 1, 2019

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