Review

Evidence-Based Framework to Manage Cyanobacteria and Cyanotoxins in Water and Sludge from Drinking Water Treatment Plants

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Abstract: Freshwater bodies and, consequently, drinking water treatment plants (DWTPs) sources are increasingly facing toxic cyanobacterial blooms. Even though conventional treatment processes including coagulation, flocculation, sedimentation, and filtration can control cyanobacteria and cell-bound cyanotoxins, these processes may encounter challenges such as inefficient removal of dissolved metabolites and cyanobacterial cell breakthrough. Furthermore, conventional treatment processes may lead to the accumulation of cyanobacteria cells and cyanotoxins in sludge. Pre-oxidation can enhance coagulation efficiency as it provides the first barrier against cyanobacteria and cyanotoxins and it decreases cell accumulation in DWTP sludge. This critical review aims to: (i) evaluate the state of the science of cyanobacteria and cyanotoxin management throughout DWTPs, as well as their associated sludge, and (ii) develop a decision framework to manage cyanobacteria and cyanotoxins in DWTPs and sludge. The review identified that lab-cultured-based pre-oxidation studies may not represent the real bloom pre-oxidation efficacy. Moreover, the application of a common exposure unit CT (residual concentration × contact time) provides a proper understanding of cyanobacteria pre-oxidation efficiency. Recently, reported challenges on cyanobacterial survival and growth in sludge alongside the cell lysis and cyanotoxin release raised health and technical concerns with regards to sludge storage and sludge supernatant recycling to the head of DWTPs. According to the review, oxidation has not been identified as a feasible option to handle cyanobacterial-laden sludge due to low cell and cyanotoxin removal efficacy. Based on the reviewed literature, a decision framework is proposed to manage cyanobacteria and cyanotoxins and their associated sludge in DWTPs.

Keywords: cyanobacteria; cyanotoxins; pre-oxidation; sludge; accumulation; management; water treatment plant

Key Contribution: Cyanobacteria and cyanotoxins management in DWTPs is a triangular activity that includes monitoring, treatment, and sludge handling and interrelation.

1. Introduction

A cyanobacterial bloom occurrence may result in metabolite (cyanotoxins and taste and odor agents) production and release, which is considered a widespread problem in drinking water resources around the world [1–10].

Conventional treatment processes, including coagulation, flocculation, sedimentation, and filtration, are widely applied to remove cyanobacterial cells and cell-bound cyanotoxins [11–14]. However, conventional treatment processes may not be able to remove
dissolved metabolites (e.g., cyanotoxins) efficiently [12,15–22]. Moreover, toxic cyanobacterial breakthrough has been reported in the effluent of conventional treatment processes and even after post-oxidation [18]. Therefore, additional treatment such as oxidation or powdered (PAC)/granular activated carbon (GAC) may be required to control dissolved metabolites [23–26].

Pre-oxidation enhances cyanobacteria cell removal during the coagulation/sedimentation process [27–34] and may decrease the cell breakthrough potential from the downflow processes. However, it is reported that pre-oxidation may cause cyanobacteria cell damage (decrease in cell viability) and cell-bound cyanotoxin release [35–37]. The level of cell lysis/damage and cyanotoxin degradation/release following pre-oxidation depends on the oxidation exposure (CT as residual concentration x contact time), and it is the driver to find the best pre-oxidation practice against cyanobacteria and cyanotoxins [38,39].

Furthermore, conventional treatment processes cause cell accumulation in drinking water treatment plants’ (DWTPs) sludge, even in DWTPs with low cyanobacterial cell numbers in the intake water [13,18,40–44]. Several studies have demonstrated that cyanobacterial cells could survive in the stored sludge and release cyanotoxins for up to 12 days [13,40,45–52]. Recent studies revealed a new challenge on the probability of extended survival time and even cyanobacterial growth during sludge storage [53–55]. Thus, recycling the supernatant of stored cyanobacteria-laden sludge to the head of the DWTPs can increase health-related concerns [55,56]. Such challenges highlight the importance of the treatment and management of cyanobacteria-laden sludge [57–59].

The objectives of this study are to: (1) critically review shreds of evidence of the pre-oxidation impact on the cultured-based and natural bloom studies, (2) perform a critical review of the fate of cyanobacteria and cyanotoxins in conventional treatment plants’ sludge and during sludge storage, and (3) develop an operational decision framework to determine the best practice to minimize risks associated with cyanobacteria and cyanotoxin presence in DWTPs.

This critical review provides insight into the fate of cyanobacteria and their associated metabolites throughout DWTPs and their sludge; furthermore, a practical decision framework to mitigate health and operational risks is developed.

2. Impact of Conventional Treatment on Cyanobacteria and Cyanotoxin Accumulation in Sludge

Different studies have reported that conventional treatment processes can remove 62–99% of the cyanobacterial cells in DWTPs [13,18,40–42,56]. It has been demonstrated that potential toxic cyanobacterial cells such as Microcystis, Dolichospermum, and Aphanocapsa can be removed using conventional processes [12,56,60,61].

The long-term monitoring of a high-risk DWTP (Lake Champlain—Quebec) during cyanobacterial bloom seasons from 2008 to 2011 showed an extreme accumulation of cyanobacteria cells (up to \(10^7\) cells/mL) and cyanotoxins (up to 60 \(\mu\)g/L microcystin-LR (MC-LR)) in the sludge of the clarifier [12,18]. Monitoring of the same DWTP in 2017 showed that cyanobacteria cell accumulation in the sludge holding tank was up to 31-fold higher than taxonomic cell counts in the intake water [56]. An investigation of four DWTPs in the Great Lakes (Ontario) with low cyanobacterial cell influx (<1000 cells/mL) revealed that cyanobacterial cells and cyanotoxins accumulated in the sludge by up to 100 and 12 times higher than the raw water, respectively [44]. Zamyadi, et al. [57] reported a 406% and 2600% cell count increase in the thickened and centrifuged sludge, respectively, in a DWTP equipped with dissolved air flotation (DAF). A similar high accumulation was also reported for the backwash of the direct filtration process [45].

Pre-oxidation may decrease the risk of cyanobacterial accumulation in the clarifier and sludge [43]. Two DWTPs (the same source for intake water with low cyanobacteria cells; maximum < 500 cells/mL) with chemically enhanced conventional treatment processes were studied, but only one of the DWTPs was equipped by pre-ozonation [43]. Pre-ozonation (initial concentration: 0.3–0.8 mg/L, contact time: 6.3 min) decreased cell
accumulation in the surface of the clarifier and filters by up to 1450 times as compared to the DWTP without pre-ozonation. Accordingly, an up to 7 times lower cell accumulation was observed in the sludge of the DWTP with pre-ozonation (Figure 1).

In many DWTPs, the supernatant of the stored sludge is recycled to the head of the plant as the spent filter backwash water [45,57,62]. A full-scale study on a low-risk DWTP (3400 cells/mL at the intake water) documented that cyanobacterial cell counts in intake water increased by up to 43% after recycling the supernatant. Surprisingly, 80% of the transferred cells from the supernatant water were viable [60]. A recent laboratory investigation on intake water that contained 1 × 10^6 cells/mL of cultured *M. aeruginosa* reported that although conventional treatment maintained the treated effluent parameters at below WHO and USEPA guidelines, recycling of the sludge supernatant resulted in an additional increase in cells and cyanotoxins levels in the influent by up to 7 × 10^4 cells/mL and 0.26 µg/L MC-LR, respectively [63].

3. Pre-Oxidation Impact on Cyanobacteria Cells, Viability, and Cyanotoxins

3.1. Impact of Pre-Oxidation on Cyanobacteria Cell Counts

Cyanobacterial entry into DWTPs can be dampened by using pre-oxidation. Pre-oxidation may cause cell lysis, damage, and cyanotoxin release and degradation. Several studies were conducted to evaluate the pre-oxidation impact on cyanobacteria (e.g., cell viability and lysis) and cyanotoxins (release and degradation). A recent study tried to map the treatment barriers against cyanobacteria cells and cyanotoxins in drinking water facilities [10]. The results showed that the efficiency of the multi-barrier approach depends on the species present, metabolite concentration, and pre-oxidation dose [10]. Tables 1–4 summarize the literature on the impact of pre-oxidation on cyanobacteria (cultured-based and natural blooms) for four common oxidants (chlorine, ozone, potassium permanganate, 

![Figure 1. Cyanobacterial accumulation in two low risk DWTPs (maximum influx cell: <500 cells/mL). Only DWTP1 had pre-ozonation. The average values are from August to October 2011, adapted from [43].](image-url)
and hydrogen peroxide). Tables 1–4 show that CT (residual concentration × contact time) is a main driver of cyanobacterial cell lysis, damage, cyanotoxin release, and degradation.

Although some studies have reported a reduction of more than 90% in taxonomic cell counts following pre-oxidation in the lab-cultured cells, Zamyadi, et al. [35] reported a cell reduction of 70% at high chlorine exposure (CT 296 mg min/L). Fan, et al. [37] showed a limited impact of chlorine exposure (CT 104 mg min/L) on the taxonomic cell counts of *Microcystis aeruginosa* (logarithmic phase). These observations might be related to the cyanobacteria stage of life and agglomeration. Furthermore, comparing the taxonomic cell count percentage in cultured-based and natural bloom studies demonstrates the lower impact of pre-oxidation during natural blooms. Figure 2 exhibits a lower impact of pre-ozonation (2 mg/L) on cell number reduction in a natural bloom in comparison with lab-cultured cyanobacteria.

![Figure 2](image-url)

**Figure 2.** Comparison of cell count reduction following ozonation (2 mg/L) in the (a) cultured *Dolichospermum, Microcystis* [64], (b) Natural bloom [65].

Tables 1–4 and Figure 2 show that pre-oxidation, even at high CTs, may not be able to cause complete cell lysis. Consequently, it is important to clarify how far the pre-oxidation can cause viability loss and cyanotoxin release.
Table 1. Summary of the literature on the impact of pre-chlorination on cyanobacteria (cultured and natural blooms). HV: high viability, LV: low viability, DV: development stage, MA: maintenance stage.

| Dominant Cyanobacteria (Cell Density) | Lab/Field | Cl₂ Dose (mg/L) | Contact Time (min) | CT (mg min/L) | Cell Count Reduction % | Cell Viability % | Toxins | Reference | Comment |
|--------------------------------------|-----------|----------------|-------------------|---------------|------------------------|----------------|--------|-----------|---------|
| Microcystis (2 × 10⁶ cells/mL)       | Lab       | 1–2            | -                 | min. 15       | max. 90                | -              | 99% degradation | [66]    | Saline solution; exact dose and contact time were not provided; no residual; CT evaluation weak; no cell-bound |
| D. circinalis (40,000 cells/mL)      | Lab       | 2              | 0–60              | min. 1.8      | max. 50                | -              | min. 15% for CT 5.8 >100% release (CT 5.8) >90% degradation (CT 50) | [67]    | River water; using fluorescein diacetate (FDA) for viability |
| Microcystis (6 × 10⁵ cells/mL)       | Lab       | 2              | 4.5               | min. 3        | max. 296.1             | max. 76%       | 100% release (CT 5) >90% degradation (CT 35) | [35]    | River water, ultrapure water; no viability was reported |
| Microcystis (7 × 10⁵ cells/mL)       | Lab       | 3, 4, 5        | 1, 2, 5, 10, 20, 30, 60 | min. 2.8      | max. 104               | Limited impact <5% (CT 4) 25% degradation (CT 2.8) Complete degradation (CT 104) | [37,66] | Ultrapure water |
| Microcystis (2 × 10⁶ cells/mL)       | Lab       | 0.5            | 0.7               | 1.5           | 5, 11, 50, 60, 120     | min. 2.5       | max. 180       | <5% (CT 180) 10% degradation 40% increase in released | [69]    | Lake water; no CT reported |
| Microcystis (10⁶ cells/mL)           | Lab       | 0.2, 0.4, 0.8  | Range 0–480      | min. 12       | max. 396               | -              | 18% (at CT 12) 0.1% (at CT 396) - | [70]    | Lake water; no CT reported; no cell count; no toxin |
| Microcystis (10⁶ cells/mL)           | Lab       | 1, 2, 4, 8     | 1, 2, 4, 8, 16, 32, 60 | HV            | min. 0.98               | max. 361       | 95–0% reduction (CT > 15) HV CT degradation Complete (CT 108) CT degradation Complete (CT > 7) No cell viability after oxidation Same as cell death | [71]    | Ultrapure water; two viability range |
| Microcystis (1 × 10⁶ cells/mL)       | Lab       | 1, 2, 4, 8     | 1, 2, 4, 8, 16, 32, 60 | DV            | min. 3.8 max. 356      | MA min 3.7       | max 293       | >95% reduction (CT > 13.3) >95% reduction (CT > 11.9) No cell viability after oxidation Same as cell death | [72]    | Ultrapure water; two stage of life |
| Microcystis-Colony (10⁵ cells/mL)    | Lab       | 0.3, 0.5, 1, 2 | Range 0–20 min   | min. 0.97     | max. 52                | -              | Depends on colony size (0–95%) Release and degradation Colony size-dependent | [73]    | Lake water; different colony size; no cell count |
| Natural bloom US: (3 × 10⁸ cells/mL) | Field     | Cl₂/DOC: 0.105–3.6 | 0–20 min        | min. 0.15     | max. 6.8               | >80% increase (CT 6.8) 88% reduction | Complete release CT 4 (Cl₂/DOC: 0.3) Complete degradation CT 11 (US), CT 7.5 (CA) | [74]    | No cell viability; no cell count; Chl-a measured as cell damage surrogate |
| Natural blooms Canada: (3 × 10⁵ cells/mL) | Field     | Cl₂/DOC: 0.05, 0.05, 0.15, 0.1, 1 | 0–20 min        | US            | min 0.13 max 15        | CA min 0.3 max 21 Complete degradation | Complete degradation CT 11 (US), CT 7.5 (CA) | [75]    | No cell viability; no cell count; Chl-a measured as cell damage surrogate |
Table 1. Cont.

| Dominant Cyanobacteria (Cell Density) | Lab/Field | Cl₂ Dose (mg/L) | Contact Time (min) | CT (mg min/L) | Cell Count Reduction % | Cell Viability % | Toxins | Reference | Comment |
|--------------------------------------|-----------|----------------|-------------------|--------------|------------------------|----------------|--------|-----------|---------|
| Natural bloom (3.3 × 10⁵ cells/mL)   | Field     | 0.2, 0.6       | 0–120 min         | min 0.15 max 3.84 | min. CT 5% decrease | min CT: 34% decrease | CT 3.84: 23% decrease | [76]  | Soft chlorination (low dose) |
| Natural bloom                        | Field     | 2, 5           | 0–60 min          | min 1.14 max 14.8 | min. < 5% reduction | max. > 50% reduction | -      | 2 mg, CT 10, >200% release 5 mg, CT 20, 200% release | [77]  | No cell viability |

Table 2. Summary of the literature on the impact of pre-ozonation on the cyanobacteria (cultured and natural bloom).

| Dominant Cyanobacteria (Cell Density) | Lab/Field | O₃ Dose (mg/L) | Contact Time (min) | CT (mg min/L) | Cell Count Reduction % | Cell Viability % (for CT) | Toxins | Reference | Comment |
|--------------------------------------|-----------|----------------|-------------------|--------------|------------------------|--------------------------|--------|-----------|---------|
| Microcystis (2 × 10⁵ cells/mL)       | Lab       | 1              | -                 | min. 1.2 max 16 | -                      | CT > 54, complete loss    | CT > 12 complete degradation | [66]  | Saline solution; exact dose and contact time not provided; no residual; CT evaluation weak |
| Microcystis (7 × 10⁵ cells/mL)       | Lab       | 2, 4, 6        | 5                 | < 0.22 max 2.29 | -                      | Min CT: 50%, Max CT: 85% | >100% release (high CT) 50% degradation | [37,68] | Ultrapure water |
| Microcystis (2.5 × 10⁶ cells/mL)     | Lab       | 0.5, 2, 4      | 0.5–10            | min. < 0.2 max 22 | 32% for 2 mg/L, 41% for 4 mg/L | Complete loss, CT < 0.2 | -      | [64]  | River water; Chl-a measured as cell damage surrogate; no toxin measurement |
| Microcystis, Dolichospermum         | Lab       | 0.63–5         | 24 h              | min 0.5 max 17 | 100% reduction (CT 0.5) | Complete loss, (CT > 2) | -      | [78]  | Ultrapure; no flow cytometry |
| Microcystis, Oscillatoria           | Lab       | 0.5, 1, 2      | 5, 10             | max. 2.5     | >95% reduction          | Complete loss            | -      | [79]  | Natural water; no toxin measurement |
| Microcystis, Dolichospermum (1.2 × 10⁵–2 × 10⁶ cells/mL) | Field     | 2, 3, 4, 5     | 0–10              | min. 1.4 max 16 | 75% reduction (CT 16.8) | CT 3.2: 45% CT 16.8: 15% | CT < 2, more than 100% release | [65]  | Natural bloom |
| Natural bloom (US (3 × 10⁶ cells/mL)—Planktothrix agardhii CA (3 × 10⁶ cells/mL)—D. spiroide) | Field | O₃/DOC: 0.05–0.75 | 0–20             | US-min. 1.5 max. 3 CA-min 0.2 max. 4.1 | -                      | -                      | >80% degradation (CT 4.1(CA)) | [75]  | No cell viability; no cell count; Chl-a measured as cell damage surrogate |
| Natural bloom                        | Field     | 0.1, 0.3       | 0–10              | max 0.86     | max CT 14% decrease     | max CT: 79%              | 14% degradation No release | [76]  | Soft ozonation (low dose) |

Table 3. Summary of the literature on the impact of potassium permanganate on cyanobacteria (cultured and natural bloom).

| Dominant Cyanobacteria (Cell Density) | Lab/Field | KMnO₄ Dose (mg/L) | Contact Time (h) | CT (mg min/L) | Cell Count Reduction % | Cell Viability % (for CT) | Toxins | Reference | Comment |
|--------------------------------------|-----------|------------------|-----------------|--------------|------------------------|--------------------------|--------|-----------|---------|
| Microcystis (2 × 10⁵ cells/mL)       | Lab       | 1–2              | -               | min 0.15 max 600 | -                      | min CT: 60%, CT > 60 complete loss | CT: 30 Complete dissolved degradation | [66]  | Saline solution; exact dose and contact time were not provided; no residual; CT evaluation weak |
Table 3. Cont.

| Dominant Cyanobacteria (Cell Density) | Lab/Field | KMnO₄ Dose (mg/L) | Contact Time (h) | CT (mg min/L) | Cell Count Reduction % | Cell Viability % (for CT) | Toxins | Reference | Comment |
|--------------------------------------|-----------|-------------------|------------------|---------------|------------------------|--------------------------|--------|-----------|---------|
| Microcystis (7 × 10⁶ cells/mL)        | Lab       | 1, 5, 10          | 0.25–7           | min. 28.7 max. 2642 | 14% cell number reduction (CT max) | CT 260: complete loss | Release at CT > 70 Complete degradation CT 2600 | [37,68] | Ultrapure water |
| Microcystis, Dolichospermum (4 × 10⁶ cells/mL) | Lab | 2, 5 | 20 | max. 456 | 10% reduction at highest CT | CT 456: 18% viability | - | [80] | Natural water; no toxin measurement |
| Microcystis Bloom from Lake Erie | Lab, Field | 0.5–8 | 1–5 | min. 120 max. 1920 | - | Cell, CT 1920: 2% Bloom, CT 1920: 40% | - | [81] | No cell count and toxin; no CT; CT with lower doses was unable to decrease viability |

Table 4. Summary of the literature on the impact of hydrogen peroxide on cyanobacteria (cultured and natural bloom). h: hour, d: day.

| Dominant Cyanobacteria (Cell Density) | Lab/Field | H₂O₂ Dose (mg/L) | Contact Time | CT (mg h/L) | Cell Count Reduction % | Cell Viability % (for CT) | Toxins | Reference | Comment |
|--------------------------------------|-----------|------------------|--------------|-------------|------------------------|--------------------------|--------|-----------|---------|
| Microcystis (3.7 × 10⁶ cells/mL)      | Lab       | 3.4, 17          | 4 h, 2 d, 4 d | min. 13.6 max. 1632 | min. CT 8% reduction max. CT 89% reduction | K⁺ release min. CT 81% max. CT 3% reduction | CT > 816 26% MC release | [82] | K release as a surrogate for cell damage; no CT provided |
| Microcystis (7 × 10⁶ cells/mL)        | Lab       | 10.2, 51, 102    | 0.1–7h       | min. 189.3 max. 17,678 | Limited change | min. CT 86% max. CT 7% reduction | No release, CT 364 >95% degradation | [37,68] | Ultrapure water |
| Pseudanabaena (10⁴ cells/mL)          | Lab       | 3, 5, 10, 20     | 2 h, 4 h, 8 h 2 d, 4 d | min. 6 max. 960 | min. CT: No change max. CT >90% reduction | CT 120: 2% | - | [83] | Reservoir water; no toxins |
| Microcystis (6 × 10⁶ cells/mL)        | Lab       | 1–15             | 0.1–7d       | min. 2.4 max. 2520 | CT 1680: 95% reduction | max. CT 3% viability | CT > 1512, 82% degradation | [84] | Culture; no CT provided |
| Microcystis, Dolichospermum (4 × 10⁶ cells/mL) | Lab | 5, 10 | 6 h | min. 13.9 max. 96.1 | <5% reduction | min. CT 39% max. CT 30% reduction | - | [79] | Natural water |
| Natural bloom: D. ovalis: (5.43 × 10⁵ cells/mL) | Field | 10 | 6 h–1 d | min. 47 max. 140.7 | max. CT 52% reduction | min. CT 60% max. CT 40% reduction | No release max., 15% MC degradation | [76] | - |

3.2. Chlorination

Figure 3 is a reconstructed graph from the cell viability results following pre-chlorination based on the oxidant exposure (CT). Parameters such as background water quality (e.g., pH and dissolved organic carbon (DOC)) which have an impact on the oxidant demand are included in the CT concept. Therefore, a comparable level of damage should be found by comparing cell viability results using oxidant exposure (CT) for lab-cultured and natural blooms. Figure 3 demonstrates that at the same level of chlorine exposure, the natural bloom is more resistant to pre-chlorination as compared to lab-cultured cells. In other words, lab-cultured studies are not representative of natural bloom pre-chlorination. Fan, et al. [73] reported that the level of cell damage and toxin release depends on the colony size. Figure 3b shows the cultured-based studies fitted with the Chick–Watson equation. Although the results from different unicellular studies are aligned with each other, the colonial *Microcystis* chlorination shows a different cell damage rate. This could be related to the agglomeration of cyanobacteria cells and increasing the mucilage sheath in colonial cyanobacteria [73]. Despite using the CT calculation to compare the results, Figure 3c demonstrates different cell damage rates for each study of real bloom after chlorination, and the same level of chlorine exposure may not result in the same level of cell damage. Figure 3c shows that cyanobacterial bloom oxidation could be site- and bloom-specific, depending on the agglomeration, cyanobacteria (bloom) stage of life, and metabolic functions. Higher cell damage following pre-oxidation (especially with higher CTs) can lead to
higher cyanotoxin release, which cannot be removed during conventional treatment. Soft chlorination showed cell damage by up to 45% and total microcystin (MC) degraded by up to 23%, while no cyanotoxin release was observed [76]. In addition, soft chlorination may cause lower disinfection by-products as a lower oxidant concentration is used in this approach.

3.3. Ozonation

Figure 4 shows the impact of pre-ozonation on cyanobacteria cell damage for cultured-based and natural bloom studies. Figure 4a demonstrates lower cyanobacteria cell damage for a specific ozone exposure for natural blooms as compared to the lab-cultured cyanobacteria. The model fit results (Figure 4b) show a higher cell damage rate for the lab-cultured cyanobacteria in comparison to the natural bloom. As per soft oxidation, soft pre-ozonation was reported to cause up to 21% of cell damage and 14% of MC degradation, while no MC release was observed simultaneously [76]. Such an observation implies the effectiveness of soft pre-ozonation to damage the cells without cyanotoxin release.

3.4. Potassium Permanganate

Figure 5 demonstrates that the viability loss of the lab-cultured studies harvested in the logarithmic phase is lower than those that harvested in the stationary phase. This observation implies the impact of the cyanobacteria stage of life on pre-oxidation efficiency. A comparison of the cell viability results of the lab-cultured with natural bloom studies following potassium permanganate pre-oxidation confirms the higher resistance of real cyanobacterial bloom cells (Figure 5). In addition, the degradation rate constant of dissolved MCs was higher than that released by MCs for high potassium permanganate doses [36,81].
Oscillatoria, Microcystis, and Lyngbya asp. [37,64,68,78,79], and natural blooms [65,76] following pre-ozonation. (b) Cell viability experimental data and fitted model for cultured-based and natural bloom samples following pre-ozonation.

Figure 4. (a) Comparison of the cell viability results of cultured-based Microcystis, Dolichospermum, Oscillatoria, Lyngbya asp. [37,64,68,78,79], and natural blooms [65,76] following pre-ozonation. (b) Cell viability experimental data and fitted model for cultured-based and natural bloom samples following pre-ozonation.

3.5. Hydrogen Peroxide

Matthijs, et al. [86] reported that a concentration of 2 mg/L H₂O₂ was able to decrease cyanobacteria (natural bloom) by two logs within 3 days. In addition, cyanobacteria remained at a low abundance level for 7 weeks following H₂O₂ addition. Figure 6 demonstrates that natural blooms are more resistant to H₂O₂ than the lab-cultured cyanobacteria, as observed for other oxidants (Figures 2–5). Foo, et al. [87] reported that the impact of H₂O₂ on cyanobacteria is dependent on the residual concentration (C) and contact time (T). In addition, the authors concluded that toxic and non-toxic Microcystis aeruginosa are impacted by H₂O₂ with the same trend. Zhou, et al. [84] stated that a low dose of H₂O₂ (<5 mg/L) would have a low and recoverable impact on the lab-cultured Microcystis. On
the other hand, the higher the H₂O₂ dose (>8 mg/L), the higher necrosis, cell death, and consequent cyanotoxin release. A medium dosage of H₂O₂ with low to medium contact time can activate apoptosis-like programmed cell death (AL-PCD) [84]. The cellular energy required for AL-PCD is provided from the transcriptional, biochemical, and structural changes. Zhou, et al. [84] documented the maximum cell death with low MC production by AL-PCD activation. Zamyadi, et al. [17] studied the impact of H₂O₂ on blooms and lab-cultured cyanobacteria (Microcystis aeruginosa). The results highlighted a delayed impact of H₂O₂ on cyanobacteria cells after complete depletion of H₂O₂ during stagnation (up to one week) [17]. Chl-a and phycocyanin (PC) fluorescence were significantly declined by 93% and 74% in natural bloom and lab-cultured samples, respectively. Additionally, the lab-cultured results revealed delayed MC release during stagnation [17].

Figure 6. Comparison of the cell viability results of cultured-based cyanobacterial cells (Microcystis, Pseudanabaena) [37,68,83,84] and natural bloom cells [76,86] after oxidation by hydrogen peroxide.

Besides the current oxidants, peracetic acid (PAA) has been used in wastewater treatment facilities as a disinfection alternative for chlorine [88]. Almuhtaram and Hofmann [89] studied the impact of PAA and PAA/UV on cyanobacteria and cyanotoxin removal. The results show that 10 mg/L of PAA with 60 min contact time was able to degrade MC-LR by 80% (3.46 M⁻¹ s⁻¹ lower reaction rate as compared to HOCl 1.2 × 10² M⁻¹ s⁻¹). In addition, the results elaborated that PAA alone can barely remove cyanobacteria, except at a high dose (10 mg/L) and with lower cyanobacterial cell counts (10⁵ cells/mL).

3.6. Considerations on the Impact of Pre-Oxidation on Downflow Processes

The impact of pre-oxidation on downflow processes should also be considered as it may influence the removal of cyanobacteria by coagulation, flocculation, and sedimentation. Previous studies have been reported that pre-oxidation has a positive impact on enhancing cyanobacterial removal through coagulation/flocculation and sedimentation [27,28,31,32,34,90]. Pre-oxidation can cause morphological deformation [82] and changes in the surface charge of the cells, leading to increased cell removal efficiency during coagulation/flocculation [37].

KMnO₄ increases the binding potential to the coagulant by oxidizing organic matter (extracellular and released cell-bound) to lower molecular weight fractions, as well as forming colloids (by MnO₂) to be adsorbed to the cyanobacterial cells and forming larger flocs [32,34,81]. Xie, et al. [27] reported that KMnO₄ exposure (CT: 10 mg min/L, estimated) could increase cyanobacteria cell removal by 22% during coagulation/flocculation. In addition, pre-ozonation with CT: 4, 10, and 20 mg min/L (estimated) led to an increase in cyanobacteria cell removal during coagulation by 14%, 20%, and 24%, respectively [27]. Cyanobacteria cell removal during coagulation was improved in a full-scale DWTP equipped by pre-ozonation systems (CT: 2.52–3.78 mg min/L (estimated)) [43]. Pre-oxidation may cause metabolite release (organic matter and cell-bound cyanotoxins) following cyanobacterial cell damage. Besides the challenge to remove dissolved cyanotoxins, coagulation efficiency can be compromised by high algal organic matter release following pre-oxidation. Xie, et al. [27] showed that due to pre-ozonation with CT > 4 mg
min/L (estimated), cyanobacteria cell viability was completely degraded, and consequently, organic matter concentration increased. Further, Barešová, et al. [91] demonstrated that pre-ozonation (CT < 40 mg min/L (estimated)) could interrupt the coagulation (Al/Fe-based) efficiency of DOC removal (in comparison with higher C Ts) due to the degradation of high molecular weight algal organic matter to low molecular weight compounds.

It is noteworthy to recall that H\textsubscript{2}O\textsubscript{2} can have a delayed impact on cyanobacteria and, potentially, cyanotoxin release after complete degradation of the oxidant [17]. This delayed cyanotoxin release should be considered in the downstream processes, as well as in sludge handling.

The oxidant exposure must be adjusted to maximize cell damage and cyanobacteria cell removal (directly or after coagulation) and minimize cyanotoxin release and cell accumulation in the sludge, simultaneously. Figure 7 summarizes the pre-oxidation (soft and normal) advantages/disadvantages of cyanobacteria and cyanotoxins during water treatment. In fact, soft pre-oxidation (low CT of Cl\textsubscript{2} and O\textsubscript{3}) can (1) partially degrade cyanobacteria cells, (2) cause low cyanotoxin release, (3) improve coagulation efficiency to remove cells, and (4) cause low cell accumulation in the downflow processes.

**Figure 7.** Summary of pre-oxidation (low and medium-high CT) impact on cyanobacteria/cyanotoxins and downflow processes (#: very low impact, +: increase, and -: decrease). Low CT for Cl\textsubscript{2} = CT < 4 mg min/L, low CT for O\textsubscript{3} = CT < 1 mg min/L, low CT for KMnO\textsubscript{4} = CT < 50 mg min/L, and low CT for H\textsubscript{2}O\textsubscript{2} = CT < 50 mg h/L.
4. Sludge Storage, Oxidation, and Handling

Cyanobacteria and cyanotoxins (cell-bound) accumulate in the sludge of clarifiers throughout the flocculation/coagulation/sedimentation processes. This cyanobacteria-laden sludge remains in the sludge holding tank before disposal. In addition, potential options to treat cyanobacteria-laden sludge need to be considered. Furthermore, safe (healthy, both operationally and environmentally) cyanobacteria-laden sludge handling approaches are required.

4.1. Fate of Cyanobacteria and Cyanotoxins during Sludge Storage

Several studies (Table 5) demonstrated that cyanobacteria cells could stay viable within 2–12 days in the stored sludge. The loss of viability and consequent cyanotoxin release caused an increase in dissolved cyanotoxin concentrations during sludge storage [13,40,45–52]. However, dissolved cyanotoxins in stored sludge can be adsorbed onto the remained PAC injected into the intake water [56], flocs [50] or it can be biodegraded by cyanotoxin degrader species [58,92].

Besides cell survival potential during sludge storage, some studies have hypothesized that cyanobacteria can also grow in stored sludge [53–55]. Water Research Foundation (WRF) and Water Research Australia [53] documented that concentrations of DOC, MC-LR, and cylindrospermopsin in stored coagulated sludge contained M. aeruginosa and C. raciborskii exceeded the expected concentrations by 4–10-fold based on the cell quota (if all cell-bound metabolites are released) within 7–16 days, respectively. Dreyfus, et al. [55] studied the fate of stored sludge that contained cultured M. aeruginosa, D. circinale, and C. raciborskii within 18 days. The authors demonstrated that DOC, MC-LR, MC-LA, and CYN concentrations increased by up to 5-, 2.2-, 1.2-, and 2.5-fold during storage, respectively. Another investigation on stored sludge containing cultured M. aeruginosa and D. circinale reported that taxonomic cell counts increased by up to 4.2-fold in sludge stored in a lagoon within 7 days [54]. The authors also reported that the concentrations of cyanobacterial metabolites increased by up to 5 times in the sludge supernatant within 20 days. In the worst case, cyanobacteria could survive by up to 35 days in the stored sludge [54]. Despite the important findings of the previous studies on cell survival and metabolite release during sludge storage, cyanobacterial cell growth during sludge storage is yet to be explored in detail. In these studies, cell and metabolite increase during sludge storage might be due to the cell growth or to either the (1) underestimation of cell quota, (2) increase of metabolite production per cell during storage, or (3) additional cell settling from the supernatant to the sludge during the storage [53–55].

Our recent study on the cyanobacteria-laden sludge of a DWTP documented cell depletion, survival, and growth in different sludge samples [58]. Cell growth was observed in four out of eight sludge samples (different sampling dates) stored in the dark for 7–38 days. In the worst-case scenario, taxonomic cell counts increased from 2.7 × 10^6 to 5.3 × 10^6 cells/mL within 16 days (96% cell growth). Cell growth was also confirmed by increasing cyanobacterial biomarkers such as the “Pentose phosphate pathway” marker, which is responsible for the heterotrophic growth of cyanobacteria [93].

Table 5. Impact of sludge storage on cyanobacteria and cyanotoxins. STX: saxitoxin, PACl: colyalumnum chloride, CTSAC: Chitosan-aluminum chloride.

| Initial Characteristics of Cyanobacteria/Coagulation/Sedimentation Process | Initial Condition of Cyanobacteria and Cyanotoxins in the Stored Sludge | Observation | Reference |
|---|---|---|---|
| Cultured M. aeruginosa (1 × 10^6 cells/mL) (Jar test, 70 mg/L alum) | 8 × 10^6 cells/mL, 2500 µg MC-LR/L | Cell survival (2 days); cell lysis and cyanotoxin release (2 days); degradation of dissolved cyanotoxins (8–10 days) | [13] |
Table 5. Cont.

| Initial Characteristics of Cyanobacteria/Coagulation/Sedimentation Process | Initial Condition of Cyanobacteria and Cyanotoxins in the Stored Sludge | Observation | Reference |
|---|---|---|---|
| Cultured *D. circinale* and *C. raciborskii* (1.0 \times 10^5 cells/mL) (Jar test, 40 mg/L Al(III)) | Sludge supernatant: *D. circinale*: 1300 cells/mL STX: 0.4 µg/L | Cells remained viable up to 7 days; cell lysis and toxin release within 3 days | [45] |
| Cultured *M. aeruginosa* (2 \times 10^6 cells/mL) (Jar test, 15 mg/L AlCl_3) | 18 µg/L dissolved MCs | Cell lysis and cyanotoxin release after 6 days | [40] |
| Cultured *M. aeruginosa* (1 \times 10^6 cells/mL) (Jar test, 4 mg/L PACl-optimum dose) | 20 µg/L dissolved MCs | Cell lysis and cyanotoxin release within 6–12 days | [46] |
| *Microcystis flos aquae* (5.2 \times 10^5 cells/mL) (Jar test, 100 mg/L Al(III)) | Sludge supernatant: MC-RR, MC-YR: < 2 µg/L | Cell survival (5 days); cell lysis and cyanotoxin release (5–10 days); degradation of dissolved cyanotoxins (up to 15 days) | [62] |
| Cultured *M. aeruginosa* (1 \times 10^6 cells/mL) (Jar test, 15 mg/L AlCl_3, 4 mg/L PACl) | ~0.9 bar vacuum pressure for dewatering the sludge 23 µg/L total MCs | Cell lysis and cyanotoxin release within 4–6 days; optimum sludge storage time for AlCl_3 and PACl was suggested to be 4 and 2 days, respectively. | [47] |
| Cultured *M. aeruginosa* (1 \times 10^6 cells/mL) (Jar test, 0–70 mg/L FeCl_3) | ~1 µg/L dissolved MCs | Cell lysis and cyanotoxin release (2–8 days); degradation of dissolved cyanotoxins (> 10 days) | [48] |
| *Myponga reservoir* Cultured *M. aeruginosa* (2.3 \times 10^5 cells/mL) Cell-bound MC-LR: 4.7 µg/L Dissolved MC-LR: 2.0 µg/L (Jar test-80 mg/L Al(III)) | Sludge supernatant after 1 day storage: Cells: 4300 cells/mL Cell-bound MC-LR: 0.5 µg/L Dissolved MC-LR: 2.5 µg/L | Cell survival (4 days); cell lysis and cyanotoxin release (4–7 days); degradation of dissolved cyanotoxins (> 4 days) | |
| *Myponga reservoir* Cultured *M. aeruginosa* (3.1 \times 10^5 cells/mL) DOC: 10.1 mg/L Cell-bound MC-LR: 5.0 µg/L Dissolved MC-LR: 2.9 µg/L (Jar test-80 mg/L Al(III)) | Sludge supernatant after 1 day storage: DOC: 5.2 mg/L Cell: 2760 cells/mL Cell-bound MC-LR: <DL Dissolved MC-LR: 2.5 µg/L | Cell growth (within 7–16 days) confirmed by DOC and MC-LR cell quota | |
| *Myponga reservoir* Cultured *C. raciborskii* (3.1 \times 10^5 cells/mL) DOC: 10 mg/L Cell-bound CYN: 2.5 µg/L Dissolved CYN: 0.7 µg/L (Jar test-80 mg/L Al(III)) | Sludge supernatant after 1 day storage: DOC: 6.0 mg/L Cell: 7080 cells/mL Cell-bound CYN: 1.0 µg/L Dissolved CYN: 0.8 µg/L | Cell growth (within 7–23 days) confirmed by DOC and CYN cell quota | |
| *River Murary* Cultured *C. raciborskii* (3.1 \times 10^5 cells/mL) DOC: 8.63 mg/L Cell-bound CYN: 2.7 µg/L Dissolved CYN: 0.3 µg/L (Jar test-80 mg/L Al(III)) | Sludge supernatant after 1 day storage: DOC: 4.9 mg/L Cell: 4140 cells/mL Cell-bound CYN: 0.3 µg/L Dissolved CYN: 0.9 µg/L | Cell growth (within 15–23 days) confirmed by DOC and CYN cell quota | |
| Initial Characteristics of Cyanobacteria/Coagulation/Sedimentation Process | Initial Condition of Cyanobacteria and Cyanotoxins in the Stored Sludge | Observation | Reference |
|---|---|---|---|
| Cultured *M. aeruginosa* (1 × 10⁶ cells/mL) (Jar test- 15 mg/L AlCl₃, 50 mg/L FeCl₃, 15 mg/L PAFC) | 20 µg/L dissolved MCs 1-4.2 mg/L dissolved polysaccharides 4 mg/L chla | Cell lysis and toxin release (2–10 days) | [49] |
| Cultured *M. aeruginosa* (2 × 10⁶ cells/mL) (Jar test, 2.6 mg/L chitosan− 7.5 mg/L AlCl₃ (CTSAC)) | 9 µg/L dissolved MCs (after coagulation) 18 µg/L dissolved MCs (without coagulation); the difference is due to adsorption in CTSAC | Toxin release (0–4 days); degradation of dissolved cyanotoxins (6–10 days) | [50] |
| *M. aeruginosa*, *D. circinale*, *C. raciborskii* (3.0 × 10⁵ cells/mL) (Jar test, 80 mg/L Alum) | Sludge supernatant after 1 day storage: DOC: 5.2–6.5 mg/L Cell: 2162–7080 cells/mL Cell-bound MC-LR: <0.5 µg/L Dissolved MC-LR: 2.5–4.0 µg/L Cell-bound CYN: 1.0 µg/L Dissolved CYN: 0.8 µg/L | Increased DOC, MC-LR, MC-LA, and CYN to higher the expected values (hypothesis: increase of the metabolite production, cell growth or both) | [55] |
| *M. aeruginosa* and *D. circinale* (8.6 × 10⁴ cells/mL) (Jar test, 80 mg/L Alum) | Non-coagulated sludge: 5.0 × 10⁶ cells/mL Coagulated sludge: 5.4 × 10⁵ cells/mL | Cell survival (up to 35 days); 4.2× increase in cell counts in the sludge lagoon within 7 days; increased metabolites to higher the expected values (up to 5×); increased cell counts in the sludge (hypothesis: cell growth, additional settling, or both) | [54] |
| Cultured *M. aeruginosa* × 10⁵ cells/mL (Jar test, 15 mg/L AlCl₃, 50 mg/L FeCl₃, 15 mg/L PAFC) | 1 µg/L dissolved MCs | Cell lysis and toxin release (4–6 days); degradation of dissolved cyanotoxins (6–10 days) | [51] |
| Cultured Oscillatoria sp. (1.0 × 10⁴ cells/mL) (Jar test, 5 and 10 mg/L PAFC) | 1.0 mg/L chla 2.3 µg/L cell-bound protein 8.6–11.4 µg/L dissolved CYN | Increase in chla level after 4 days, suggesting cell growth; loss of cell integrity after 2 days, while cells remained viable up to 8 days; increase in dissolved CYN, showing toxin release within 4 days | [94] |
| Cultured *C. raciborskii* (1 × 10⁶ cells/mL at late exponential phase) (Jar test, 10 mg/L PAFC) | 1.1 µg/L dissolved CYN 2 mg/L cell-bound protein | Cell lysis and toxin release after 6 days; degradation of dissolved cyanotoxins after 10 days | [52] |
| n/a | Sludge of a DWTP containing natural cyanobacterial blooms stored for 7–35 days in the darkness (8 samples). 0.7 × 10⁵–5.6 × 10⁶ cells/mL 25–7130 ng/L cell-bound MCs 38–349 µg/L dissolved MCs | Cell growth in 4/8 samples after 9–35 stagnation days; cell death in the rest 4/8 samples; degradation of dissolved cyanotoxins after 8 days | [58] |
4.2. Cyanobacteria-Laden Sludge Treatment

A summary of studies on the treatment of cyanobacteria-laden sludge is presented in Table 6. The available data demonstrated that sludge oxidation could not completely remove cyanobacteria cells and metabolites from the sludge [57,59]. Sludge is often stored after oxidation, while its supernatant can be recycled to the head of the DWTP. Thus, the impact of oxidation on sludge storage should be investigated. Recent findings showed no remarkable benefits in sludge oxidation followed by sludge storage as compared to only sludge storage [59]. The maximum additional taxonomic cell count decreased by a combination of oxidation (KMnO$_4$ or H$_2$O$_2$) and storage was 32% as compared to storage only. However, oxidation/storage could cause a remarkable cell growth (by up to 145%) and toxic gene copy numbers of mcyD increase (by up to 13.0×) in some sludge samples [59]. This phenomenon can be attributed to gene expression regulation due to the presence of oxidative stresses [58,59,95,96]. Similarly, sludge oxidation could not completely remove cyanobacteria and cyanotoxins from the supernatant sludge [59]. Finally, the costs and by-product formation during the oxidation of organic-matter-rich sludge should be considered [58].

Table 6. Data of cyanobacteria-laden sludge treatment. MIB: 2-Methylisoborneol.

| Source of Sludge | Scale | Treatment Agent/Dosage | Contact Time | Initial Conditions | Cell Count Reduction | Metabolite Reduction | Reference |
|------------------|-------|------------------------|--------------|-------------------|---------------------|---------------------|-----------|
|                  |       | Laboratory             | 3 mg/L KMnO$_4$ | 2 h               | $5.0 \times 10^4$ cells/mL $Pseudanabaena$ | >95% | - |
| Sludge thickener |       | Laboratory 10–100 mg/L PAC | 1 h          | 100/L MIB | - | 42–100% MIB | [57] |
|                  |       | Full-scale             | 10 mg/L KMnO$_4$ | 15 h (max.) | $4.3 \times 10^5$ cells/mL (natural blooms) | 13–98% total and $Pseudanabaena$ cell counts | - |
|                  |       | Full-scale             | 10 mg/L KMnO$_4$ | 24–72 h | KMnO$_4$: 120 ng/L MIB (natural blooms) | 40–52% in total and $Pseudanabaena$ cell counts | 20–22% MIB |
|                  |       | Laboratory             | 5 mg/L KMnO$_4$ | 60 min | $2.3–2.7 \times 10^6$ cells/mL | 46–55% total cell counts | 0.3–24% MCs |
|                  |       | Laboratory             | 10 mg/L KMnO$_4$ | 63–161 ng/L MCs (natural blooms) | 59–62% total cell counts | 2–32% MCs |
|                  |       | Sludge holding tank    | 10 mg/L H$_2$O$_2$ | 24 h | 58% total cell counts | 27% MCs |
|                  |       | Full-scale             | 10 mg/L KMnO$_4$ | 24–72 h | $2.4 \times 10^6$ cells/mL 88–1083 ng/L MCs (natural blooms) | 24–43% total cell counts (31% cell count increase after 48 h in one sample) | MCs: 3–25% decrease in one sample 37–589% increase in one sample | [59] |

4.3. Sludge Handling Challenges

In general, the sludge supernatant is recycled to the head of the DWTP or is discharged into the source [13,56,97]. The solid phase either can be transferred to the WWTP or is ap-
plied for landfilling [98–100]. Less environmentally friendly approaches such as untreated residual discharge into lakes or ponds can be also applied in some circumstances [101]. The re-use of DWTPs’ residuals is growing [99,102–104]. However, an investigation demonstrated that the half-life of MC analogs varies from 8 to 18 days in soil [105]. Since there is a risk of soil and groundwater contamination, landfill and field applications of cyanobacteria-laden sludge should be avoided. Overall, cyanobacteria-laden sludge should be treated before disposal either in situ or via sending it to wastewater treatment plants.

Flocs may have a protective role during sludge storage for *M. aeruginosa* [40,46–48,52]. In contrast, Li, et al. [52] documented that polyaluminium ferric chloride (PAFC) can stimulate the lysis of *C. raciborskii* and CYN release by up to 94% during sludge storage. This may occur in the sludge and lead to cyanotoxin release. However, all studies have been conducted in laboratory conditions and on cultured-based cyanobacteria. In fact, due to complex parameters such as the presence of various cyanobacterial cells in various forms (aggregated, multicellular), ages, and viabilities, the design of such experiments in full scale is complex.

Stresses such as oxidation and storage can shift cyanobacterial communities towards resistant genera (e.g., *Microcystis* and *Aphanocapsa*), which can produce MCs [56,58,59]. Thus, the survival probability of MC producer species can increase during sludge oxidation or storage. The fate of cyanotoxins in the sludge is complex due to the simultaneous occurrence of various phenomena such as cell survival, growth, lysis, cell-bound cyanotoxin release, and released cyanotoxin degradation [33–56,99,105,106]. Based on the increased risk of cell lysis and cyanotoxin release during sludge storage, some studies have suggested that cyanobacteria-laden sludge should be disposed of prior to 4 days to avoid metabolite release [47,51,108]. However, these studies only focused on metabolite release and not on cell survival/growth phenomena. Additionally, the possibility of sludge disposal can be a technical and financial challenge in large DWTPs.

5. Decision Framework to Manage Cyanobacteria and Cyanotoxins in Drinking Water Treatment Plants

5.1. Framework Basis

Since cyanobacterial cells and their associated metabolites, including cyanotoxins, as well as taste and odor agents such as geosmin and 2-Methylisoborneol (MIB), affect water and sludge quality, monitoring should be applied for the evaluation of the water treatment chain and sludge handling.

Microscopy taxonomic cell count techniques have been widely applied to evaluate the water and sludge in previous studies [12,18,43,44,56,77]. Previous cyanobacterial monitoring guidelines were prepared based on taxonomic cell counts and biovolumes [109–111]. A recent study suggested 0.3 mm$^3$/L biovolumes as the vigilance level [109]. However, bias related to human error [112], the negative impact of Lugol’s iodine on biovolumes [113], cell underestimation/overestimation due to the presence of aggregated cells [114], and the presence of debris and sediments, especially in the sludge samples [56], may affect the results. More importantly, the significant time required for taxonomic cell counts is a major barrier in using them for a real-time/practical approach.

In situ fluorometry using on-line probes is a compromising technique for measurement of PC based on relative fluorescence units (RFU) in water resources [18,115–119]. However, the correlation between RFU and biovolume is complex and site-specific [119–121]. Previous investigations have reported that various RFUs ranging from 0.7 to 1.8 could correlate with a 0.3 mm$^3$/L biovolume in different sources and bloom events [116,119]. Therefore, it is recommended to perform a correlation between on-line probe readings (RFU) and biovolumes for each water resource. It is noteworthy that the limits of detection and quantification of on-line probes should be considered [116,118].

MC concentration has been introduced in several guidelines such as those of the WHO (1.0 µg/L MC-LR) [122,123] and Health Canada (1.5 µg/L) [124]. Geosmin and MIB negatively affect water quality, raise complaints about taste and odor, and decrease
the public’s confidence about the treated water safety [125–129]. Thus, they should be monitored throughout the treatment chain and the recycled sludge supernatant. Using enzyme-linked immunoassay (ELISA) tests for cyanotoxins measurement has been accepted for cyanotoxin monitoring [44,130–132]. The reported thresholds for geosmin and MIB are 1.3–4.0 ng/L and 6.3–15 ng/L, respectively [133–135]. Since the taste and odor agents are not harmful, but increase complaints and concerns about the water quality [126,127,129,134], olfactory detection can be considered for monitoring and detection [128,136].

5.2. Decision Framework

A decision framework to manage cyanobacteria and cyanotoxins in DWTPs is presented in Figure 8. The objective of this framework is to minimize cell breakthrough and accumulation throughout DWTPs and sludge. The three steps, (i) source water risk assessment, (ii) treatment breakthrough assessment and management, and (iii) sludge and supernatant risk assessment and management, should be taken for cyanobacteria and cyanotoxin control in DWTPs and sludge. This framework can help water utilities to understand appropriate approaches/strategies against cyanobacteria and cyanotoxins in DWTPs.

Overall, taxonomic cell counts, MCs, and taste and odor agents should be monitored in the (i) intake water, (ii) treatment chain, and (iii) sludge supernatant. These points are subjected to cyanobacteria and cyanotoxin accumulation, leading to a negative impact on water quality.

The optimization of conventional processes may include coagulant dose adjustment, applying aid-flocculants, and lowering the sedimentation and filtration rate during cell breakthrough [12,137–139]. Secondary barriers such as pre-oxidation, PAC injection, and GAC, in case of metabolite breakthrough, should be applied [15,25,39,65,140–144]. The impact of supernatant recycling or discharging on the source/intake water quality should be considered during toxic cyanobacterial blooms. Supernatant treatment may be required in the presence of cyanotoxins or taste and odor agents. MC concentration levels should be monitored in the sludge (solids) in case of landfelling or land application. In the case of elevated concentrations of MCs, sludge treatment is required.
6. Conclusions

- Using the exposure unit (CT) is recommended for cyanobacteria and cyanotoxins oxidation studies, rather than using dose or contact time individually.
- Regardless of the oxidant type, lab-cultured studies cannot depict the complete picture of natural cyanobacterial bloom behavior during oxidation and may overestimate the oxidation efficiency. In addition, cyanobacterial bloom oxidation is site- and bloom-specific, which could be related to the level of agglomeration, cyanobacteria (bloom) stage of life, and metabolic functions.
- Soft pre-chlorination and pre-ozonation can compromise cell viability with no or limited cyanotoxin release. Overall, soft pre-oxidation may cause lower disinfection by-products compared to normal pre-oxidation.
- The cyanobacteria in stored sludge can not only survive, but also grow and release cyanotoxins, even in the dark. Although dissolved cyanotoxins can be degraded during sludge storage, the potential risk of growth and cyanotoxin release should be considered. In fact, the cell growth/depletion in stored sludge is complex and not easy to predict. Therefore, the worst-case scenario should be considered during sludge handling.
- Due to the low efficacy of sludge oxidation as compared to only stored sludge, as well as the occurrence of cell growth, and gene expression regulation during oxidation/storage, oxidation cannot be a reliable approach in sludge treatment and management.
- Management of cyanobacteria and cyanotoxins in sludge should be initiated with the minimization of cyanobacteria and cyanotoxin accumulation throughout DWTPs.
- To control the negative impacts of cyanobacterial accumulation in DWTPs, recycling sludge supernatant to the head of the DWTPs should be regulated during cyanobacterial seasons. Suitable treatment and disposal approaches should be set into guidance and regulations for sludge-containing cyanotoxins.

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