



(11) **EP 3 601 157 B9**

(12) **CORRECTED EUROPEAN PATENT SPECIFICATION**

(15) Correction information:
Corrected version no 1 (W1 B1)
Corrections, see
Claims EN 3

(51) Int Cl.:
C01B 11/02 (2006.01)

(86) International application number:
PCT/US2018/023784

(48) Corrigendum issued on:
25.08.2021 Bulletin 2021/34

(87) International publication number:
WO 2018/175732 (27.09.2018 Gazette 2018/39)

(45) Date of publication and mention
of the grant of the patent:
26.05.2021 Bulletin 2021/21

(21) Application number: **18718019.5**

(22) Date of filing: **22.03.2018**

(54) **LOW RISK CHLORINE DIOXIDE ONSITE GENERATION SYSTEM**

SYSTEM ZUR ERZEUGUNG VOR ORT VON CHLORDIOXID MIT GERINGEM RISIKO

SYSTÈME DE GÉNÉRATION SUR SITE DE DIOXYDE DE CHLORE À FAIBLE RISQUE

(84) Designated Contracting States:
AL AT BE BG CH CY CZ DE DK EE ES FI FR GB
GR HR HU IE IS IT LI LT LU LV MC MK MT NL NO
PL PT RO RS SE SI SK SM TR

(30) Priority: **24.03.2017 US 201762476463 P**
13.11.2017 US 201762585316 P

(43) Date of publication of application:
05.02.2020 Bulletin 2020/06

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Description

BACKGROUND

5 1. Field of the Invention

[0001] The present disclosure generally relates to producing chlorine dioxide. More particularly, the disclosure relates to a chlorine dioxide reactor and methods of producing chlorine dioxide and treating aqueous systems.

10 2. Description of the Related Art

[0002] With the decline of gaseous chlorine as a microbiocide, various alternatives have been explored; including bleach, bleach with bromide, bromo-chloro dimethylhydantoin, ozone, and chlorine dioxide (ClO₂). Of these, chlorine dioxide has generated a great deal of interest for control of microbiological growth in a number of different industries, including the dairy industry, the beverage industry, the pulp and paper industry, the fruit and vegetable processing industries, various canning plants, the poultry industry, the beef processing industry and miscellaneous other food processing applications. Chlorine dioxide is also seeing increased use in municipal potable water treatment facilities and in industrial waste treatment facilities, because of its selectivity towards specific environmentally-objectionable waste materials, including phenols, sulfides, cyanides, thiosulfates, and mercaptans. In addition, chlorine dioxide is being used in the oil and gas industry for downhole applications as a well stimulation enhancement additive.

[0003] Unlike chlorine, chlorine dioxide remains a gas when dissolved in aqueous solutions and does not ionize to form weak acids. This property is at least partly responsible for the biocidal effectiveness of chlorine dioxide over a wide pH range. Moreover, chlorine dioxide is a highly effective microbiocide at concentrations as low as 0.1 parts per million (ppm) over a wide pH range.

[0004] The biocidal activity of chlorine dioxide is believed to be due to its ability to penetrate bacterial cell walls and react with essential amino acids within the cell cytoplasm to disrupt cell metabolism. This mechanism is more efficient than other oxidizers that "burn" on contact and is highly effective against *legionella*, algae and amoebal cysts, giardia cysts, coliforms, *salmonella*, *shigella*, and *cryptosporidium*.

[0005] Unfortunately, chlorine dioxide in solution is unstable with an extremely short shelf life and thus, is not commercially available. Chlorine dioxide solutions must typically be generated at its point of use such as, for example, by a reaction between a metal chlorate or metal chlorite in aqueous solution and a liquid phase strong acid. US 2005/244328 A1 and US 2010/155341 A1 disclose such processes for the production of chlorine dioxide. However, the use of liquid phase strong acids poses handling issues and safety concerns.

[0006] Production of chlorine dioxide using chlorate, hydrogen peroxide, and acid generates heat and may result in inadvertent explosive decomposition of chlorine dioxide if the temperature is not properly controlled.

BRIEF SUMMARY

[0007] A method of producing chlorine dioxide is disclosed. The method may include feeding a first solution comprising an acid through a first feed line into a mixing device; feeding a second solution comprising chlorite through a second feed line into the mixing device; mixing the first solution and the second solution in the mixing device to form a reaction mixture; feeding the reaction mixture into the reactor; reacting the acid and the chlorite in the reactor, wherein the reactor comprises a proximal portion in fluid communication with the mixing device and a distal portion in fluid communication with a motive water line; and mixing chlorine dioxide with motive water in the motive water line at the distal portion of the reactor. The mixing device is in fluid communication with a reactor, and the mixing device, the reactor, a portion of the first feed line, and a portion of the second feed line are positioned within the motive water line. The reactor comprises a coiled configuration having a coil plane angle of 1 degree to 60 degrees.

[0008] In some embodiments, the acid is hydrochloric acid, phosphoric acid, sulfuric acid, hypochlorous acid, or any combination thereof.

[0009] Another of producing chlorine dioxide is also disclosed. The method may include feeding a first solution comprising an acid through a first feed line into a mixing device; feeding a second solution comprising chlorate and hydrogen peroxide through a second feed line into the mixing device; mixing the first solution and the second solution in the mixing device to form a reaction mixture, wherein the mixing device is in fluid communication with a reactor; feeding the reaction mixture into the reactor; reacting the acid, the chlorate, and the hydrogen peroxide in the reactor, wherein the reactor comprises a proximal portion in fluid communication with the mixing device and a distal portion in fluid communication with a motive water line, wherein the mixing device, the reactor, a portion of the first feed line, and a portion of the second feed line are positioned within the motive water line; and mixing chlorine dioxide with motive water in the motive water line at the distal portion of the reactor. The reactor comprises a coiled configuration having a coil plane angle of 1 degree

to 60 degrees.

[0010] In some embodiments, the acid is sulfuric acid.

[0011] In other embodiments, a method of treating process water is disclosed. The method may include producing chlorine dioxide as described herein and injecting the chlorine dioxide and the motive water into the process water.

[0012] In some embodiments, the motive water line includes at least two longitudinal channels through which motive water may flow.

[0013] In some embodiments, the reaction mixture has a residence time in the reactor of at least about 0.1 minute, and a precursor velocity is at least about 25 cm/min, wherein the precursor velocity is a velocity of the first and second solutions.

[0014] In some embodiments, the methods described herein may include cooling the mixing device and the reactor with the motive water.

[0015] In some embodiments, the methods described herein may include producing the chlorine dioxide at a rate ranging from 0.0005 kg.h⁻¹ (0.001 lb/ hour) to 9 kg.h⁻¹ (20 lb/hour).

[0016] In some embodiments, the methods described herein may include operating the reactor at a temperature ranging from 2 °C to 80 °C.

[0017] In some embodiments, the method may include determining a chlorine dioxide demand in the process water using measurements selected from a chlorine dioxide sensor, an oxidation and reduction potential, a flow meter, a microbiological measurement, and any combination thereof.

[0018] In some embodiments, a cooling tower comprises the process water.

[0019] In other embodiments, a reactor for producing chlorine dioxide is disclosed. The reactor may include a mixing device; a first feed line in fluid communication with the mixing device; a second feed line in fluid communication with the mixing device; a reactor comprising a proximal portion in fluid communication with the mixing device and a distal portion in fluid communication with a motive water line; wherein the mixing device, the reactor, a portion of the first feed line, and a portion of the second feed line are positioned within the motive water line.

[0020] The reactor includes a coiled configuration having a coil plane angle of 1 degree to 60 degrees.

[0021] In some embodiments, the first feed line and the second feed line are opposite to and directed against each other.

[0022] In some embodiments, the reactor is operated at a pressure about equal to or greater than atmospheric pressure. Atmospheric pressure includes pressures ranging from about 101.3 kPa to about 33.7 kPa. In some embodiments, the reactor is operated at a pressure of about 101.3 kPa.

[0023] The foregoing has outlined rather broadly the features and technical advantages of the present disclosure in order that the detailed description that follows may be better understood. Additional features and advantages of the disclosure will be described hereinafter that form the subject of the claims of this application.

BRIEF DESCRIPTION OF THE SEVERAL VIEWS OF THE DRAWINGS

[0024] A detailed description of the invention is hereafter described with specific reference being made to the drawings in which:

FIG. 1 shows a conceptual diagram of an embodiment of a method of treating cooling tower water by generating chlorine dioxide onsite;

FIG. 2 shows an embodiment of a chlorine dioxide mixer and reactor;

FIG. 3 shows an embodiment of a chlorine dioxide mixer and reactor;

FIG. 4 shows a cross-sectional view of an embodiment of a chlorine dioxide mixer and reactor;

FIG. 5 shows an embodiment of a coiled reactor;

FIG. 6 shows a cross-sectional view of an embodiment of the motive water line and the reactor; and

FIG. 7 shows a conceptual diagram of an embodiment of a method of treating cooling tower water and other targets by generating chlorine dioxide onsite.

DETAILED DESCRIPTION

[0025] Various embodiments are described below with reference to the drawings in which like elements generally are referred to by like numerals. The relationship and functioning of the various elements of the embodiments may better be understood by reference to the following detailed description. However, embodiments are not limited to those illustrated in the drawings. It should be understood that the drawings are not necessarily to scale, and in certain instances details may have been omitted that are not necessary for an understanding of embodiments disclosed herein, such as - for example-conventional fabrication and assembly.

[0026] Traditionally, an eductor withdraws the chlorine dioxide solution from the reactor using reduced pressure. Reduced pressure is produced by feeding motive water through the eductor; however, the quality and quantity of the

motive water must be carefully controlled to avoid producing varying pressures. Varying pressures can lead to inefficiencies in chlorine dioxide production.

[0027] A method is disclosed for treating process water. The method may include feeding a first solution comprising an acid through a first feed line into a mixing device and feeding a second solution comprising chlorate and hydrogen peroxide through a second feed line into the mixing device. The first and second solutions are then mixed in the mixing device to form a reaction mixture. A reactor is in fluid communication with the mixing device. After mixing, the reaction mixture is fed into the reactor. The acid, chlorate, and hydrogen peroxide react in the reactor, thereby producing chlorine dioxide. The reactor includes a proximal portion in fluid communication with the mixing device and a distal portion in fluid communication with a motive water line. The mixing device, the reactor, a portion of the first feed line, and a portion of the second feed line are positioned within the motive water line. As the reaction mixture exits the reactor, the reaction mixture mixes with motive water in the motive water line at the distal portion of the reactor. The mixed chlorine dioxide and motive water may be injected into the process water.

[0028] In some embodiments, the first solution may consist of or comprise sulfuric acid and water, where the sulfuric acid concentration ranges from about 50% by weight to about 98% by weight. The sulfuric acid concentration in the first solution may be about 78%. In some embodiments, the second solution may comprise or consist of chlorate, hydrogen peroxide, and water. The second solution may comprise a concentration of chlorate that is in a range of about 25% by weight to about 60% by weight. The concentration of chlorate in the second solution may be about 40% by weight. The concentration of hydrogen peroxide in the second solution may be in the range of about 2% by weight to about 30% by weight. The concentration of hydrogen peroxide in the second solution may be about 8% by weight. In some embodiments, the acid may be sulfuric acid and the chlorate may be sodium chlorate.

[0029] In some embodiments, the first solution may consist of or comprise hydrochloric acid and water, where the hydrochloric acid concentration ranges from about 5% by weight to about 38% by weight. The hydrochloric acid concentration in the first solution may be about 37%. In some embodiments, the second solution may consist of or comprise a chlorite and water. The second solution may have a chlorite concentration that ranges from about 5% by weight to about 60% by weight. The chlorite concentration in the second solution may be about 25% by weight. In some embodiments, the chlorite may be sodium chlorite.

[0030] In certain embodiments, at least two feed lines carry precursor chemicals to the mixing device. The precursors may include the first and second solutions. In other embodiments, a third feed line may feed water or other chemicals to the mixing device. In some embodiments, the feed lines to the mixing device may consist of a first feed line and a second feed line.

[0031] In other embodiments, the method may further include the step of cooling the mixing device and the reactor with the motive water. The motive water passing through the motive water line may serve to control the temperature of the solutions in the feed lines and the temperature of the reaction mixture.

[0032] The reactor comprises a coiled configuration, which may further increase mixing and enhance mixing. The reactor having a coiled configuration includes a coil plane angle in a range from 1 to 60 degrees. In some embodiments, the coil plane angle may be from about 5 to about 30 degrees. The reactor having a coiled configuration may include a coil diameter. The coil diameter may be in a range from about 0.1 to about 16.0 inches, about 1 inch to about 16 inches, about 2 inch to about 16 inches, about 2 inch to about 10 inches, or about 2 inches to about 4 inches.

[0033] In some embodiments, the motive water line may comprise at least two channels. The channels may be longitudinal. In some embodiments, the motive water line may be a double containment pipe. The motive water line may have an inner channel and an outer channel through which motive water may flow, and the reactor may be positioned within the inner channel. The motive water flow rate in the outer channel may be different than the flow rate of the motive water in the inner channel. The motive water flow rate in the inner channel may be such that the temperature of the reaction mixture is controlled within a certain range. The reaction mixture temperature may be 2 °C to 80 °C, 2 °C to 70 °C, 35 °C to 70 °C, 40 °C to 70 °C, or 50 °C to 70 °C. The reaction mixture temperature may be 60 °C. The temperature of the motive water in the outer channel may be different from the temperature of the motive water in the inner channel. The outer channel may contain motive water having a flow rate sufficient to dilute the chlorine dioxide to a safe concentration such as less than about 3,000 ppm, less than about 2,000 ppm, less than about 1,500 ppm, less than about 1,000, less than about 750 ppm, less than about 500 ppm, less than about 250 ppm, less than about 100 ppm, or less than about 50 ppm.

[0034] In some embodiments, the first feed line and the second feed line may be opposite to and directed against each other. In this configuration, the solutions fed through the first and second feed lines may collide and mix.

[0035] In some embodiments, the reactor may include at least one contact zone. In some embodiments, the reactor may include at least two, three, four, or more contact zones. In other embodiments, the contact zone may comprise an inner diameter at least about two times larger than an inner diameter of the reactor. In some embodiments, the contact zone may comprise an inner diameter at least about three, four, or five times larger than an inner diameter of the reactor.

[0036] In some embodiments, the acid, the chlorate, and the hydrogen peroxide may form a reaction mixture in the mixing device. The first and second solutions combine to form the reaction mixture. At least one advantage of using a

solution of chlorate and hydrogen peroxide to react with an acid solution is that no chlorine gas is produced. The absence of chlorine gas provides a more safe and economical process.

[0037] In certain embodiments, the reaction mixture may have a residence time in the reactor of at least 0.1 minute. In some embodiments, the reaction mixture may have a residence time in the reactor of at least about 1 minute. In other
5 embodiments, the reaction mixture may reside in the reactor for at least about 3 minutes, at least about 4 minutes, at least about 5 minutes, at least about 6 minutes, at least about 7 minutes, at least about 8 minutes, at least about 9 minutes, or at least about 10 minutes. In some embodiments, the reaction mixture may have a residence time in the reactor of about 3.8 minutes. Residence time may be calculated by dividing total reactor volume by total precursor flow rate.

[0038] In some embodiments, the precursors (acid and chlorate/hydrogen peroxide or chlorite) may be fed into the
10 mixing device at a flow velocity of 25 cm/minute. In some embodiments, the precursors may be fed into the mixing device at a flow velocity in a range from about 20 cm/minute to about 200 cm/minute, about 20 cm/minute to about 65 cm/minute, or about 20 cm/minute to about 50 cm/minute. In some embodiments the precursor velocity may be 25 cm/minute, 50 cm/minute, or 65 cm/min. Precursor velocity can be calculated by dividing total precursor flow rate by the cross-sectional area of the reactor.

[0039] In other embodiments, the reactor may be operated at a pressure equal to or greater than atmospheric pressure.

[0040] In some embodiments, the method may include withdrawing chlorine dioxide using an eductor. In some emb-
odiments, the methods disclosed do not use an eductor to withdraw chlorine dioxide from the reactor. In some embod-
iments, the reactor does not comprise an eductor.

[0041] In some embodiments, the method may include operating the reactor at a temperature in a range from 2 °C to
20 80 °C. In some embodiments, the method may include operating the reactor at a temperature of 60 °C.

[0042] In some embodiments, the temperature of the motive water may be in a range from 2 °C to 80 °C. The motive
water temperature may be in range from about 15 °C to about 70 °C, about 30 °C to about 70 °C, about 40 °C to about
70 °C, about 40 °C to about 60 °C, or about 50 °C to about 70 °C.

[0043] In some embodiments, the method may include determining a chlorine dioxide demand in the process water
25 using measurements from a sensor, such as a chlorine dioxide sensor, an oxidation and reduction potential, a flow
meter, a microbiological measurement, or any combination thereof. In some embodiments, the chlorine dioxide may be
added to the process water in a cooling tower.

[0044] In some embodiments, the chlorine dioxide may be added to an aqueous system. The present disclosure is
not limited to adding chlorine dioxide to process water in cooling towers.

[0045] In some embodiments, the method may include a process control system including a Programmable Logic
30 Controller (PLC), a chlorine dioxide analyzer, a pressure transmitter (PT) and a flow transmitter (FT), controls the feed
pumps for the chemicals to the reactor and for the motive water.

[0046] In some embodiments, the method may include a distribution system. The distribution system may include at
least two independently controlled dosing points. The distribution system may include, for example, 3, 4, 5, 6, 7, or 8
35 independently controlled dosing points. When the different ClO₂ production set points are entered into the controller,
the PLC automatically calculates and adjusts the required production of the chlorine dioxide reactor. A distribution system
may allow chlorine dioxide solution to be distributed to multiple points directly from the discharge chlorine dioxide reactor,
thus avoiding a storage tank system. The distribution system may incorporate a ClO₂ solution pump and flowmeters and
control valves to manage the distribution among various dosing points.

[0047] The flow meters that may be used with the disclosed methods may be any suitable flow meter, such as but not
40 limited to clamp-on or pulse flow meters. Pulse-based flow meters detect the kinetic energy of flowing fluid using optical
or magnetic sensors and converting it into electrical energy in the form of digital pulses. The flow rate can be determined
by measuring the periods of the pulses.

[0048] The pumps used to feed the first and second solutions may be in communication with the PLC. The PLC may
45 also be in communication with any pumps placed on a product feed line.

[0049] In some embodiments, the reactor may be in fluid communication with at least one product feed line. Pumps
may be placed on each one of the product feed lines, or in some embodiments, multiple product feed lines may share
a single pump.

[0050] In some embodiments, any of the methods disclosed herein may include determining a flow rate of the first
50 solution in the first feed line, a flow rate of the second solution in the second feed line, a flow rate of a chlorine dioxide
solution in a product feed line, or a flow rate of the motive water in the motive water line.

[0051] In some embodiments, the flow rate of any solutions being fed into the reactor may be determined using at
least two flow meters that determine the flow rate based on different principles to ensure that an accurate amount of
precursor chemical is being fed into the reactor.

[0052] In some embodiments, any of the methods disclosed herein may include sensing a reactor pressure. In some
55 embodiments, the product feed lines may be equipped with a check valve, flow meter, and feed pump.

[0053] In certain embodiments, any of the methods disclosed herein may include calculating a chlorine dioxide dose.
The chlorine dioxide dose may be calculated using the flow rates of the, first solution, second solution, motive water,

chlorine dioxide solution in the product feed lines, and the chlorine dioxide concentration. For example, the chlorine dioxide concentration in the main line can be calculated using the formula: Main Line Product Rate = $(F_p + F_a + F_w - F_{\text{productline}}) \cdot C_{\text{ClO}_2}$ where F_p is the flow rate of hydrogen peroxide and chlorate, F_a is the flow rate of the acid, F_w is the flow rate of the motive water, $F_{\text{productline}}$ is the sum of flow rates in any product feed lines connected to the main line, and C_{ClO_2} is the concentration of chlorine dioxide.

[0054] Multiple alarms and failsafe settings may be set using the PLC. For example, pump issues may be detected by monitoring the differential between the flow meter readings and the pump rate calculations. Some failsafes may include pressure relief valves on the reactor or product feed lines. Any of the feed lines may have a pressure sensor that communicates readings to the main control device. By way of example, the main control device may include a PLC and an Adam AI/AO (analog input/analog output) module. If the reactor is housed in a cabinet, then the cabinet may have leak detection sensors and a cabinet door lock.

[0055] In other embodiments of the present disclosure, a method for producing chlorine dioxide is provided. The method includes feeding a first solution comprising an acid through a first feed line into a mixing device and feeding a second solution comprising chlorate and hydrogen peroxide through a second feed line into the mixing device. The method includes mixing the first solution and the second solution in the mixing device. The mixing device is connected to a reactor. After mixing, the first solution and the second solution are fed into the reactor where the acid, the chlorate, and the hydrogen peroxide react to form chlorine dioxide. The reactor includes a proximal portion in fluid communication with the mixing device and a distal portion in fluid communication with a motive water line. The mixing device, the reactor, a portion of the first feed line, and a portion of the second feed line are positioned within the motive water line. The method includes mixing the chlorine dioxide with motive water in the motive water line at the distal portion of the reactor. The reactor comprises a coiled configuration having a coil plane angle of 1 degree to 60 degrees.

[0056] In other embodiments, a method of producing chlorine dioxide is provided. The method includes feeding a first solution, such as an acid, through a first feed line into a mixing device. A second solution, such as chlorite, is fed through a second feed line into the mixing device. The mixing device is in fluid communication with a reactor. The method includes mixing the first solution and the second solution in the mixing device to form a reaction mixture. The reaction mixture is fed into the reactor where the acid and the chlorite react. The reactor includes a proximal portion in fluid communication with the mixing device and a distal portion in fluid communication with a motive water line. The mixing device, the reactor, a portion of the first feed line, and a portion of the second feed line are positioned within the motive water line. The method includes mixing chlorine dioxide with motive water in the motive water line at the distal portion of the reactor. The reactor comprises a coiled configuration having a coil plane angle of 1 degree to 60 degrees. In some embodiments, the acid may be hydrochloric acid, phosphoric acid, sulfuric acid, hypochlorous acid, or any combination thereof. In some embodiments, the acid may be hydrochloric acid.

[0057] A reactor for producing chlorine dioxide is provided. The reactor may include a mixing device, a first feed line in fluid communication with the mixing device, and a second feed line in fluid communication with the mixing device. The reactor may include a proximal portion in fluid communication with the mixing device and a distal portion in fluid communication with a motive water line. The mixing device, the reactor, a portion of the first feed line, and a portion of the second feed line may be positioned within the motive water line. The reactor comprises a coiled configuration having a coil plane angle of 1 degree to 60 degrees.

[0058] In some embodiments, the diameter of the motive water line may be larger than the diameter of the reactor and the first and second feed lines such that the reactor and the feed lines may be positioned in the lumen of the motive water line.

[0059] In some embodiments, the first feed line and the second feed line may be opposite to and directed against each other as in a "T" configuration, see, for example, the orientation of components corresponding to reference numerals 11 and 12 in Fig. 4.

[0060] In certain embodiments, the reactor may include at least one contact zone. The contact zone may include an inner diameter at least two times larger than an inner diameter of the reactor. As the reaction mixture flows through the reactor it may reach a contact zone where the reaction mixture may further mix. The outlet of the contact zone may narrow to the diameter of the reactor.

[0061] In some embodiments, the reactor may have a length in a range from about 25 inches to about 300 inches. In some embodiments, the reactor may have a length in a range from about 30 inches to about 90 inches, about 30 inches to about 80 inches, about 30 inches to about 70 inches, about 30 inches to about 60 inches, about 30 inches to about 50 inches, about 40 inches to about 90 inches, about 40 inches to about 80 inches, about 35 inches to about 45 inches, or about 35 inches to about 60 inches. In other embodiments, the reactor may be about 40 inches in length.

[0062] The reactor may be positioned horizontally, vertically, or any angle in between. In some embodiments, the reactor may be positioned vertically. In embodiments where the reactor is a coiled reactor, the reactor may spiral upward.

[0063] In some embodiments, the reactor may have an inner diameter in a range from about 0.1 inches to about 4 inches. In some embodiments, the reactor may have an inner diameter in a range from about 1 inch to about 1.25 inches, or about 0.25 inches to about 1.25 inches. In some embodiments, the reactor may have an inner diameter of about 0.25

inches, about 2 inches, about 3 inches, or about 4 inches. In other embodiments, the reactor may have an inner diameter of about 1.25 inches, about 1 inch, about 0.75 inches, about 0.375 inches, or about 0.1875 inches.

[0064] In some embodiments, the chlorine dioxide may be produced at a rate in a range from 0.0005 kg.h⁻¹ (0.001 lb/hour) to 9 kg.h⁻¹ (20 lb/hour). In some embodiments, the chlorine dioxide may be produced at a rate in a range from 0.0009 kg.h⁻¹ (0.002 lb/hour) to 6.8 kg.h⁻¹ (15 lb/hour). In some embodiments, the chlorine dioxide may be produced at a rate in a range from 2.3 kg.h⁻¹ (5 lb/hour) to 6.8 kg.h⁻¹ (15 lb/hour). As chlorine dioxide is produced at a higher rate, a larger diameter reactor may be used to ensure that the residence time in the reactor is at least 3 minutes. In some embodiments, the residence time in the reactor may be at least 2 minutes.

[0065] Referring to the figures, FIG. 1 shows a conceptual diagram of certain embodiments for a method of treating cooling tower water onsite. Precursor chemicals may be stored in a tank **10** onsite from which the chemicals may be fed through the first feed line **11** and the second feed line **12** into a chlorine dioxide mixer and reactor **15**. Water may be fed into the mixer to dilute the precursor chemicals being fed into the chlorine dioxide mixer and reactor **15**. Water may be fed into the mixer and reactor **15** through a motive feed line **14**. A water line **13** may supply water to the motive water line **14** and/or it may supply water to dilute acid before or after entering the mixer. Chlorine dioxide may be injected into process water **16** that may be fed into a cooling tower **19**. A pump **17** may feed the process water **16** through a heat exchanger **18**.

[0066] FIG. 2 and FIG. 3 show a closer view of some embodiments of the chlorine dioxide mixer and reactor **15**. The first feed line **11** and the second feed line **12** may introduce precursor chemicals into the mixing device **20**. In some embodiments, a water line **13** may feed water into the mixing device **20**. A motive feed line **14** may supply water to the motive water line **24**. In some embodiments, the water in the motive water line **24** may cool the reactor **21**. In other embodiments, the water in the motive water line **24** dilutes and mixes with the chlorine dioxide at the distal portion of the reactor **23**. The proximal portion of the reactor **22** may be connected to the mixing device **20**. Fluid in the mixing device **20** may flow out of the mixing device **20** and into the reactor **21**. In some embodiments, the reactor **21** may be in a coiled configuration. In some embodiments, a chlorine dioxide feed line **25** carries the chlorine dioxide mixed with motive water to the process water. Some embodiments may include at least one contact zone **30** (see FIG. 3, for example). The contact zone **30** may allow increased mixing of the reaction mixture or increased residence time in the reactor in order to increase reaction efficiency.

[0067] FIG. 4 shows an embodiment where the first feed line **11** and the second feed line **12** are opposite to and directed against each other. The solutions mix in the mixing device **20** and then enter the proximal portion of the reactor **22**. The reaction mixture may continue to flow through the reactor **21**, exit the distal portion of the reactor **23**, and mix with the motive water in the motive water line **24**. Water may be fed into the motive water line **24** from the motive feed line **14**. In some embodiments, the mixing device **20**, the reactor **21**, a portion of the first feed line **40**, and a portion of the second feed line **41** are positioned within the motive water line **24**.

[0068] FIG. 5 shows an embodiment where the reactor **21** may comprise a coiled configuration. The reactor having a coiled configuration may include a coil plane angle **50** ranging from about 5 to about 30 degrees. The reactor having a coiled configuration may include a coil diameter **51**. The coil diameter may range from about 0.1 inch to about 4.0 inches. The inner diameter of the coil reactor may be as described above for the reactor.

[0069] FIG. 6 show an embodiment where the motive water line comprises an outer channel **60** and an inner channel **61**. The reactor **21** may be disposed within the lumen of the inner channel **61**. In this configuration, the motive water may be fed into the motive water line via two separate channels. The outer channel **60** and inner channel **61** may carry motive water at difference flow rates. The motive water in the outer channel **60** may have a different temperature than the motive water in the inner channel **61**.

[0070] FIG. 7 shows a conceptual diagram of certain embodiments for a method of treating cooling tower water and multiple targets. Precursor chemicals may be stored in a tank **10** onsite from which the chemicals may be fed through the first feed line **11** and the second feed line **12** into a chlorine dioxide mixer and reactor **15**. Water may be fed into the mixer via line **13** to dilute the precursor chemicals being fed into the chlorine dioxide mixer and reactor **15**. Water may be fed into the mixer and reactor **15** through a motive feed line **14**. Chlorine dioxide may be injected into process water **16** that may be fed into a cooling tower **19**. A pump **17** may feed the process water **16** through a heat exchanger **18**. The chlorine dioxide generated in the reactor **15** can be added to process water **16** or diverted into a product feed line **70** for treating other targets. The product feed line **70** may include a valve **71**, a flow meter **72**, and a pump **73**. All valves and pumps may be controlled using a PLC (not pictured).

Examples

Example 1

[0071] A solution containing about 40% sodium chlorate and about 8% hydrogen peroxide and an aqueous solution of about 78% sulfuric acid were each fed at a rate of about 63 mL/h through 0.25 inch tubing into a 0.25 inch Kynar

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Union Tee that was inserted inside a polyvinyl chloride cross tube with about a 2 inch inner diameter.

[0072] The precursors flowing through the Kynar Union Tee then flowed through about 40 inches of 0.25 inch tubing (8 mL volume) that was inserted inside 0.5 inch tubing. Motive water flowed into the polyvinyl chloride cross tube and out of the 0.5 inch tubing (around the 0.25 inch tubing) to provide cooling and dilution. The reaction tube length and precursor velocity of about 26.5 cm/min provided a residence time of about 3.81 minutes.

[0073] Each experiment ran for about 15 minutes, and the chlorine dioxide concentration was measured about every 5 minutes. The motive water served two purposes: dilution and cooling. Water to precursor ratio of about 30 or less was required to establish the heat needed for the reaction. This generated chlorine dioxide concentrations of greater than about 5,500 mg/L.

[0074] Flow velocity of the precursors was set at about 26.5 cm/min. Flow velocities may be greater than about 25 cm/min to allow the gaseous oxygen to remain as nano- and micro-bubbles and to accelerate the movement of nano- and micro-bubbles out of the reactor tube.

[0075] The concentration of chlorine dioxide was determined using a spectrophotometer. Readings from the spectrophotometer were used to calculate the chlorine dioxide concentration using a calibration curve. The curve was prepared according to Iodometric Method 4500-ClO₂ in Standard Methods for the Examinations of Water and Wastewater 20th edition 1998.

[0076] The amount of solution consisting of about 40% sodium chlorate and about 7.99% hydrogen peroxide required for 100% reaction efficiency was about 2.86 mL to produce about 1 gram of chlorine dioxide. The reaction or conversion efficiency was calculated from the percentage ratio of the total chlorine dioxide generated to the theoretical chlorine dioxide production based on the amount of Purate consumed: $\eta_{\text{current}} = (C_{\text{ClO}_2} \cdot Q_{\text{ClO}_2}) / (Q_{\text{Purate}} / 2.86)$ where C_{ClO_2} is concentration of chlorine dioxide (g/L), Q_{ClO_2} flowrate of chlorine dioxide (L/hour), and Q_{Purate} flow rate of Purate (chlorate about 40% and hydrogen peroxide about 7.99% mixture) (mL/hour).

Table 1. Reaction efficiency

Motive Water (L/h)	7.50	7.50	7.50	3.76	3.76	3.76	3.14	3.14	3.14
Purate Flow Rate (mL/h)	63.0	63.0	63.0	63.0	63.0	63.0	63.0	63.0	63.0
78% Sulfuric Acid Flow Rate (mL/h)	63.0	63.0	63.0	63.0	63.0	63.0	63.0	63.0	63.0
Residence Time (min)	3.81	3.81	3.81	3.81	3.81	3.81	3.81	3.81	3.81
Water:Prec. Ratio	60	60	60	30	30	30	25	25	25
Precursor Flow Velocity (cm/m)	26.5	26.5	26.5	26.5	26.5	26.5	26.5	26.5	26.5
Chlorine Dioxide Conc. (g/L)	2.49	2.36	2.35	5.51	5.53	6.62	6.69	6.73	6.70
Chlorine Dioxide Flow Rate (L/h)	7.63	7.63	7.63	3.89	3.89	3.89	3.27	3.27	3.27
Actual Output (g/h)	18.7	17.7	17.6	21.4	21.5	21.8	21.8	22.0	21.9
Theoretical Output (g/h)	22.0	22.0	22.0	22.0	22.0	22.0	22.0	22.0	22.0
Reaction Efficiency (%)	84.9	80.3	79.9	97.2	97.6	99.1	99.1	99.8	99.3

Example 2

[0077] In this experiment, a 3/8" inch Tee connected to about 91.4 cm of 3/8" inch tubing was used as a mixing device and reactor, respectively. Different residence times (calculated by dividing reactor volume by total reactant flow rate) and motive water temperatures were tested. The conversion rate was calculated as described above. Purate was a mixture of chlorate and hydrogen peroxide as described above.

Table 2. Residence time and temperature effects on conversion.

Residence Time	Motive Water Temp	Purate	H ₂ SO ₄	ClO ₂	H ₂ SO ₄ /Purate	Conversion Rate
min	°C	mole/min	mole/min	lb/hr	mole/mole	%
15.6	34	0.0095	0.0250	0.085	2.64	88.4
15.6	37	0.0095	0.0250	0.085	2.64	93.4
8.6	36	0.0172	0.0455	0.154	2.64	87.1
8.6	43	0.0172	0.0455	0.154	2.64	88.1
8.6	53	0.0172	0.0455	0.154	2.64	99.7
8.6	51	0.0172	0.0366	0.154	2.13	86.2
15.6	50	0.0095	0.0250	0.085	2.64	95.9
7.8	51	0.0189	0.0500	0.169	2.64	99.9
3.9	50	0.0379	0.1001	0.338	2.64	89.0
5.2	48	0.0284	0.0751	0.254	2.64	96.0

Example 3

[0078] A 1/4" inch Tee connected to a 12 ml chamber by about 5.72 cm of 1/4" inch tubing was used in this experiment. The chamber was also connected to about 26 cm of 1/4" tubing. The configuration was similar to the reactor depicted in FIG. 3 where the 12 ml chamber may be a contact zone.

[0079] Table 3 shows the effects of motive water and the ratio of H₂SO₄/Purate on conversion rate. Table 4 shows the effects of reduced amount of acid at a motive water temperature of about 62 °C and a residence time of about 14 min. The residence time for these trials was fixed at about 14 min.

Table 3. Effects of motive water temperature and H₂SO₄/Purate on conversion

Temp °C	Purate mole/min	H ₂ SO ₄ mole/min	ClO ₂ lb/hr	H ₂ SO ₄ /Purate mole/mole	Conversion Rate %
16	0.0095	0.0250	0.085	2.64	64.8
30	0.0095	0.0250	0.085	2.64	92.6
41	0.0095	0.0250	0.085	2.64	95.1
53	0.0095	0.0250	0.085	2.64	94.4
62	0.0095	0.0250	0.085	2.64	97.7
41	0.0095	0.0205	0.085	2.16	87.8
47	0.0095	0.0208	0.085	2.20	95.4
62	0.0095	0.0208	0.085	2.20	97.4

Table 4. Effect of reduced acid at 62 °C.

Temp °C	Purate mole/min	H ₂ SO ₄ mole/min	ClO ₂ lb/hr	H ₂ SO ₄ /Purate mole/mole	Conversion Rate %
62	0.0095	0.0254	0.085	2.68	97.1
	0.0095	0.0208	0.085	2.20	99.8
	0.0095	0.0162	0.085	1.71	92.7
	0.0095	0.0127	0.085	1.34	65.9

[0080] Table 5 shows the effects of residence time and motive water temperature on conversion rate.

Table 5. Effect of residence time and temperature on conversion rate.

Residence Time	Temp	Purate	H ₂ SO ₄	ClO ₂	H ₂ SO ₄ /Purate	Conversion Rate
min	°C	mole/min	mole/min	lb/hr	mole/mole	%
3	53	0.0095	0.0159	0.085	1.68	52.7
14	47	0.0095	0.0162	0.085	1.71	78.9
14	62	0.0095	0.0162	0.085	1.71	92.7
17	45	0.0095	0.0159	0.085	1.68	71.7
17	46	0.0095	0.0162	0.085	1.71	83.7
17	46	0.0189	0.0324	0.170	1.71	68.5
17	62	0.0095	0.0162	0.085	1.71	92.1

Example 4

[0081] In this experiment, a 1/4" inch Tee connected to a 12 ml chamber by about 2 feet of 1/4" inch tubing was used. The chamber was also connected to about 10 feet of 1/4" tubing. The configuration was similar to the reactor depicted in FIG. 3 where the 12 ml chamber may be a contact zone. The volume of the reactor was about 78.2 ml.

Table 6. Effect of residence time, motive water temperature, and acid on conversion rate.

Residence Time	Temp	Purate	H ₂ SO ₄	ClO ₂	H ₂ SO ₄ /Purate	Conversion Rate
min	°C	mole/min	mole/min	lb/hr	mole/mole	%
21.1	31.2	0.0095	0.0250	0.085	2.64	95.0
21.1	29.3	0.0095	0.0205	0.085	2.16	90.0
21.1	35.6	0.0095	0.0205	0.085	2.16	88.5
21.1	37.5	0.0095	0.0205	0.085	2.16	93.0
21.1	44.1	0.0095	0.0205	0.085	2.16	99.1
10.6	43.9	0.0189	0.0409	0.169	2.16	87.6
10.6	43.5	0.0189	0.0455	0.169	2.40	89.3
11.6	43.3	0.0172	0.0455	0.154	2.64	97.1
11.6	33.8	0.0172	0.0455	0.154	2.64	85.7

Example 5

[0082] Chlorine dioxide production was tested using the reactor disclosed herein and chlorite as a precursor instead of chlorate. A solution of about 25 wt% of chlorite and a solution of about 29 wt% hydrochloric acid was fed to a reactor. The reactor included a 1/4" inch Tee (mixing device) connected to a 12 ml chamber by about 5.72 cm of 1/4" inch tubing. The chamber was also connected to about 26 cm of 1/4" tubing. Table 7 shows the reaction efficiency as a function of time.

Table 7. Chlorine dioxide production using chlorite and hydrochloric acid.

Time	ClO ₂	ClO ₂ Vol.	ClO ₂ Output	NaClO ₂ Rate	mL NaClO ₂ /g	Rxn. Eff.
(min)	(mg/L)	(L/h)	(g/h)	(L/h)	ClO ₂	(%)
0	162	7.59	1.23	0.013	10.25	54.5
2	178	7.59	1.35	0.013	9.33	59.9
4	202	7.59	1.53	0.013	8.22	68.0
6	214	7.59	1.62	0.013	7.76	72.1
8	225	7.59	1.71	0.013	7.38	75.8
10	227	7.59	1.72	0.013	7.31	76.4
12	229	7.59	1.74	0.013	7.25	77.1
14	264	7.59	2.00	0.013	6.29	88.9
16	280	7.59	2.12	0.013	5.93	94.3
18	282	7.59	2.14	0.013	5.89	94.9
20	289	7.59	2.19	0.013	5.75	97.3

(continued)

Time (min)	ClO ₂ (mg/L)	ClO ₂ Vol. (L/h)	ClO ₂ Output (g/h)	NaClO ₂ Rate (L/h)	mL NaClO ₂ /g ClO ₂	Rxn. Eff. (%)
22	291	7.59	2.21	0.013	5.71	98.0
24	289	7.59	2.19	0.013	5.75	97.3
26	291	7.59	2.21	0.013	5.71	98.0
28	292	7.59	2.22	0.013	5.69	98.3
30	290	7.59	2.20	0.013	5.73	97.6

[0083] Chlorine dioxide production using chlorite and hydrochloric acid was tested in a larger scale reactor. The reactor included a 3/8" inch Tee connected to about 91.4 cm of 3/8" inch tubing. Table 8 compares the theoretical production rate with the actual production rate.

Table 8. Chlorine dioxide conversion efficiency in large scale reactor.

Theoretical ClO ₂ Production Rate (lb/h)	Average Motive Water Flow Rate (gpm)	Average ClO ₂ Concentration in Motive Water (ppm)	Actual ClO ₂ Production Rate (lb/hr)	Average Conversion Efficiency (%)	Standard Deviation (%)
0.2	1.48	258.8	0.192	96.1	21.0
2.0	4.87	807.8	1.970	98.4	12.0
4.0	6.00	1336.8	4.019	100.4	5.1

Claims

1. A method of producing chlorine dioxide, comprising:

feeding a first solution comprising an acid through a first feed line (11) into a mixing device (20);
 feeding a second solution comprising chlorite through a second feed (12) line into the mixing device (20);
 mixing the first solution and the second solution in the mixing device (20) to form a reaction mixture, wherein the mixing device is in fluid communication with a reactor (21);
 feeding the reaction mixture into the reactor (21);
 reacting the acid and the chlorite in the reactor (21), wherein the reactor comprises a proximal portion (22) in fluid communication with the mixing device (20) and a distal portion (23) in fluid communication with a motive water line (24), wherein the mixing device (20), the reactor (21), a portion of the first feed line (11), and a portion of the second feed line (12) are positioned within the motive water line (24); and
 mixing chlorine dioxide with motive water in the motive water line (24) at the distal portion of the reactor (21), wherein the reactor (21) comprises a coiled configuration having a coil plane angle of 1 degree to 60 degrees.

2. The method of claim 1, wherein the acid is hydrochloric acid, phosphoric acid, sulfuric acid, hypochlorous acid, or any combination thereof.

3. A method of producing chlorine dioxide, comprising:

feeding a first solution comprising an acid through a first feed line (11) into a mixing device (20);
 feeding a second solution comprising chlorate and hydrogen peroxide through a second feed line (12) into the mixing device (20);
 mixing the first solution and the second solution in the mixing device (20) to form a reaction mixture, wherein the mixing device is in fluid communication with a reactor (21);
 feeding the reaction mixture into the reactor (21);
 reacting the acid, the chlorate, and the hydrogen peroxide in the reactor (21), wherein the reactor comprises a proximal portion (22) in fluid communication with the mixing device and a distal portion (23) in fluid communication with a motive water line (24), wherein the mixing device (20), the reactor (21), a portion of the first feed line (11), and a portion of the second feed line (12) are positioned within the motive water line (24), and

mixing chlorine dioxide with motive water in the motive water line (24) at the distal portion (23) of the reactor, wherein the reactor (21) comprises a coiled configuration having a coil plane angle of 1 degree to 60 degrees.

4. The method of claim 3, wherein the acid is sulfuric acid.

5. A method of treating process water, comprising: producing chlorine dioxide according to any one of claims 1-4; and injecting the chlorine dioxide and the motive water into the process water.

6. The method of any one of claims 1-5, further comprising cooling the mixing device and the reactor with the motive water.

7. The method of any one of claims 1-6, wherein the first feed line and the second feed line are opposite to and directed against each other.

8. The method of any one of claims 1-7, wherein the motive water line comprises at least two longitudinal channels through which motive water may flow, and/or wherein the reactor is operated at a pressure equal to or greater than atmospheric pressure.

9. The method of any one of claims 1-8, wherein the reaction mixture has a residence time in the reactor of at least 0.1 minute, and a precursor velocity is at least 25 cm/min, wherein the precursor velocity is a velocity of the first and second solutions.

10. The method of any one of claims 1-9, further comprising operating the reactor at a temperature ranging from 2 °C to 80 °C.

11. The method of any of claims 5-10, wherein a cooling tower comprises the process water.

12. A reactor for producing chlorine dioxide, comprising:

a mixing device (20);

a motive water line (24);

a first feed line (11) in fluid communication with the mixing device;

a second feed line (12) in fluid communication with the mixing device; and

a reactor (21) comprising a proximal portion (22) in fluid communication with the mixing device and a distal portion (23) in fluid communication with a motive water line;

wherein the mixing device, the reactor, a portion of the first feed line, and a portion of the second feed line are positioned within the motive water line and

wherein the reactor comprises a coiled configuration having a coil plane angle of 1 degree to 60 degrees.

13. The reactor of claim 12, wherein the first feed line and the second feed line are opposite to and directed against each other.

Patentansprüche

1. Verfahren zum Erzeugen von Chlordioxid, das Folgendes umfasst:

Zuführen einer ersten Lösung, die eine Säure umfasst, durch eine erste Zuführungsleitung (11) in eine Mischvorrichtung (20);

Zuführen einer zweiten Lösung, die Chlorit umfasst, durch eine zweite Zuführungsleitung (12) in die Mischvorrichtung (20);

Mischen der ersten Lösung und der zweiten Lösung in der Mischvorrichtung (20), um ein Reaktionsgemisch auszubilden, wobei die Mischvorrichtung in Fluidverbindung mit einem Reaktor (21) steht;

Zuführen des Reaktionsgemisches in den Reaktor (21);

Umsetzen der Säure und des Chlorits in dem Reaktor (21), wobei der Reaktor einen proximalen Abschnitt (22) in Fluidverbindung mit der Mischvorrichtung (20) und einen distalen Abschnitt (23) in Fluidverbindung mit einer Treibwasserleitung (24) umfasst,

wobei die Mischvorrichtung (20), der Reaktor (21), ein Abschnitt der ersten Zuführungsleitung (11) und ein

Abschnitt der zweiten Zuführungsleitung (12) innerhalb der Treibwasserleitung (24) positioniert sind; und Mischen von Chlordioxid mit Treibwasser in der Treibwasserleitung (24) an dem distalen Abschnitt des Reaktors (21), wobei der Reaktor (21) eine gespulte Konfiguration umfasst, die einen Winkel einer Spulenebene von 1 Grad bis 60 Grad aufweist.

2. Verfahren nach Anspruch 1, wobei die Säure Chlorwasserstoffsäure, Phosphorsäure, Schwefelsäure, Hypochlorsäure oder eine beliebige Kombination davon ist.

3. Verfahren zum Erzeugen von Chlordioxid, das Folgendes umfasst:

Zuführen einer ersten Lösung, die eine Säure umfasst, durch eine erste Zuführungsleitung (11) in eine Mischvorrichtung (20);

Zuführen einer zweiten Lösung, die Chlorat und Wasserstoffperoxid umfasst, durch eine zweite Zuführungsleitung (12) in die Mischvorrichtung (20);

Mischen der ersten Lösung und der zweiten Lösung in der Mischvorrichtung (20), um ein Reaktionsgemisch auszubilden, wobei die Mischvorrichtung in Fluidverbindung mit einem Reaktor (21) steht;

Zuführen des Reaktionsgemisches in den Reaktor (21);

Umsetzen der Säure, des Chlorats und des Wasserstoffperoxids in dem Reaktor (21),

wobei der Reaktor einen proximalen Abschnitt (22) in Fluidverbindung mit der Mischvorrichtung und einen distalen Abschnitt (23) in Fluidverbindung mit einer Treibwasserleitung (24) umfasst, wobei die Mischvorrichtung (20), der Reaktor (21), ein Abschnitt der ersten Zuführungsleitung (11) und ein Abschnitt der zweiten Zuführungsleitung (12) innerhalb der Treibwasserleitung (24) positioniert sind, und

Mischen von Chlordioxid mit Treibwasser in der Treibwasserleitung (24) an dem distalen Abschnitt (23) des Reaktors, wobei der Reaktor (21) eine gespulte Konfiguration umfasst, die einen Winkel der Spulenebene von 1 Grad bis 60 Grad aufweist.

4. Verfahren nach Anspruch 3, wobei die Säure Schwefelsäure ist.

5. Verfahren zum Behandeln von Betriebswasser, das Folgendes umfasst: Erzeugen von Chlordioxid nach einem der Ansprüche 1-4; und Injizieren des Chlordioxids und des Treibwassers in das Betriebswasser.

6. Verfahren nach einem der Ansprüche 1-5, das ferner ein Kühlen der Mischvorrichtung und des Reaktors mit dem Treibwasser umfasst.

7. Verfahren nach einem der Ansprüche 1-6, wobei die erste Zuführungsleitung und die zweite Zuführungsleitung sich gegenüber liegen und gegeneinander gerichtet sind.

8. Verfahren nach einem der Ansprüche 1-7, wobei die Treibwasserleitung wenigstens zwei Längskanäle umfasst, durch die Treibwasser fließen kann, und/oder wobei der Reaktor bei einem Druck betrieben wird, der gleich oder größer als Luftdruck ist.

9. Verfahren nach einem der Ansprüche 1-8, wobei das Reaktionsgemisch eine Verweilzeit in dem Reaktor von wenigstens 0,1 Minuten aufweist und eine Vorläufergeschwindigkeit wenigstens 25 cm/min beträgt, wobei die Vorläufergeschwindigkeit eine Geschwindigkeit der ersten und der zweiten Lösung ist.

10. Verfahren nach einem der Ansprüche 1-9, das ferner das Betreiben des Reaktors bei einer Temperatur umfasst, die von 2 °C bis 80 °C reicht.

11. Verfahren nach einem der Ansprüche 5-10, wobei ein Kühlturm das Betriebswasser umfasst.

12. Reaktor zum Erzeugen von Chlordioxid, der Folgendes umfasst:

eine Mischvorrichtung (20);

eine Treibwasserleitung (24);

eine erste Zuführungsleitung (11) in Fluidverbindung mit der Mischvorrichtung;

eine zweite Zuführungsleitung (12) in Fluidverbindung mit der Mischvorrichtung; und

einen Reaktor (21), der einen proximalen Abschnitt (22) in Fluidverbindung mit der Mischvorrichtung und einen distalen Abschnitt (23) in Fluidverbindung mit einer Treibwasserleitung umfasst;

wobei die Mischvorrichtung, der Reaktor, ein Abschnitt der ersten Zuführungsleitung und ein Abschnitt der zweiten Zuführungsleitung innerhalb der Treibwasserleitung positioniert sind und
wobei der Reaktor eine gespulte Konfiguration umfasst, die einen Winkel der Spulenebene von 1 Grad bis 60 Grad aufweist.

13. Reaktor nach Anspruch 12, wobei die erste Zuführungsleitung und die zweite Zuführungsleitung sich gegenüber liegen und gegeneinander gerichtet sind.

Revendications

1. Procédé de production de dioxyde de chlore, comprenant :

l'introduction d'une première solution comprenant un acide à travers une première conduite d'alimentation (11) dans un dispositif de mélange (20) ;
l'introduction d'une seconde solution comprenant du chlorite à travers une seconde conduite d'alimentation (12) dans le dispositif de mélange (20) ;
le mélange de la première solution et de la seconde solution dans le dispositif de mélange (20) pour former un mélange réactionnel, le dispositif de mélange étant en communication fluïdique avec un réacteur (21) ;
l'introduction du mélange réactionnel dans le réacteur (21) ;
la mise en réaction de l'acide et du chlorite dans le réacteur (21), le réacteur comprenant une partie proximale (22) en communication fluïdique avec le dispositif de mélange (20) et une partie distale (23) en communication fluïdique avec une conduite d'eau motrice (24), le dispositif de mélange (20), le réacteur (21), une partie de la première conduite d'alimentation (11) et une partie de la seconde conduite d'alimentation (12) étant positionnés à l'intérieur de la conduite d'eau motrice (24) ; et
le mélange du dioxyde de chlore avec de l'eau motrice dans la conduite d'eau motrice (24) au niveau de la partie distale du réacteur (21),
le réacteur (21) comprenant une configuration enroulée ayant un angle de plan de bobine de 1 degré à 60 degrés.

2. Procédé selon la revendication 1, dans lequel l'acide est l'acide chlorhydrique, l'acide phosphorique, l'acide sulfurique, l'acide hypochloreux ou toute combinaison de ceux-ci.

3. Procédé de production de dioxyde de chlore, comprenant :

l'introduction d'une première solution comprenant un acide à travers une première conduite d'alimentation (11) dans un dispositif de mélange (20) ;
l'introduction d'une seconde solution comprenant du chlorate et du peroxyde d'hydrogène à travers une seconde conduite d'alimentation (12) dans le dispositif de mélange (20) ;
le mélange de la première solution et de la seconde solution dans le dispositif de mélange (20) pour former un mélange réactionnel, le dispositif de mélange étant en communication fluïdique avec un réacteur (21) ;
l'introduction du mélange réactionnel dans le réacteur (21) ;
la mise en réaction de l'acide, du chlorate et du peroxyde d'hydrogène dans le réacteur (21), le réacteur comprenant une partie proximale (22) en communication fluïdique avec le dispositif de mélange et une partie distale (23) en communication fluïdique avec une conduite d'eau motrice (24), le dispositif de mélange (20), le réacteur (21), une partie de la première conduite d'alimentation (11) et une partie de la seconde conduite d'alimentation (12) étant positionnés dans la conduite d'eau motrice (24), et le mélange du dioxyde de chlore avec de l'eau motrice dans la conduite d'eau motrice (24) au niveau de la partie distale (23) du réacteur, le réacteur (21) comprenant une configuration enroulée ayant un angle de plan de bobine de 1 degré à 60 degrés.

4. Procédé selon la revendication 3, dans lequel l'acide est l'acide sulfurique.

5. Procédé de traitement d'eau de process, comprenant : la production de dioxyde de chlore selon l'une quelconque des revendications 1 à 4 ; et l'injection du dioxyde de chlore et de l'eau motrice dans l'eau de process.

6. Procédé selon l'une quelconque des revendications 1 à 5, comprenant en outre le refroidissement du dispositif de mélange et du réacteur avec l'eau motrice.

7. Procédé selon l'une quelconque des revendications 1 à 6, dans lequel la première conduite d'alimentation et la

seconde conduite d'alimentation sont opposées et dirigées l'une contre l'autre.

8. Procédé selon l'une quelconque des revendications 1 à 7, dans lequel la conduite d'eau motrice comprend au moins deux canaux longitudinaux à travers lesquels l'eau motrice peut s'écouler, et/ou dans lequel le réacteur fonctionne à une pression égale ou supérieure à la pression atmosphérique.

9. Procédé selon l'une quelconque des revendications 1 à 8, dans lequel le mélange réactionnel a un temps de séjour dans le réacteur d'au moins 0,1 minute et une vitesse de précurseur est d'au moins 25 cm/min, la vitesse du précurseur étant une vitesse des première et seconde solutions.

10. Procédé selon l'une quelconque des revendications 1 à 9, comprenant en outre le fonctionnement du réacteur à une température comprise entre 2 °C et 80 °C.

11. Procédé selon l'une quelconque des revendications 5 à 10, dans lequel une tour de refroidissement comprend l'eau de process.

12. Réacteur de production de dioxyde de chlore, comprenant :

un dispositif de mélange (20) ;

une conduite d'eau motrice (24) ;

une première conduite d'alimentation (11) en communication fluide avec le dispositif de mélange ;

une seconde conduite d'alimentation (12) en communication fluide avec le dispositif de mélange ; et

un réacteur (21) comprenant une partie proximale (22) en communication fluide avec le dispositif de mélange et une partie distale (23) en communication fluide avec une conduite d'eau motrice ;

dans lequel le dispositif de mélange, le réacteur, une partie de la première conduite d'alimentation et une partie de la seconde conduite d'alimentation sont positionnés dans la conduite d'eau motrice et

dans lequel le réacteur comprend une configuration enroulée ayant un angle de plan de bobine de 1 degré à 60 degrés.

13. Réacteur selon la revendication 12, dans lequel la première conduite d'alimentation et la seconde conduite d'alimentation sont opposées et dirigées l'une contre l'autre.

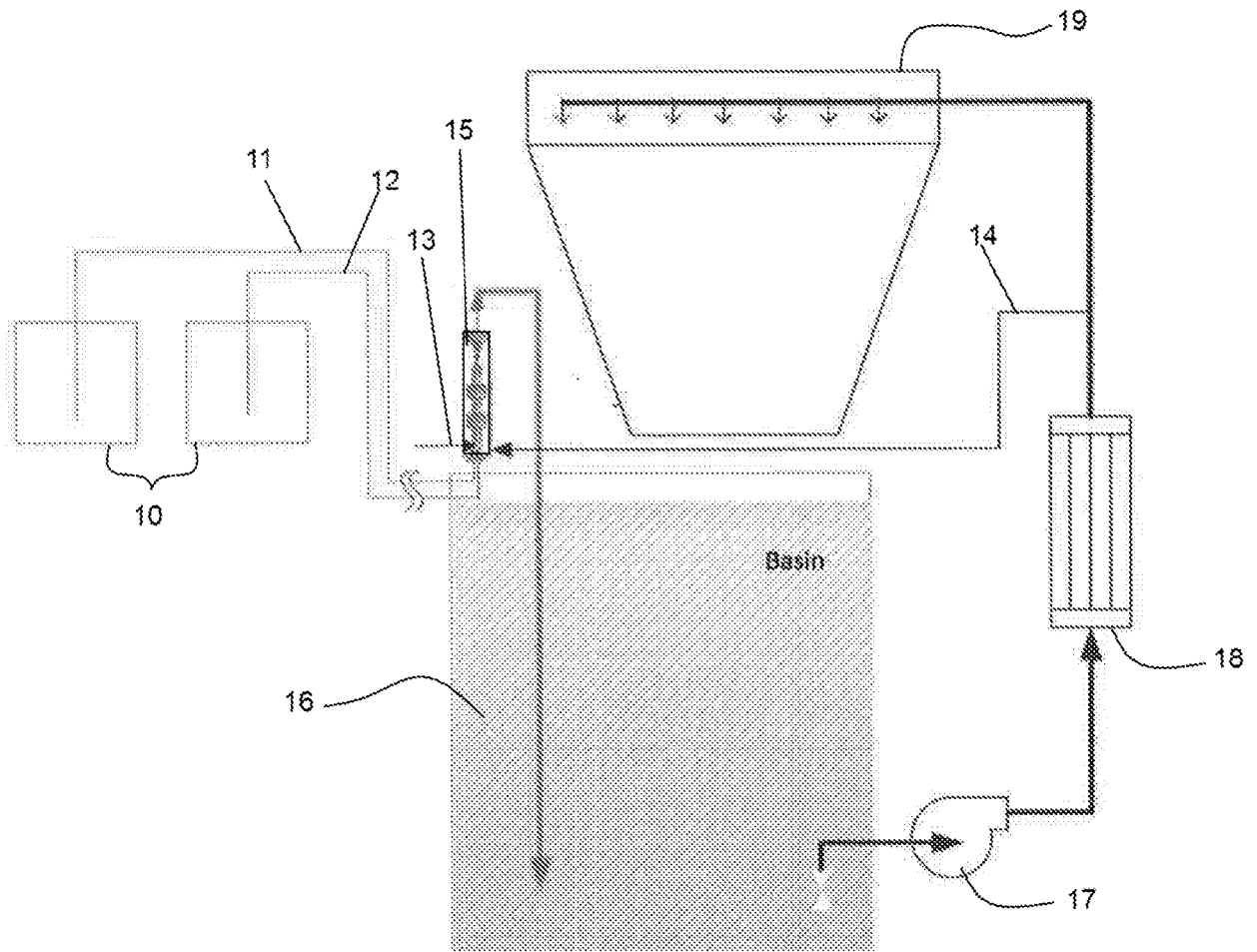


FIG. 1

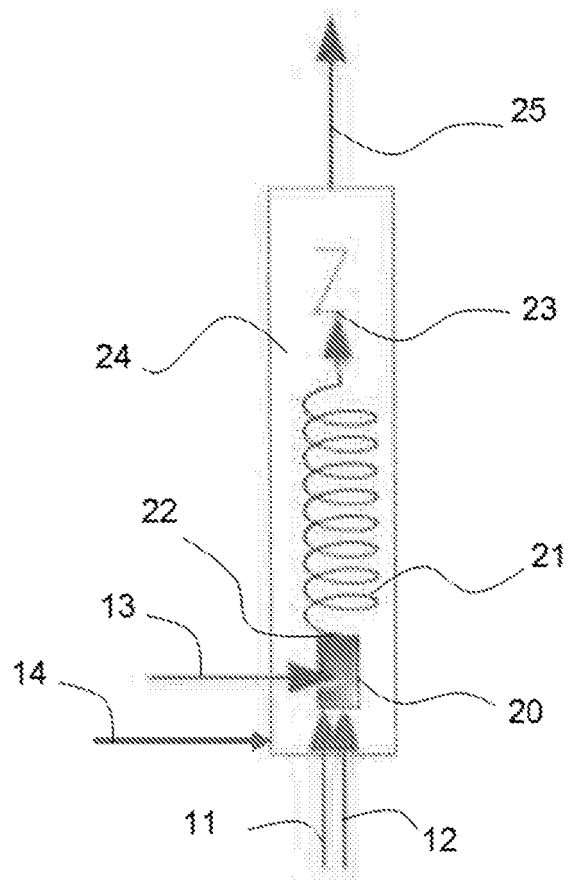


FIG. 2

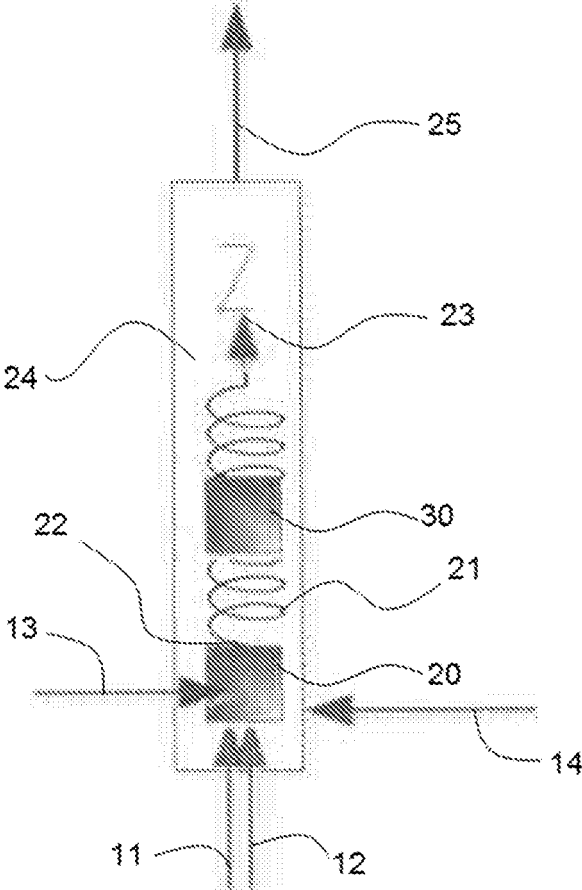


FIG. 3

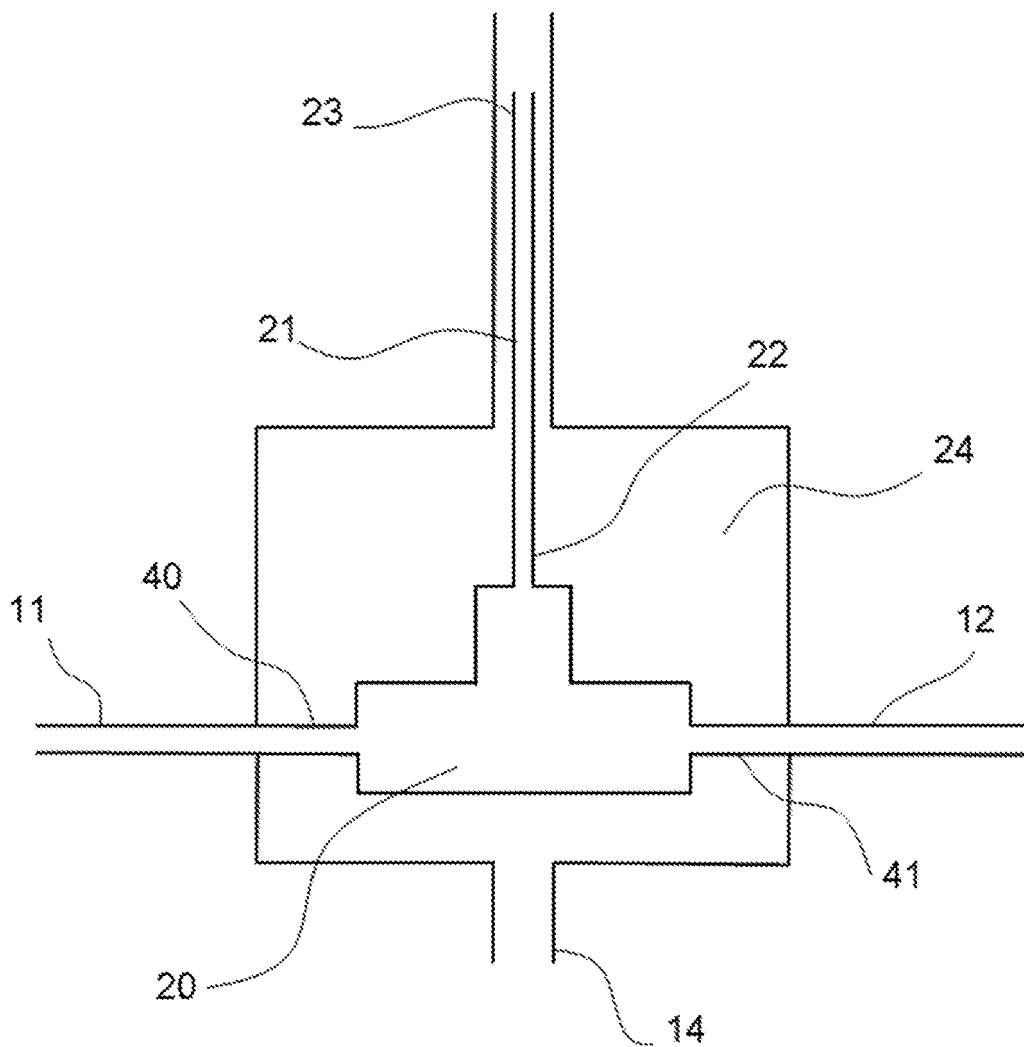


FIG. 4

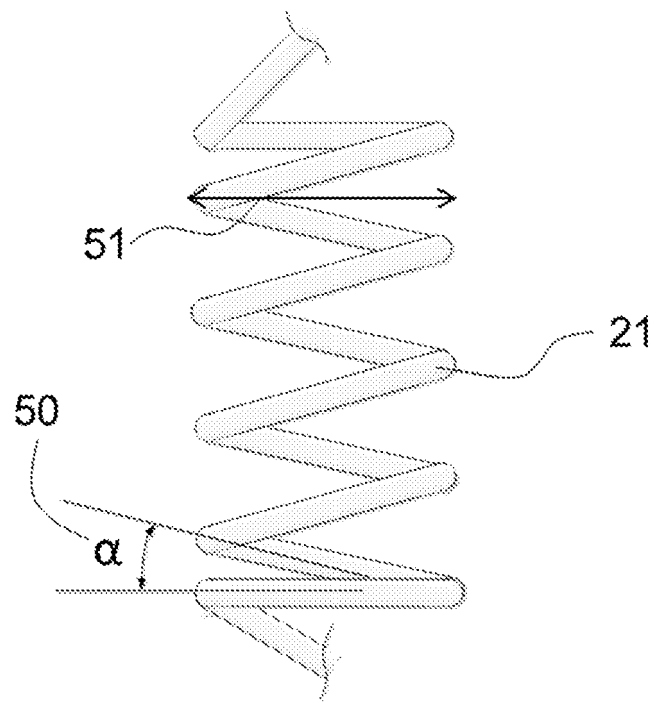


FIG. 5

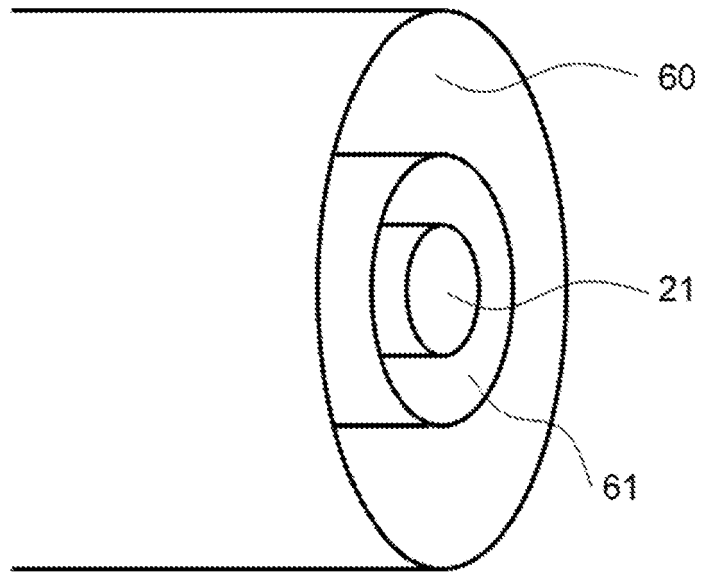


FIG. 6

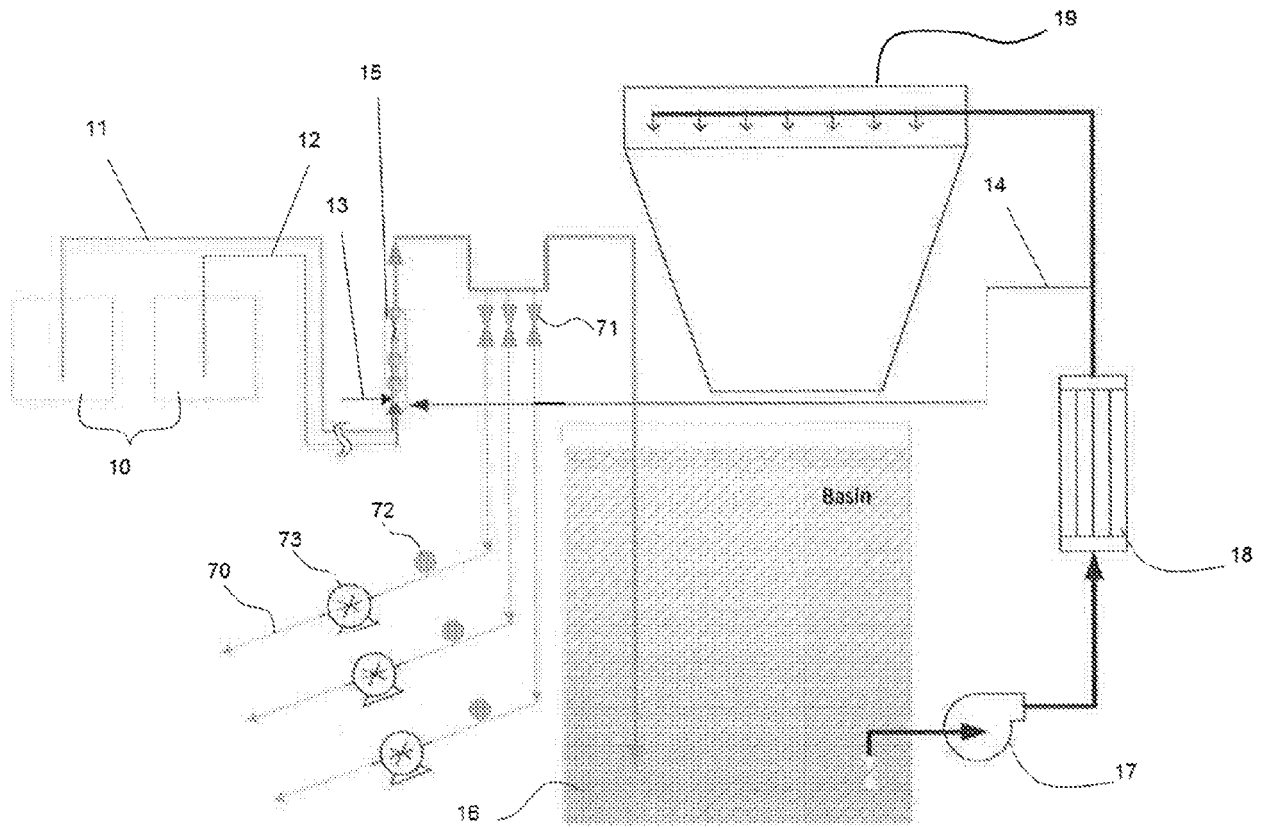


FIG. 7

REFERENCES CITED IN THE DESCRIPTION

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