<table>
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<th>(45) Date of publication and mention of the grant of the patent:</th>
<th>23.07.2003 Bulletin 2003/30</th>
</tr>
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<tbody>
<tr>
<td>(21) Application number:</td>
<td>98106792.9</td>
</tr>
<tr>
<td>(22) Date of filing:</td>
<td>15.04.1998</td>
</tr>
</tbody>
</table>
| (54) **Flue gas treating process**  
Verfahren zur Abgasbehandlung  
Procédé pour le traitement de gaz de fumée | |
| (84) Designated Contracting States: | DE DK ES IT |
| (43) Date of publication of application: | 28.10.1998 Bulletin 1998/44 |
| (60) Divisional application: | 01124366.4 / 1 182 400 |
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| DE-A- 2 752 288 | DE-A- 3 636 554 |

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FIELD OF THE INVENTION AND RELATED ART STATEMENT

[0001] This invention relates to a flue gas treating technique for effecting at least the denitration and desulfurization of flue gas. More particularly, it relates to a flue gas treating technique which makes it possible to reduce the size of the equipment and enhance the performance of the equipment.

[0002] Conventionally, in order to remove nitrogen oxides, sulfur oxides (typically sulfur dioxide) and dust (e.g., fly ash) present in flue gas discharged from a boiler of a thermal electric power plant or the like, an exemplary flue gas treating system or process as illustrated in FIGs. 9 and 10 is widely employed. This flue gas treating technique is described hereinafter.

[0003] As illustrated in FIG. 9, untreated flue gas A discharged from a boiler (not shown in FIG. 9) is first introduced into a denitrator 2 installed in a boiler house 1, so that nitrogen oxides present in the flue gas are decomposed. Denitrator 2 functions to decompose nitrogen oxides according to the catalytic ammonia reduction method using a catalyst. In this step, it has been conventional practice to inject ammonia B into the flue gas in an amount almost equal to the equivalent amount required for denitration. The amount of ammonia slipping to the downstream side of denitrator 2 is as slight as about 5 ppm.

[0004] Then, the flue gas is introduced into an air heater (or heat exchanger) 3 also installed in boiler house 1. Thus, heat is recovered from the flue gas and used to heat air C supplied to the boiler. Conventionally, a heat exchanger of the so-called Ljungström type has been used as air heater 3.

[0005] Then, the flue gas leaving this air heater 3 is subsequently conducted out of boiler house 1 by a flue 4 and introduced into a dry electrostatic precipitator 5 installed outside boiler house 1. In this electrostatic precipitator 5, dust present in the flue gas is captured and removed.

[0006] In the case of an oil-fired boiler, ammonia may also be injected into the flue gas in flue 4 so that sulfur trioxide (SO₃) present in the flue gas may be captured as ammonium sulfate [(NH₄)₂SO₄] in electrostatic precipitator 5. On the other hand, in the case of a coal-fired boiler, a large amount of dust such as fly ash is present in the flue gas, so that SO₃ present in the flue gas does not form a harmful mist (submicron particles), but remains in a state condensed on the dust particles and captured in electrostatic precipitator 5 and an absorption tower 8 which will be described later. Accordingly, in the case of a coal-fired boiler, the injection of ammonia in flue 4 is generally omitted.

[0007] Then, the flue gas leaving electrostatic precipitator 5 is conducted through a flue 6 and introduced into the heat recovery section (or heat exchanger) 7 of a gas-gas heater where heat is recovered therefrom. Thereafter, the flue gas is introduced into an absorption tower 8 serving as a desulfurizer. In this absorption tower 8, the flue gas is brought into gas-liquid contact with an absorbing fluid having an absorbent (e.g., limestone) suspended therein (hereinafter referred to as the absorbent slurry), so that chiefly SO₂ present in the flue gas is absorbed into the absorbent slurry and, moreover, the residual dust is also captured by the absorbent slurry. In a tank provided at the bottom of absorption tower 8, the slurry having SO₂ absorbed therein is oxidized to form gypsum as a by-product according to the following reactions including a neutralization reaction.

(Absorption tank)

\[
\text{(Absorption tank)} \quad \text{(Tank)}
\]

[0008] $\text{SO}_2 + \text{H}_2\text{O} \rightarrow \text{H}^+ + \text{HSO}_3^-$  \hspace{1cm} (1)

[0009] $\text{H}^+ + \text{HSO}_3^- + \frac{1}{2}\text{O}_2 \rightarrow 2\text{H}^+ + \text{SO}_4^{2-}$  \hspace{1cm} (2)

$2\text{H}^+ + \text{SO}_4^{2-} + \text{CaCO}_3 + \text{H}_2\text{O} \rightarrow \text{CaSO}_4\cdot2\text{H}_2\text{O} + \text{CO}_2$  \hspace{1cm} (3)

[0010] Then, the flue gas from which SO₂ and the like have been removed in absorption tower 8 serving as a desulfurizer is passed through the reheating section 9 of the gas-gas heater, where it is heated to a temperature favorable
ammonium sulfate (NH₄ HSO₄) will be produced especially in this air heater. An investigation conducted by the present ordinary air heater. Consequently, when ammonia remains in the flue gas leaving denitrator 2, a large amount of acid ° ordinary conditions in this type of equipment, while the flue gas is cooled from about 350 ° according to the reaction formula (4) given below. The dew point of acid ammonium sulfate is about 230 °.

[0017] That is, if ammonia remains in the flue gas, highly adherent acid ammonium sulfate (NH₄HSO₄) is produced according to the reaction formula (4) given below. The dew point of acid ammonium sulfate is about 230°C under ordinary conditions in this type of equipment, while the flue gas is cooled from about 350°C to about 130°C in an ordinary air heater. Consequently, when ammonia remains in the flue gas leaving denitrator 2, a large amount of acid ammonium sulfate (NH₄HSO₄) will be produced especially in this air heater. An investigation conducted by the present inventors has revealed that, in a conventional air heater of the Ljungström type, this acid ammonium sulfate tends to become deposited in the gaps of the heat reservoir within the air heater and hence requires frequent maintenance operations such as cleaning.

\[ \text{SO}_3 + \text{NH}_3 + \text{H}_2\text{O} \rightarrow \text{NH}_4\text{HSO}_4 \]  

(4)

[0018] Accordingly, a first object of the present invention is to provide a flue gas treating process in which the arrangement and construction of the equipment is improved so as to achieve a reduction in the size and cost of the equipment.

[0019] A second object of the present invention is to provide a flue gas treating process which makes it possible to
enhance the flue gas treating performance and thereby achieve a reduction in the size of the equipment and the like.

[0020] A third object of the present invention is to provide a flue gas treating process which can achieve the aforesaid reduction in equipment size and enhancement in performance without detracting from its maintainability.

[0021] A fourth object of the present invention is to provide a flue gas treating process in which the arrangement and construction of the equipment is improved and the flue gas treating performance is enhanced, whereby a marked reduction in the size and cost of the equipment, including a reduction in the size of the stack, can be achieved.

[0022] In order to accomplish the above objects, the present invention provides a flue gas treating process comprising a denitration step of injecting ammonia into flue gas containing at least nitrogen oxides and sulfur oxides to decompose the nitrogen oxides present in the flue gas, and a desulfurization step of introducing the flue gas leaving the denitration step into an absorption tower where it is brought into gas-liquid contact with an absorbing fluid to remove at least the sulfur oxides from the flue gas by absorption into the absorbing fluid, wherein ammonia is injected into the flue gas as required at a point downstream of the denitration step, and the amount of ammonia injected in the denitration step and/or the amount of ammonia injected at the point downstream of the denitration step are determined so as to be on such an excessive level that ammonia or an ammonium salt will remain in the flue gas introduced into the desulfurization step, wherein a region in which a liquid having higher acidity than the absorbing fluid is sprayed so as not to allow ammonia to be easily released into the gaseous phase is created on the downstream side of the region of said absorption tower in which the flue gas is brought into gas-liquid contact with the absorbing fluid, whereby the ammonia remaining in the flue gas introduced into said desulfurization step is absorbed in said absorption tower without allowing it to remain in the flue gas leaving said absorption tower.

[0023] In the flue gas treating process of the present invention, the amount of ammonia injected in the denitration step may be determined so that the concentration of ammonia remaining in the flue gas leaving the denitration step is not less than 30 ppm.

[0024] The flue gas treating process of the present invention may further comprise a heat recovery step of introducing the flue gas leaving the denitration step into a heat exchanger on the upstream side of the absorption tower and thereby recovering heat from the flue gas, and a non-leakage type heat exchanger of shell-and-tube structure may be employed as the heat exchanger.

[0025] The flue gas treating process of the present invention may further include the heat recovery step of introducing the flue gas leaving the denitration step into a heat exchanger on the upstream side of the absorption tower and thereby recovering heat from the flue gas, and the amount of ammonia injected in the denitration step and/or the amount of ammonia injected at the point downstream of the denitration step may be determined so that the concentration of ammonia remaining in the flue gas introduced into the heat exchanger will be in excess of the SO$_3$ concentration in this flue gas by 13 ppm or more.

[0026] The flue gas treating process of the present invention may further include a first dust removal step of introducing the flue gas into a dry electrostatic precipitator on the upstream side of the absorption tower and thereby removing dust present in the flue gas, and a second dust removal step of introducing the flue gas into a wet electrostatic precipitator on the downstream side of the absorption tower and thereby removing the dust remaining in the flue gas.

[0027] In the flue gas treating process of the present invention, ammonia is injected into the flue gas as required at a point downstream of the denitration step, and the amount of ammonia injected in the denitration step and/or the amount of ammonia injected at the point downstream of the denitration step are determined so as to be on such an excessive level that ammonia or an ammonium salt will remain in the flue gas introduced into the desulfurization step.

[0028] And when a region in which a liquid having higher acidity than the absorbing fluid is sprayed so as not to allow ammonia to be easily released into the gaseous phase is created on the downstream side of the region of the absorption tower in which the flue gas is brought into gas-liquid contact with the absorbing fluid, whereby the ammonia remaining in the flue gas introduced into the desulfurization step is absorbed in the absorption tower without allowing it to remain in the flue gas leaving the absorption tower, an adverse effect due to the excessive injection of ammonia (i.e., the emission of ammonia into the atmosphere) can be avoided. This serves to cope with future ammonia emission standards and also contributes to further purification of flue gas.

[0029] Consequently, at least the degree of desulfurization in the desulfurization step is enhanced and this eventually contributes to a reduction in the sizes of the absorption tower and the stack.

[0030] In particular, when the amount of ammonia injected in the denitration step is determined so that the concentration of ammonia remaining in the flue gas leaving the denitration step will be not less than 30 ppm, especially the degree of denitration in the denitration step is markedly enhanced and this eventually contributes to a reduction in the size of the stack.

[0031] Moreover, when the flue gas treating process of the present invention further includes the heat recovery step of introducing the flue gas leaving the denitration step into a heat exchanger on the upstream side of the absorption tower and thereby recovering heat from the flue gas, and a non-leakage type heat exchanger of shell-and-tube structure is employed as the heat exchanger, the heat of the flue gas can be effectively utilized in the conventional manner to preheat air for use in the boiler or reheat the treated flue gas, and troubles due to scale formation can be lessened.
That is, even if acid ammonium sulfate formed by the reaction of the injected ammonia with SO₃ present in the flue gas, and sulfuric acid mist formed from SO₃ present in the flue gas condense in the aforesaid heat exchanger, a non-leakage type heat exchanger of shell-and-tube structure is less subject to the deposition of such materials on the heat transfer surfaces and the like or clogging with them, as compared with a conventionally used heat exchanger of the Ljungström type.

Moreover, in this case, the heat exchanger does not allow air to leak into the flue gas. This can reduce the amount of flue gas to be treated and hence achieve a reduction in cost.

Furthermore, when the flue gas treating process of the present invention further includes the heat recovery step of introducing the flue gas leaving the denitrification step into a heat exchanger on the upstream side of the absorption tower and thereby recovering heat from the flue gas, and the amount of ammonia injected in the denitrification step and/or the amount of ammonia injected at the point downstream of the denitrification step is determined so that the concentration of ammonia remaining in the flue gas introduced into the heat exchanger will be in excess of the SO₃ concentration in this flue gas by 13 ppm or more, the condensation of acid ammonium sulfate in the aforesaid heat exchanger is minimized and a fine powder of neutral ammonium sulfate is chiefly produced. Consequently, the formation of scale due to acid ammonium sulfate is markedly suppressed and this makes the maintenance of the heat exchanger very easy.

Furthermore, when the flue gas treating process of the present invention further includes the first dust removal step of introducing the flue gas into a dry electrostatic precipitator on the upstream side of the absorption tower and thereby removing dust present in the flue gas, and the second dust removal step of introducing the flue gas into a wet electrostatic precipitator on the downstream side of the absorption tower and thereby removing the dust remaining in the flue gas, the dedusting capability of the whole system is markedly improved.

BRIEF DESCRIPTION OF THE DRAWINGS

FIG. 1 is a schematic view illustrating a flue gas treating system in accordance with a first embodiment of the present invention;
FIG. 2 is a schematic view illustrating a flue gas treating system in accordance with a second embodiment of the present invention;
FIG. 3 is a schematic view illustrating the details of the desulfurizer included in the flue gas treating system of FIG. 2;
FIG. 4 is a schematic view illustrating a no-waste water disposal system suitable for use with the flue gas treating system of FIG. 2;
FIG. 5 is a graph showing data which demonstrate an effect of the present invention (i.e., an improvement in degree of desulfurization);
FIG. 6 is a schematic view illustrating an experimental apparatus for demonstrating an effect of the present invention (i.e., the minimization of a deposit due to SO₃ in flue gas);
FIG. 7 is a graph showing experimental results (changes in the gas pressure loss in the heat exchanger) for demonstrating an effect of the present invention (the minimization of a deposit due to SO₃ in flue gas);
FIG. 8 is a graph showing experimental results (changes in the overall heat transfer coefficient of the heat exchanger) for demonstrating an effect of the present invention (the minimization of a deposit due to SO₃ in flue gas);
FIG. 9 is a schematic view illustrating a conventional flue gas treating system; and
FIG. 10 is a schematic view illustrating the denitrator and other apparatus included in the conventional flue gas treating system.

DETAILED DESCRIPTION OF PREFERRED EMBODIMENTS

Several embodiments of the present invention will be described hereinbelow with reference to the accompanying drawings.

First embodiment

A first embodiment of the present invention is described with reference to FIG. 1. The same elements as included in the conventional system of FIG. 9 are designated by the same reference numerals, and the duplicate explanation thereof is omitted.

The flue gas treating system of this embodiment is characterized in that, below the main body 13a of a stack, an absorption tower 21, the reheating section 22 of a gas-gas heater, and a fan 23 are arranged in line on the vertical axis of the stack so as to form part of the stack.
[0040] The heat recovery section 24 of the gas-gas heater is installed at a position which is on the way of a flue 25 connecting a dry electrostatic precipitator 5 with absorption tower 21 and on the inside of the framework 14 for the stack. Eventually, all of absorption tower 21, the heat recovery section 24 and reheating section 22 of the gas-gas heater, and fan 23 are all installed in an unoccupied space on the inside of the framework 14 for the stack.

[0041] In this embodiment, absorption tower 21 is such that the flue gas is introduced thereinto through an inlet formed in the lower lateral part thereof, brought into gas-liquid contact with an absorbing fluid in a countercurrent manner to remove at least sulfur oxides from the flue gas by absorption into the absorbing fluid, and discharged from an outlet formed at the upper end thereof. In the same manner as described previously in connection with the conventional system, gypsum is formed as a by-product by using, for example, limestone as the absorbent.

[0042] The reheating section 22 of the gas-gas heater is directly connected to the upper end of absorption tower 21. Thus, the flue gas discharged from the upper outlet of absorption tower 21 is introduced into reheating section 22 from the bottom side, heated to a temperature favorable for emission into the atmosphere by using the heat recovered in heat recovery section 24, and discharged from the top side.

[0043] In this case, a gas-gas heater of the heating medium circulation type is used, and its heat recovery section 24 and reheating section 22 comprise non-leakage type heat exchangers of shell-and-tube structure. These non-leakage type heat exchangers are advantageous in that, even if SO$_3$ present in the flue gas reacts with ammonia present therein according to the aforementioned reaction formula (4) to produce acid ammonium sulfate (NH$_4$HSO$_4$), they are less subject to the deposition of acid ammonium sulfate which tends to be responsible for scale formation.

[0044] Fan 23 is an axial-flow fan installed above the aforesaid reheating section 22 and functioning to suck the flue gas from the bottom side and discharge it from the top side. The motor is disposed on its internal axis.

[0045] In the arrangement and construction of the flue gas treating system according to this embodiment, the installation space of the whole equipment is significantly reduced as compared with the conventional system illustrated in FIG. 9, so that a marked reduction in the size of the equipment in horizontal directions can be achieved. Specifically, all of the space required for the installation of absorption tower 8, fan 11 and flues 6 and 10 in the system of FIG. 9 becomes unnecessary. Moreover, the installation space of the stack is the same as before, because a space allowing absorption tower 21 and the like to be installed has conventionally been left in the framework 12 for the stack.

[0046] Furthermore, flues 6 and 10 themselves and their supporting members become unnecessary, and the main body of stack 13a becomes much shorter than before. Eventually, a marked reduction in equipment cost is achieved.

Second embodiment

[0047] Next, a second embodiment of the present invention is described with reference to FIG. 2. The same elements as included in the first embodiment are designated by the same reference numerals, and the duplicate explanation thereof is omitted.

[0048] This embodiment is characterized in that it is equipped with an air heater (or heat exchanger) 31 comprising a non-leakage type heat exchanger of shell-and-tube structure, and a wet electrostatic precipitator 32 is installed between absorption tower 21a and reheating section 22. In this embodiment, the aforesaid air heater 31 and the previously described heat recovery section 24 of the gas-gas heater constitute a heat exchanger for carrying out the heat recovery step of the present invention. Moreover, the dry electrostatic precipitator 5 described in connection with the conventional system serves to carry out the first dust removal step of the present invention, and the aforesaid wet electrostatic precipitator 32 serves to carry out the second dust removal step of the present invention.

[0049] The above-described construction has the following advantages. First of all, since air heater 31 comprises a non-leakage type heat exchanger, the amount of scale deposited is relatively small even if acid ammonium sulfate is produced in the flue gas introduced into air heater 31 as described previously. This is very advantageous from the viewpoint of maintenance.

[0050] More specifically, as described previously, when ammonia remains in the flue gas leaving denitrator 2, acid ammonium sulfate is produced especially in air heater 31. According to an investigation conducted by the present inventors, it has been found that, in the case of a conventional air heater of the Ljungström type, such acid ammonium sulfate tends to become deposited in the gaps of the heat reservoir within the air heater and requires frequent maintenance operations such as cleaning.

[0051] However, an investigation conducted by the present inventors has revealed that non-leakage type heat exchangers of shell-and-tube structure are less subject to such deposition of acid ammonium sulfate or clogging with it. It has also been known that the problem concerning the production of acid ammonium sulfate and its deposition on the inner surfaces of the heat exchanger can also be lightened by injecting an excessive amount of ammonia into the flue gas, and this will be specifically described later.

[0052] Moreover, when air heater 31 comprises a non-leakage type heat exchanger, air C fed to the boiler does not leak into the flue gas. This causes a decrease in the flow rate of the flue gas being treated and hence a corresponding reduction in the capacities of fan 23 and the flues and in power consumption.
Furthermore, since wet electrostatic precipitator 32 is installed, the fine dust and other foreign matter which were not captured in absorption tower 21a can be removed. Thus, the residual dust concentration in treated flue gas D is reduced. This enhances the performance of the equipment in that respect and also contributes to a reduction in stack height.

Next, the construction and effects of the characteristic parts of the flue gas treating process of the present invention, which is carried out by using the flue gas treating system of the above-described embodiment, are described hereinbelow.

According to this process, instead of using an ammonia decomposition catalyst, the amount of ammonia B injected in denitrator 2 is determined so as to be on such an excessive level that a large amount of ammonia or an ammonium salt will remain in the flue gas introduced into absorption tower 21a.

If ammonia or an ammonium salt remains in the flue gas introduced into absorption tower 21a, this ammonia or ammonium salt is dissolved in the slurry within absorption tower 21a as a result of gas-liquid contact between the flue gas and the absorbent slurry. This raises the ammonium salt concentration (in other words, ammonium ion concentration) in the liquid phase of the slurry circulating through absorption tower 21a.

An investigation conducted by the present inventors has revealed that, when the ammonium salt concentration (or ammonium ion concentration) in the circulating fluid of the absorption tower is increased to 150 mmol/liter or more, the degree of removal of sulfur dioxide from the flue gas (i.e., the degree of desulfurization) in the absorption tower rises to the vicinity of 95% as shown in FIG. 5, even if the other conditions remain constant. Consequently, according to this embodiment in which the amount of ammonia B injected in denitrator 2 is determined so as to be on an excessive level as described above, the size of absorption tower 21a can be reduced as compared with the prior art. Moreover, the concentration of sulfur oxides (typically sulfur dioxide) remaining in treated flue gas D can further be reduced and, therefore, the height of the stack can be reduced.

Furthermore, since the amount of ammonia B injected is naturally in excess of the equivalent amount required for denitrification, the denitrification capability of denitrator 2 (or the denitrification step) is enhanced. According to an investigation conducted by the present inventors, it has been found that, if the amount of ammonia B injected is determined so that it is in excess of the equivalent amount required for denitrification and, moreover, the concentration of ammonia remaining in the flue gas leaving the denitrification step (i.e., slip ammonia) will be not less than 30 ppm, the degree of denitrification is enhanced from the conventional level of about 80% to about 90% and the concentration of nitrogen oxides remaining in treated flue gas D can be reduced to half.

In this connection, trial calculations made by the present inventors with respect to an electric power plant of the 150 MW class indicate that, if the degree of desulfurization and the degree of denitrification are enhanced as described above, and the degree of dust removal is also enhanced by the installation of wet electrostatic precipitator 32, the stack height (L1) can be markedly reduced from the conventional value of about 150 m to about 90 m. Moreover, in consequence of that, the width (W1) of the installation space of framework 14b for the stack can be markedly reduced from the conventional value of about 38 m to about 25 m.

In this case, the particular amount of ammonia B injected must be determined not only so as to be in excess of the equivalent amount required for denitrification, but also with consideration for the SO₃ concentration in the flue gas. Specifically, at least a portion of the ammonia remaining in the flue gas leaving denitrator 2 (or the denitrification step) reacts with SO₃ present in this flue gas to form ammonium salts such as the above-described ammonium sulfate (NH₄)₂SO₄ and the production of highly adherent acid ammonium sulfate tending to form scale becomes relatively little. Moreover, according to this embodiment using air heater 31 of shell-and-tube structure, troubles due to scale formation are less as compared with the conventional system using an air heater of the Ljungström type. Eventually, the problem of scale formation due to acid ammonium sulfate can be practically solved and the denitrator
need not be provided with an ammonia decomposition catalyst.

[0065] In this embodiment, an excessive amount of ammonia is positively injected so that a large amount of ammonia or an ammonium salt will remain in the flue gas introduced into absorption tower 21a. Accordingly, the disposal of gas-liquid contact absorbed into the slurry within absorption tower 21a and the ammonia leaking into treated flue gas D pose problems. However, these problems can be solved by employing the existing no-waste water disposal technique (the so-called AWMT) in which ammonia is recovered and reused by mixing dust from the electrostatic precipitator with waste water from the desulfurizer, or an ammonia absorption technique newly devised by the present inventors.

[0066] Some embodiments of these techniques are described below with reference to FIGs. 3 and 4. FIG. 3 is a schematic view showing, in particular, the detailed construction of a desulfurizer suitable for use in the flue gas treating system of this embodiment as illustrated in FIG. 2, and FIG. 4 is a schematic view showing the construction of an exemplary no-waste water disposal system suitable for use with the flue gas treating system of this embodiment (in the case of an oil-fired boiler).

[0067] In this case, as illustrated in FIG. 3, absorption tower 21a serving as a desulfurizer is an absorption tower of the liquid column type which is provided at the bottom with a tank 41 for holding an absorbing fluid E having an absorbent (i.e., limestone) suspended therein (hereinafter referred to as absorbent slurry E) and which has a gas-liquid contact region extending above tank 41 and serving to bring flue gas into gas-liquid contact with the slurry within tank 41.

[0068] This absorption tower 21a is a so-called counterflow absorption tower in which a flue gas inlet section 42 for introducing flue gas is formed in its lower part and a flue gas outlet section 43 for discharging the desulfurized flue gas A1 is formed in its upper end, so that the flue gas enters from the lower part of the absorption tower and flows upward.

[0069] A mist eliminator 43a is installed in flue gas outlet section 43. This mist eliminator 43a serves to collect any mist produced as a result of gas-liquid contact and entrained by the flue gas, so that a large amount of mist containing sulfur dioxide, ammonia and the like may not be discharged together with the desulfurized flue gas A1. In this embodiment, the mist collected by this mist eliminator 43a is allowed to flow down from its lower end and return directly to tank 41.

[0070] Moreover, in absorption tower 21a, a plurality of spray pipes 44 are disposed in parallel. In these spray pipes 44, a plurality of nozzles (not shown) for injecting the slurry within tank 41 upward in the form of liquid columns are formed.

[0071] Furthermore, a circulating pump 45 for withdrawing and raising the absorbent slurry within tank 41 is installed on the outside of tank 41. Thus, the slurry is fed to spray pipes 44 through a circulation line 46.

[0072] In the embodiment illustrated in FIG. 3, tank 41 is provided with a means for blowing air F for oxidizing use in the form of fine bubbles while agitating the slurry within tank 41. This means comprises an agitator 47 and an air supply pipe 48 for blowing air F into the slurry in the vicinity of the agitating blades of agitator 47. Thus, the absorbent slurry having sulfur dioxide absorbed therein is brought into efficient contact with the air in tank 41 and thereby completely oxidized to form gypsum.

[0073] More specifically, the absorbent slurry injected from spray pipes 44 within absorption tower 21a flows downward while absorbing sulfur dioxide and dust (containing ammonium salts such as ammonium sulfate) and, moreover, ammonia gas as a result of gas-liquid contact with flue gas, and enters tank 41 where it is oxidized by contact with a large number of air bubbles blown thereinto while being agitated by means of agitator 47 and air supply pipe 48, and then undergoes a neutralization reaction to become a slurry containing gypsum at a high concentration. The dominant reactions occurring in the course of these treatments are represented by the aforementioned reaction formulas (1) to (3).

[0074] Thus, a large amount of gypsum, a small amount of limestone (used as the absorbent), and a slight amount of dust and ammonium collected from the flue gas are steadily suspended or dissolved in the slurry within tank 41. In this embodiment, the slurry within tank 41 is withdrawn and fed to a solid-liquid separator 49 through a pipe line 46a branching from circulation line 46. The slurry is filtered in solid-liquid separator 49, so that gypsum G having a low water content is recovered. On the other hand, a portion H1 of the filtrate from solid-liquid separator 49 is fed to a slurry preparation tank 52 as water constituting absorbent slurry E, and the remainder is discharged as desulfurization waste water H2 in order to prevent the accumulation of impurities.

[0075] Since the ammonia and ammonium salts (such as ammonium sulfate) absorbed from the flue gas have high solubilities, most of them are contained in the liquid phase of slurry E and eventually discharged together with desulfurization waste water H2.

[0076] In this embodiment, a slurry containing limestone as the absorbent is fed from slurry preparation tank 52 to tank 41 during operation. This slurry preparation tank 52 is equipped with a stirrer 53 and serves to prepare absorbent slurry E by mixing powdered limestone I introduced from a silo (not shown) with filtrate H1 fed as described above, and stirring this mixture. Absorbent slurry E within slurry preparation tank 52 is suitably fed to tank 41 by means of a slurry pump 54. Moreover, in order to make up for the water gradually lost owing to evaporation in absorption tower 21a or the like, make-up water (such as industrial water) is suitably supplied, for example, to tank 41 or slurry preparation tank 52.

[0077] During operation, the flow rate of the aforesaid make-up water supplied to tank 41, the flow rate of the slurry
withdrawn through pipe line 46a, and the like are suitably controlled. Thus, tank 41 is maintained in such a state that the slurry containing gypsum and the absorbent at predetermined concentrations is always stored therein at a level within certain limits.

[0078] Also during operation, in order to maintain the degree of desulfurization and the purity of gypsum at a high level, the boiler load (i.e., the flow rate of flue gas A), the sulfur dioxide concentration in the flue gas introduced into absorption tower 21a, the pH and limestone concentration of the absorbent slurry within tank 41, and the like are detected with sensors. On the basis of the results of detection, the feed rate of limestone to tank 41 and other parameters are suitably controlled by means of a controller (not shown). Conventionally, the pH of the absorbent slurry within tank 41 is usually adjusted to about 6.0 so that highly pure gypsum may be formed by accelerating the above-described oxidation reaction while maintaining the high capacity to absorb sulfur dioxide.

[0079] Furthermore, as a means for preventing the ammonia injected in excess from remaining in the desulfurized flue gas A1, spray pipes 55 are installed above spray pipes 44. These spray pipes 55 serve to inject a liquid J (which may be in the form of a slurry) having a lower pH value than the slurry within tank 41 into absorption tower 21a and thereby create, in the upper part of absorption tower 21a, a region which does not allow ammonia to be easily released into the gaseous phase.

[0080] For example, liquid J injected from spray pipes 55 comprises a dilute solution of sulfuric acid, and its pH is adjusted to a value (e.g., 4.0 to 5.0) at which ammonia is not easily released into the flue gas.

[0081] In the above-described construction, the flue gas introduced into absorption tower 21a through flue gas inlet section 42 is first brought into gas-liquid contact with the slurry injected from spray pipes 44 in the form of liquid columns, and then brought into gas-liquid contact with the liquid injected from spray pipes 55. Thus, dust and ammonia, together with sulfur dioxide, are absorbed or captured.

[0082] During this process, the liquid injected from spray pipes 55 in the outlet section (or upper end) of absorption tower 21a is adjusted to a pH value at which ammonia is not easily released into the flue gas. Consequently, the partial pressure of ammonia is depressed in the upper part of absorption tower 21a, so that the phenomenon in which the ammonia once dissolved in the liquid phase of the slurry is reversely released into the flue gas in the upper part of the absorption tower is avoided.

[0083] Thus, desulfurized flue gas A1 having very low concentrations of sulfur dioxide, dust and ammonia is ultimately discharged from flue gas outlet section 43 formed at the upper end of absorption tower 21a. In this case, calculations made by the present inventors have revealed that the degree of removal of ammonia is about 90%. Consequently, in spite of the construction in which an excessive amount of ammonia is positively injected, little ammonia is contained in treated flue gas D (FIG. 2) and no problem with ammonia emission into the atmosphere arises.

[0084] However, it is desirable from the viewpoint of air pollution prevention to minimize the ammonia concentration in treated flue gas D emitted into the atmosphere. Accordingly, there has been a demand for a flue gas treating technique which can achieve a reduction in equipment size and a high degree of desulfurization and, moreover, can minimize the amount of ammonia emitted.

[0085] Next, the construction of the exemplary no-waste water disposal system illustrated in FIG. 4 is described below. This is an example relating to the treatment of flue gas from an oil-fired boiler. In this case, dust K collected from flue gas by dry electrostatic precipitator 5 illustrated in FIGs. 2 and 3 contains, in addition to unburned carbon constituting its principal component, impurities such as vanadium (which is a toxic heavy metal) and magnesium, ammonium sulfate formed from the injected ammonia and SO3 present in the flue gas, and the like.

[0086] In this system, as illustrated in FIG. 4, desulfurization waste water H2 shown in FIG. 3 is first introduced into a mixing tank 61 where it is stirred and mixed with dust K fed from dry electrostatic precipitator 5 to form a mixed slurry S1. In this step, ammonia and ammonium sulfate contained in dust K are dissolved in the liquid phase of slurry S1, and most of them exist as sulfate ion or ammonium ion, similarly to those contained in waste water H2. Then, mixed slurry S1 is transferred to a precipitation tank 63 having undergone the reduction of vanadium is transferred to a precipitation tank 63 where ammonia B3 (which will be described later) is added to and mixed with the slurry. In this step, tetravalent vanadium present in the slurry reacts with ammonia according to the following reaction formula (5), so that the vanadium is dissolved in the liquid phase.

\[ 2VO_2^+ + SO_3^{2-} + 2H^+ \rightarrow 2VO_2^{2+} + SO_4^{2-} + H_2O \]  

(5)

Then, mixed slurry S2 having undergone the reduction of vanadium is transferred to a precipitation tank 63 having undergone the reduction of vanadium is transferred to a precipitation tank 63 where ammonia B3 (which will be described later) is added to and mixed with the slurry. In this step, tetravalent vanadium present in the slurry reacts with ammonia according to the following reaction formula (6), and the resulting product precipitates.
After being treated for the precipitation of vanadium, mixed slurry S3 is withdrawn from precipitation tank 63 and transferred to a solid-liquid separator 65 comprising a flocculating settler and/or a vacuum type belt filter by means of a slurry pump 64. Thus, solid matter N is separated therefrom in the form of sludge or cake. The separated solid matter N consists essentially of unburned carbon present in dust K, and additionally contains the precipitated vanadium.

Then, waste liquor S4 from which solid matter containing vanadium has been removed is transferred to a neutralization tank 66 where a chemical agent O [e.g., slaked lime (Ca(OH)₂)] and return waste liquor P (which will be described later) are added thereto with stirring. Thus, sulfur ion and ammonium ion present in the waste liquor are converted into gypsum or ammonium hydroxide.

The resulting slurry S5, which now contains gypsum as a solid component and ammonium hydroxide, is then transferred to a primary concentrator 67 where ammonia B1 is separated therefrom by evaporation. The resulting slurry S6 containing gypsum and other solid matter at high concentrations is withdrawn therefrom by means of a slurry pump 68.

Primary concentrator 67 consists of an evaporator 67a, a heater 67b and a circulating pump 67c, and functions to heat the slurry with hot steam W1 generated, for example, in a boiler of an electric power plant and thereby evaporate water B1 containing ammonia.

Besides gypsum, the solid matter contained in slurry S6 includes mainly magnesium hydroxide [Mg(OH)₂]. This magnesium hydroxide is formed by the combination of magnesium present in dust K as an impurity with hydroxide ion present in the slurry.

Subsequently, this slurry S6 is introduced into a solid separator 69 comprising a cyclone or a sedimentation centrifuge, where slurry S6 is separated into a slurry S7 containing chiefly gypsum (coarsely particulate solid matter) and a slurry S8 containing other finely particulate solid matter (including mainly the above-described magnesium hydroxide). Slurry S7 is returned to absorption tower tank 41 constituting the desulfurizer illustrated in FIG. 3. On the other hand, a portion of slurry S8 is dehydrated in a dehydrator (or secondary concentrator) 70, and the resulting solid matter containing chiefly magnesium hydroxide is discharged as sludge Q.

Of slurry S8, the remaining portion not fed to dehydrator 70 is returned to neutralization tank 66 as return waste liquor P.

Ammonia water B1 produced by evaporation in primary concentrator 67 is cooled and condensed in a cooler 71 using cooling water W2 as the coolant, and stored in a storage tank 72.

Since ammonia water B1 within storage tank 72 usually has a low concentration of about 3-6%, it is fed to an ammonia concentrator 74 by means of a pump 73 and concentrated therein to yield ammonia water having a concentration of 10-20%. A portion of this ammonia water is gasified in a vaporizer 75, and the resulting ammonia B2 is injected into the flue gas in the previously described denitrator 2 as ammonia B containing steam W3. The remaining portion of the ammonia water is fed to the aforesaid precipitation tank 63 as ammonia B3.

In the above-described no-waste water disposal system, desulfurization waste water H2 resulting from the desulfurization operation is treated by mixing it with the removed dust K, so that an improvement in handleability is achieved. The resulting mixed slurry is subjected to a series of treatments for the reduction, precipitation and solid-liquid separation of vanadium present therein, and the separated vanadium is discharged as sludge. Moreover, the mixed slurry concentrated and treated in such a way that gypsum, water and ammonia are finally returned to the flue gas or to the upstream side of the system (e.g., the absorption tower constituting the desulfurizer). Thus, it becomes possible to use ammonia in a circulating manner and realize a so-called no-waste water closed system in which no waste water to be discharged is produced. This eliminates the need of waste water treatment prior to its discharge and enables the effective utilization of ammonia.

Experiments

Now, some experiments carried out by the present inventors are described below. The purpose of these experiments is to demonstrate that the formation of scale on the inner surfaces (e.g., heat transfer surfaces) of the heat exchanger for the heat recovery step due to acid ammonium sulfate is suppressed when an excessive amount of ammonia is injected according to one feature of the present invention and a non-leakage type heat exchanger of shell-and-tube structure is used as the heat exchanger.

First of all, an experimental apparatus illustrated in FIG. 6 was used. Specifically, an air heater 82 and a cooler 83 were installed downstream of a combustion furnace 81. Moreover, a cyclone 84 for separating and removing dust such as unburned carbon was installed downstream of them. The flue gas leaving this cyclone 84 was introduced into a non-leakage type heat exchanger 85 of shell-and-tube structure. In this case, the flue gas was passed through heat...
exchanger 85 on its shell side (i.e., on the outside of the heating tubes), while a heating medium was passed through the heating tubes of heat exchanger 85. The heating medium heated by the heat of the flue gas in heat exchanger 85 was cooled and regenerated in a cooler 86 using cooling water.

[0100] The SO$_3$ concentration in the flue gas was regulated by injecting SO$_3$ into the flue gas at a position downstream of air heater 82 and upstream of cooler 83. Moreover, the ammonia concentration in the flue gas was regulated by injecting ammonia into the flue gas at a position downstream of cyclone 84 and upstream of heat exchanger 85. Heat exchanger 85 was capable of so-called steel ball cleaning by scattering steel balls continuously on its shell side, and was subjected to steel ball cleaning tests as required.

[0101] Other experimental conditions were as follows.

(Fuel)

[0102] Type: Fuel oil A.
Combustion rate: 15 liters/hr.

(Flue gas)

SO$_3$ concentration: 25 ppm.
NH$_3$ concentration: 63 ppm.
Temperature at the outlet of the cyclone: 170°C.
Temperature at the inlet/outlet of the heat exchanger: 130/90°C.

(Heating medium)

[0104] Inlet temperature: 75°C.

(Steel balls)

[0105] Scattering rate: 2,280 kg/m$^2$·h.

[0106] In this case, the ammonia concentration in the flue gas was determined to be 63 (= 50+13) ppm so that it would be in excess of the equivalent amount required for the formation of ammonium sulfate [(NH$_4$)$_2$SO$_4$] as a result of reaction with SO$_3$ by 13 ppm. The aforesaid equivalent amount is equal to twice the number of moles of SO$_3$ and, in this case, corresponded to a concentration of 50 (= 25x2) ppm.

[0107] Under the above-described conditions, the experimental apparatus was continuously operated for 83 hours without steel ball cleaning. Thereafter, steel ball cleaning was carried out for 2 hours.

[0108] FIG. 7 shows the results of actual measurement of changes in the gas pressure loss in heat exchanger 85, and FIG. 8 shows the results of actual measurement of changes in the overall heat transfer coefficient of heat exchanger 85 which serves as an indication of its heat transfer capability.

[0109] As can be seen from these results, the changes in gas pressure loss and overall heat transfer coefficient were relatively slight even after 83 hours' continuous operation. Moreover, they can be completely restored to their initial levels by steel ball cleaning.

[0110] Moreover, when the surfaces of the heating tubes of heat exchanger 85 after 83 hours' continuous operation were photographed and visually observed with the naked eye, the accumulation of a deposit was slight. On analysis, this deposit includes mainly an ammonium sulfate type compound having an NH$_4$/SO$_4$ molar ratio of 1.5 to 1.9.

[0111] Thus, it can be seen that, if ammonia is injected in such an amount as to be in excess of the SO$_3$ concentration by 13 ppm or more, the production of acid ammonium sulfate is suppressed and this greatly facilitates the operation for removing a deposit.

[0112] It is to be understood that the present invention is not limited to the above-described embodiments, but may
also be practiced in various other ways.

[0113] For example, the injection of ammonia need not only be carried out in the denitrator (or the denitration step), but may also be carried out at any point downstream of the denitrator and upstream of the absorption tower. By way of example, in the system of FIG. 2, ammonia may be injected into the flue gas in flue 4 for the purpose of capturing SO3 and enhancing the desulfurization capability, or ammonia may be injected into the flue gas in flue 25 (on the downstream side of dry electrostatic precipitator 5) for the purpose of enhancing the desulfurization capability and the like.

[0114] In this case, in order to enhance the desulfurization capability of the absorption tower, the amount of ammonia injected in the denitration step and/or the amount of ammonia injected at the point downstream of the denitration step may be determined so that it is in excess of the equivalent amount required for denitration or the equivalent amount for SO3 and, therefore, ammonia or an ammonium salt (e.g., ammonium sulfate) will remain in the flue gas introduced into the desulfurization step.

[0115] Moreover, in order to fully suppress the formation of scale due to SO3 in air heater 31 and the heat recovery section 24 of the gas-gas heater, the amount of ammonia injected in the denitration step and/or the amount of ammonia injected at the point downstream of the denitration step may be determined so as to be on such an excessive level that the concentration of ammonia remaining in the flue gas introduced into these heat exchangers will be in excess of the SO3 concentration in this flue gas by 13 ppm or more.

[0116] Furthermore, air heater 31 and the heat recovery section 24 of the gas-gas heater are separately installed, for example, in the previously described embodiment of FIG. 2. However, they may be combined into a single unit. That is, the system may be constructed in such a way that air C fed to the boiler is heated by the heat recovered in a heat exchanger installed, for example, at the position of air heater 31 (FIG. 2), and a portion of the heating medium is conducted to the reheating section 22 of the gas-gas heater and used to heat treated flue gas D.

[0117] Even where the air heater and the heat recovery section of the gas-gas heater are separately installed, the heat recovery section of the gas-gas heater may be installed at a position upstream of electrostatic precipitator 5.

[0118] In this connection, if heat recovery from the flue gas is fully carried out on the upstream side of electrostatic precipitator 5 and the temperature of the flue gas introduced into electrostatic precipitator 5 is further reduced, this is advantageous especially in the case of flue gas from a coal-fired boiler, because the degree of removal of dust (e.g., fly ash) in electrostatic precipitator 5 is markedly improved on the basis of its increased resistivity.

Claims

1. A flue gas treating system comprising a denitration step of injecting ammonia into flue gas (A) containing at least nitrogen oxides and sulfur oxides to decompose the nitrogen oxides present in the flue gas, and a desulfurization step of introducing the flue gas leaving said denitration step into an absorption tower (21) where it is brought into gas-liquid contact with an absorbing fluid (E) to remove at least the sulfur oxides from the flue gas by absorption into the absorbing fluid, wherein ammonia (B) is injected into the flue gas as required at a point downstream of said denitration step, and the amount of ammonia injected in said denitration step and/or the amount of ammonia injected at the point downstream of said denitration step are determined so as to be on such an excessive level that ammonia or an ammonium salt remains in the flue gas introduced into said desulfurization step, wherein a region in which a liquid (J) having higher acidity than the absorbing fluid (E) is sprayed so as not to allow ammonia to be easily released into the gaseous phase is created on the downstream side of the region of said absorption tower in which the flue gas is brought into gas-liquid contact with the absorbing fluid, whereby the ammonia remaining in the flue gas introduced into said desulfurization step is absorbed in said absorption tower without allowing it to remain in the flue gas (D) leaving said absorption tower.

2. A flue gas treating process as claimed in Claim 1, wherein the amount of ammonia injected in said denitration step is determined so that the concentration of ammonia remaining in the flue gas leaving said denitration step is not less than 30 ppm.

3. A flue gas treating process as claimed in Claim 1 or Claim 2, further comprising a heat recovery step of introducing the flue gas leaving said denitration step into a heat exchanger (24) on the upstream side of said absorption tower and thereby recovering heat from the flue gas and wherein a non-leakage type heat exchanger (85) of shell-and-tube structure is employed as said heat exchanger.

4. A flue gas treating process as claimed in Claim 1 or Claim 3 which further includes the heat recovery step of
introducing the flue gas leaving said denitration step into a heat exchanger (24) on the upstream side of said absorption tower and thereby recovering heat from the flue gas and wherein the amount of ammonia injected in said denitration step and/or the amount of ammonia injected at the point downstream of said denitration step are determined so that the concentration of ammonia remaining in the flue gas introduced into said heat exchanger will be in excess of the SO₃ concentration in this flue gas by 13 ppm or more.

5. A flue gas treating process as claimed in any of Claims 1 to 3, further comprising a first dust removal step of introducing the flue gas into a dry electrostatic precipitator (5) on the upstream side of said absorption tower (21) and thereby removing dust present in the flue gas, and a second dust removal step of introducing the flue gas into a wet electrostatic precipitator (32) on the downstream side of said absorption tower and thereby removing the dust remaining in the flue gas.

Patentansprüche

1. Rauchgas-Behandlungssystem, das einen Denitrierungsschritt unter Injektion von Ammoniak in Rauchgas (A), das zumindest Stickoxide und Schwefeloxide enthält, zur Zersetzung der in dem Rauchgas vorhandenen Stickoxide, und einen Entschwefelungsschritt unter Einführung des Rauchgases, das den Denitrierungsschritt verlässt, in einen Absorptionsturm (21), worin es in einen Gas-Flüssigkeits-Kontakt mit einem absorbierenden Fluidum (E) zur Entfernung von zumindest der Schwefeloxide aus dem Rauchgas durch Absorption in das absorbierende Fluidum umfasst,
worin Ammoniak (B) nach Bedarf an einem Punkt stromabwärts des Denitrierungsschrittes in das Rauchgas injiziert wird, und
die Menge an in dem Denitrierungsschritt injizierten Ammoniak und/oder die Menge an Ammoniak, die an dem Punkt stromabwärts des Denitrierungsschrittes injiziert wird, werden auf einen solchen Überschuss festgelegt, dass Ammoniak oder ein Ammoniumsalz in dem Rauchgas, das in den Entschwefelungsschritt eingeführt wird, zurückbleibt, worin ein Bereich, in dem eine Flüssigkeit (J) mit einer höheren Acidität als das absorbierende Fluidum (E) so eingespritzt wird, dass das Ammoniak nicht leicht in die Gasphase abgegeben wird, auf der stromabwärtigen Seite des Bereichs des Absorptionsturms erzeugt wird, indem das Rauchgas in einen Gas-Flüssig-Kontakt mit dem absorbierenden Fluidum gebracht wird, wodurch das in dem Rauchgas zurückbleibende Ammoniak, das in den Entschwefelungsschritt eingeführt wird, in dem Absorptionsturm absorbiert wird, ohne dass es in dem Rauchgas (D), das den Absorptionsturm verlässt, zurückbleibt.

2. Rauchgas-Behandlungsverfahren gemäß Anspruch 1, worin die Menge an Ammoniak, das in den Denitrierungsschritt eingeführt wird, so bestimmt wird, dass die Konzentration an Ammoniak, die in dem Rauchgas, das den Denitrierungsschritt verlässt, zurückbleibt, nicht weniger als 30 ppm beträgt.


4. Rauchgas-Behandlungsverfahren gemäß Anspruch 1 oder 3, das ferner den Wärmerückgewinnungsschritt unter Einführung des Rauchgases, das den Denitrierungsschritt verlässt, in einen Wärmeaustauscher (24) auf der stromaufwärtigen Seite des Absorptionsturms umfasst, wodurch Wärme aus dem Rauchgas zurückgewonnen wird, und worin die Menge an Ammoniak, die im Denitrierungsschritt injiziert wird, und/oder die Menge an Ammoniak, die an dem Punkt stromabwärts des Denitrierungsschrittes injiziert wird, so bestimmt wird, dass die Konzentration des Ammoniaks, das im Rauchgas zurückbleibt, das in den Wärmeaustauscher eingeführt wird, in einem Überschuss gegenüber der SO₃-Konzentration in diesem Rauchgas von 13 ppm oder mehr vorhanden ist.

Revendications

1. Système de traitement d'effluents gazeux comprenant une étape de dénitrification en injectant de l'ammoniac dans les effluents gazeux (A) contenant au moins des oxydes d'azote et des oxydes de soufre pour décomposer les oxydes d'azote présents dans les effluents gazeux, et une étape de désulfuration en introduisant les effluents gazeux quittant ladite étape de dénitrification dans une tour d'absorption (21) où ils sont amenés dans un contact gaz-liquide avec un fluide d'absorption (E) pour enlever au moins les oxydes de soufre des effluents gazeux par absorption dans le fluide d'absorption, dans lequel de l'ammoniac (B) est injecté dans les effluents gazeux tel que requis au niveau d'un point en aval de ladite étape de dénitrification, et la quantité d'ammoniac injecté dans ladite étape de dénitrification et/ou la quantité d'ammoniac injecté au niveau du point en aval de ladite étape de dénitrification sont déterminées de façon à avoir un niveau si élevé que de l'ammoniac ou un sel d'ammonium subsiste dans les effluents gazeux introduits dans ladite étape de désulfuration, où une zone dans laquelle un liquide (J) ayant une acidité supérieure au fluide d'absorption (E) est pulvérisé afin d'empêcher l'ammoniac d'être libéré de façon aisé dans la phase gazeuse qui est créée du côté aval de la zone de ladite tour d'absorption dans laquelle les effluents gazeux sont amenés en contact gaz-liquide avec le fluide d'absorption, l'ammoniac restant dans les effluents gazeux introduits dans ladite étape de désulfuration étant absorbé dans ladite tour d'absorption sans lui permettre de rester dans les effluents gazeux (D) quittant ladite tour d'absorption.

2. Processus de traitement d'effluents gazeux selon la revendication 1, dans lequel la quantité d'ammoniac injecté dans ladite étape de dénitrification est déterminée de sorte que la concentration d'ammoniac restant dans les effluents gazeux quittant ladite étape de dénitrification n'est pas inférieure à 30 ppm.

3. Processus de traitement d'effluents gazeux selon la revendication 1 ou la revendication 2, comprenant de plus une étape de récupération de chaleur en introduisant les effluents gazeux quittant ladite étape de dénitrification dans un échangeur thermique (24) du côté amont de ladite étape de désulfuration, et récupérant de ce fait de la chaleur provenant des effluents gazeux, et dans lequel un échangeur thermique du type sans fuite (85) ayant une structure multitubulaire à calandre est utilisé en tant que dit échangeur thermique.

4. Processus de traitement d'effluents gazeux selon la revendication 1 ou la revendication 3, qui comprend de plus l'étape de récupération de chaleur en introduisant les effluents gazeux quittant ladite étape de dénitrification dans un échangeur thermique (24) du côté amont de ladite étape de désulfuration, et récupérant de ce fait de la chaleur provenant des effluents gazeux, et dans lequel la quantité d'ammoniac injecté dans ladite étape de dénitrification et/ou la quantité d'ammoniac injecté au niveau du point en aval de ladite étape de dénitrification sont déterminées de sorte que la concentration d'ammoniac restant dans les effluents gazeux introduits dans ladit échangeur thermique va dépasser la concentration en SO₃ dans ces effluents gazeux de 13 ppm ou plus.

5. Processus de traitement d'effluents gazeux selon l'une quelconque des revendications 1 à 3, comprenant de plus une première étape d'enlèvement de poussière en introduisant les effluents gazeux dans un dépoussiéreur électrostatique à sec (5) du côté amont de ladite étape d'absorption (21), et enlevant de ce fait la poussière présente dans les effluents gazeux, et une seconde étape d'enlèvement de poussière en introduisant les effluents gazeux dans un dépoussiéreur électrostatique humide (32) du côté aval de ladite étape d'absorption, et enlevant de ce fait la poussière restant dans les effluents gazeux.
FIG. 5

DEGREE OF REMOVAL OF SO₂ IN ABSORPTION TOWER [%]

AMMONIUM SALT CONCENTRATION IN CIRCULATING FLUID OF ABSORPTION TOWER [mmol/l]
FIG. 6

STEEL BALLS

FLUE GAS

RETURN OF STEEL BALLS

COOLING WATER

NH₃

SO₃

AIR

FUEL

FLUE GAS

HEATING MEDIUM
FIG. 7

GAS PRESSURE LOSS IN WHOLE HEAT EXCHANGER [mmH₂O]
FIG. 8

OVERALL HEAT TRANSFER COEFFICIENT OF HEAT EXCHANGER
[kcal/m²·h·°C]
FIG. 10

PRIOR ART