Observations of production and emission of greenhouse gases and ammonia during storage of solids separated from pig slurry: Effects of covering

M.N. Hansen\textsuperscript{a,*}, K. Henriksen\textsuperscript{b}, S.G. Sommer\textsuperscript{a}

\textsuperscript{a}Danish Institute of Agricultural Sciences, Department of Agricultural Engineering, Schüttesvej 17, DK-8700 Horsens, Denmark
\textsuperscript{b}Aalborg University, Institute of Life Sciences, Environmental Engineering Section, Sohngaardsholmvej 57, 9000, Aalborg

Received 6 September 2005; received in revised form 26 January 2006; accepted 2 February 2006

Abstract

Separation of slurry produces a solid fraction that is stored in manure heaps before being used as a fertiliser in crop production. Considerable amounts of ammonia (NH\textsubscript{3}) and greenhouse gases may be emitted during storage, which has deleterious environmental effects. The emission levels can be expected to depend on oxygenation level inside the bulk of the stored manure and therefore on storage conditions. An experiment was thus set up to study gaseous emissions during storage of the solid fraction, and the effects of the oxygenation of manure heaps on emissions of NH\textsubscript{3} and various greenhouse gases. Emissions of NH\textsubscript{3} and the greenhouse gases methane (CH\textsubscript{4}), nitrous oxide (N\textsubscript{2}O), and carbon dioxide (CO\textsubscript{2}) from an uncovered and covered heap of solids separated from pig slurry were compared, and related to the oxygenation level inside the manure heap. Approximately 15% of the initial nitrogen content was lost when separated solids were stored uncovered. Of the initial nitrogen content, 4.8% was lost as N\textsubscript{2}O, 0.3% was lost as NH\textsubscript{3}, while the 9.6% unaccounted for was assumed lost as dinitrogen (N\textsubscript{2}). Of the initial carbon content, 28% was lost during uncovered storage; the majority of this was emitted as CO\textsubscript{2} (25%), while 1.3% was emitted as CH\textsubscript{4}. Oxygenation level inside the heap was found to influence the production and emission of greenhouse gases. Covering the heap with an airtight material delayed aeration of the bulk of the stored manure, which reduced the internal heat production, degradation of organic matter, and emission of NH\textsubscript{3} and greenhouse gases. Emissions of NH\textsubscript{3}, N\textsubscript{2}O, and CH\textsubscript{4} were reduced by 12%, 99%, and 88%, respectively, when the manure heap was covered with an airtight material.

\textsuperscript{c} 2006 Elsevier Ltd. All rights reserved.

Keywords: Solid manure; Separation; Ammonia; Methane; Nitrous oxide

1. Introduction

Slurry separation is a technology that separates slurry into a minor solid fraction containing a majority of the dry matter and phosphorus content of slurry, and a larger liquid fraction containing most of the ammonium–nitrogen and potassium content (Møller et al., 2002). In Denmark, slurry separation is mainly performed to reduce the risk of phosphorus accumulation on agricultural land in areas of dense livestock production, as it allows the transport of slurry phosphorus at a lower cost due
to the volume reduction (Møller et al., 2000a). The solid fraction produced by slurry separation has a high phosphorus content and is therefore used as a fertilizer in crop production. Before being used it is stored in heaps, in which aerobic conditions normally allow composting (Eghball et al., 1997; Hao and Chang, 2001). Composting is caused by high microbial activity, which increases the temperature, pH, and decomposition of organic matter of the manure (Møller et al., 2000b; Amon et al., 2001; Beck-Friis, 2001; Sommer, 2001). As the potential for ammonia emission is increased by the elevation of temperature and pH taking place during the composting of solid manure (Dewes, 1996), a significant part the initial nitrogen (N) content of solid manure may be lost during storage (Eghball et al., 1997; Petersen et al., 1998; Sommer and Dahl, 1999; Møller et al., 2000b; Missetbrook et al., 2001; Webb et al., 2001). It has been shown that the emission of ammonia from solid manure heaps can be reduced by covering (Webb et al., 2001, Sommer, 2001; Chadwick, 2005; Sagoo et al., 2005), so solid manure heaps in Denmark have to be covered with either air-permeable or airtight covering material. However, a side-effect of such airtight covering is a reduction of the oxygen (O₂) concentration inside the heap, which may affect the production and emission of greenhouse gases.

Agriculture contributes significantly to greenhouse-gas emissions, primarily due to the emission of methane (CH₄) and nitrous oxide (N₂O). Scandinavian inventories have estimated that greenhouse-gas emissions from agriculture account for between 7% and 16% of total national emissions, of which manure handling contributes 11% (Rypdal, 2002). Development of effective manure-management technologies is thus crucial if countries are to meet their Kyoto Protocol commitments to reduce greenhouse-gas emissions (UNFCCC, 1999). Significant amounts of CH₄ and N₂O are emitted during the composting of solid manure (Hellebrand, 1998; Sommer and Møller, 2000; Hellebrand and Kalk, 2001a). Livestock manure may account for 6% to 10% of annual global emissions of CH₄ (Safely et al., 1992), whose global-warming potential has been estimated to be 21 times higher than that of CO₂ (Houghton et al., 1996). Nitrous oxide is produced by nitrification processes taking place in aerobic environments (Bremner and Blackmer, 1989) and by denitrification of nitrate in anaerobic environments (Hüther et al., 1997). Emission of both CH₄ and N₂O will therefore be influenced by the oxygenation level of the stored solid manure, which again can be expected to be influenced by covering. The present study was carried out to investigate how covering of heaps of solids separated from pig slurry affects production and emission of greenhouse gases and ammonia.

2. Materials and methods

2.1. Manure

Anaerobically digested pig slurry was separated using a decanting centrifuge (model NX 309, Alfa Laval Corp., Glinde, Germany). Thirteen tonnes of the solid fraction thus produced was immediately transferred into two identical cone-shaped heaps (height = 1.5 m, diameter = 4.8 m), which were stored for 4 months. The bulk density of the heaps was calculated to be 0.7 t m⁻³. One heap was stored uncovered (uncover) while the other was covered with a plastic sheet (0.15-mm polyethylene) (cover). To reduce inflow of air to the covered heap, the edges of the plastic sheet were covered with a sand layer. The impact of rain was simulated by weekly irrigation of the uncovered heap. Water was added in quantities corresponding to the registered weekly rainfall; however, to avoid leaching, addition did not exceed the absorbent capacity of the heap. The composition of manure was determined both before and after the storage period by analysing 8 representative samples per treatment. Each sample consisted of 10 subsamples of ca. 100 g each representatively taken by hand from the different regions of the heap. Each sample was thoroughly mixed in plastic bags to avoid volatilisation of NH₃ before ca. 100 g was analysed using standard techniques. Dry matter (DM) was determined after drying samples at 100 °C for 24 h, ash content by combustion analyses (Carbolite, EOF 71/250), total nitrogen (N) and total carbon (C) by the Dumas procedure (LECO, CN 2000, Michigan, USA), total ammoniacal N (TAN) with an Lachat Autoanalyser (DS/EN, ISO 11732), and pH of the samples, mixed with demineralised water (1:1 V:V), was measured with a standard electrode (Type 704, Metrohm, 1988). Nitrous oxide is another important greenhouse gas which has a global-warming potential 310 times higher than that of CO₂ (Houghton et al., 1996). Nitrous oxide is produced by nitrification processes taking place in aerobic environments (Bremner and Blackmer, 1989) and by denitrification of nitrate in anaerobic environments (Hüther et al., 1997).
Herisau, Switzerland). Total loss, during storage, of wet weight (ww), DM, ash, N, and C were determined by the differences of mass determined before and after the storage period.

2.2. Dynamic chambers

The heaps were stored in identical dynamic chambers (15.6 m × 7.3 m and 4.5 m tall). The concrete floor of each chamber sloped to a central drain to allow for collection of effluent. The emission chambers were equipped with internal ventilators to simulate wind and to ensure mixing of internal air. A ventilator situated in the ceiling of each chamber drew air through the chambers at a preset flow rate of 0.7 m$^3$ s$^{-1}$ allowing that chamber air was changed 4.9 times per hour; however, actual air flow was determined continuously throughout the measurement period by a pre-calibrated free-running impeller mounted in each ventilating duct. Samples of inflowing and outflowing air were transported through heated and insulated Teflon tubes (id = 3.0 mm) to a Photoacoustic Multi-gas Monitor (Model 1312, INNOVA, Copenhagen, Denmark), which measured concentrations of CO$_2$, CH$_4$, N$_2$O, NH$_3$, and air humidity every 10 min throughout the 120 d storage period. Emissions of the different gases were quantified using the following equation:

$$E = \sum_{j=1}^{n} (C_{\text{out}} - C_{\text{in}}) V_j$$

where $E$ is the total emission in mg of a specified gas, $j$ is the number of observations, $n$ is the total number of observations, $V$ is volume of air flow in m$^3$, and $C_{\text{out}}$ and $C_{\text{in}}$ are, respectively, the concentration of gas in outlet and inlet air in mg m$^{-3}$ air.

2.3. Composition of gas phase inside the bulk of the heaps

For determining CO$_2$, CH$_4$, N$_2$O, and oxygen (O$_2$) concentration inside the heaps, gas samples were collected from different depths in the heaps using a modification of the technique described by Petersen et al. (1998). Two flexible, but rigid plastic tubes ($l = 4.0$ m, id = 10 mm) were placed inside each heap at establishment. Both tubes were positioned 0.2 m above the base of the heaps, one 0.1 m below the surface of the heaps (surface), and the other in the centre of the heaps (centre). Air could diffuse into the tubes through 3.0-mm holes per cm length. Both ends of the tubes were connected with two 2-m lengths of gas-tight FEB tubes to gas-tight diaphragm pumps (Rietshle Thomas, type 5002, Maribo, Denmark), situated outside the heaps. This allowed circulation of air inside the closed tube systems prior to air sampling. With syringes, two samples of 50 ml were sucked from the tubes and transferred to 5-ml glass bottles fitted with butyl rubber septa. Sampling was performed on daily basis the first week after heap establishment, thereafter twice weekly throughout the storage period. When transferring a sample to a glass bottle, an extra needle was inserted through the rubber seal and the bottle was flushed with 35 ml of the gas from the syringes; then the needle penetrating the septa was removed and the 15 ml remaining in the syringes was injected into the glass bottle. Gas concentrations in the glass bottles were determined by means of gas chromatographic analyses (Hewlett-Packard 5890 Series II gas chromatograph, Minnesota, USA) equipped with an electron capture detector (ECD) detector for determination of N$_2$O concentrations, and a flame ionization detector (FID) for determination of CH$_4$ concentrations, and (Perkin Elmer, Clarus 500, Boston, USA) equipped with a thermal conductivity detector (TCD) for determination of O$_2$ and CO$_2$ concentrations. Temperature was automatically measured and recorded by Testostor data loggers (Model 175, Testo Inc., Wien, Austria) at 10-min intervals at the same depths as the sampling tubes, and in addition, 0.7 m below the heap surface (intermediate).

2.4. Statistical analysis

The influence of storage and storage treatment on manure composition was statistically analysed using the generalised linear model (GLM) procedure (SAS Institute, 1988). Assumption of equal variance of different groups was tested using Bartlett’s test prior to analysis. Where significant treatment effects within groups were found by analysis, Duncan’s multiple range test was used to test for significant differences of means. For all the statistics, a significance level of $\alpha = 0.05$ was applied. As production of gases inside heaps and emission from the heaps were not replicated, no statistical analyses were applied to gas concentration inside the manure heaps and to emission of gases from the differently treated manure heaps.
3. Results

3.1. Temperature and concentrations of O\textsubscript{2}, N\textsubscript{2}O and CH\textsubscript{4} inside the heaps

Immediately after establishment of the heaps the entire mass of stored manure was anaerobic (Fig. 1). In the uncovered heap, the O\textsubscript{2} concentration near the surface increased rapidly within 1–2 d, but did not reach ambient concentrations until 60 d; whereas O\textsubscript{2} concentrations remained low in the centre part throughout the measurement period, except for a periodic increase taking place after 90–120 d storage (Fig. 1a). In the covered heap, low concentrations of O\textsubscript{2} were observed both in the centre and near the surface until 90 to 120 d of storage, when O\textsubscript{2} concentrations temporarily increased (Fig. 1b).

Temperature inside the uncovered heap increased rapidly after establishment. Temperatures were observed to be highest near the surface of the heap; however, in this area temperature declined to near ambient temperature within 30 d of commencement of storage. In the intermediate and centre zones of the uncovered heap, temperatures increased to between 30 and 40 °C during the first weeks of storage and remained high thereafter (Fig. 2a).

In the covered heap, temperatures increased initially to ca. 30°C in all zones, but after 7–30 d temperatures gradually declined to ambient levels (Fig. 2b).

The concentration of N\textsubscript{2}O in the uncovered heap increased in the outer layers after 14 d of storage to reach maximum concentration after 20–40 d, where after N\textsubscript{2}O concentrations gradually declined to reach ambient levels after 40–120 d of storage (Fig. 3a). A minor increase in N\textsubscript{2}O concentrations in the centre of the uncovered heap took place after 60 d of storage. In contrast, no increase in N\textsubscript{2}O concentration was observed inside the covered manure heap (Fig. 3b).

Concentrations of CH\textsubscript{4} in the centre of the uncovered manure heap increased during the first 30 d of storage, after which they gradually declined toward ambient levels. The highest concentrations were observed in the central part of the heap (Fig. 4a). Methane concentrations inside the manure were substantially decreased by covering;

Fig. 1. Oxygen concentrations in solids separated from pig slurry when stored in an uncovered (a) and covered (b) heap. Concentrations were determined 0.1 m below heap surface — x —, in the centre of heaps — ▲ —, and in ambient air — —.

Fig. 2. Development of temperatures in different regions in solids separated from pig slurry when stored in an uncovered (a) and covered (b) heap. Temperatures were determined 0.1 m below heap surface — x —, 0.7 m below heap surface — ○ —, in the centre of the heap — ▲ —, and in ambient air — —.
however a minor increase in CH₄ concentration was observed in the centre of the covered heap toward the end of the storage period (Fig. 4b).

3.2. Storage loss of mass, carbon and nitrogen determined by mass balance

Uncovered material had a higher mass loss (water and organic matter) than covered material. The average pH of the material was reduced during storage either with or without covering, but the pH of the covered material after storage (pH 8.3) was significantly higher than that of the uncovered material (pH 7.1). (Table 1). Covering reduced storage loss of DM, whereas concentration of DM was unchanged by storage (Table 2). Ammonium (NH₄) concentration was significantly reduced by storage in an uncovered heap, while it was increased by 20% in a covered heap (Table 2). Approximately 30% of the initial carbon content was lost from the uncovered heap during storage, while only 7% of the initial carbon was lost from the covered heap (Table 3). Of the initial total nitrogen content 15% was lost from the uncovered heap, while only 6% was lost from the covered heap (Table 3).

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Weight, kg</th>
<th>PH</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Uncover</td>
<td>Cover</td>
</tr>
<tr>
<td>Start of storage</td>
<td>6540</td>
<td>6880</td>
</tr>
<tr>
<td>End of storage</td>
<td>5860</td>
<td>6620</td>
</tr>
<tr>
<td>Loss of initial content</td>
<td>680 (10.4%)</td>
<td>260 (3.8%)</td>
</tr>
</tbody>
</table>

Values in parentheses represent loss in per cent of initial weight. Values followed by same letter are not significantly different (P<0.05%).

3.3. Emission of N₂O, CH₄ and NH₃

The emission rate of N₂O from the uncovered heap peaked after 14 d of storage; thereafter the emission rate gradually decreased to 2 g N₂O–N d⁻¹ t⁻¹ after 120 d of storage. An increase in N₂O emission was observed following irrigation during the first 80 d of storage (Fig. 5a). The emission rate of CH₄ from the uncovered heap peaked after 30 d of storage, whereafter the
emission rate dropped to low levels after 40 d of storage (Fig. 5b). Ammonia emission from the uncovered heap was high for the first 20 d of storage. Covering the heap reduced the initial emission of NH₃; however, NH₃ was emitted at a low rate until 40 d after start of storage (Fig. 5c).

Emission rates of N₂O and CH₄ during storage of the separated manure solids were considerably reduced by covering the heap (Fig. 5). The cumulated emissions of CH₄ and CO₂ were reduced by 88% and 93%, respectively, by covering the heap (Table 3). For the uncovered heap, the carbon emission loss was almost equal to the carbon loss measured by mass balance, whereas it accounted for less than 30% in the covered heap. Ammonia emission was reduced by 12% by covering, while nitrogen loss via the N₂O pathway was reduced by 99%. Ammonia and N₂O emissions comprised approximately one-third of the total N loss from the uncovered heap (determined by mass balance analysis), but only 6% of the measured total nitrogen loss from the covered heap.

### 4. Discussion

#### 4.1. Aeration of heaps

Covering was found to have a major impact on composting activity and on the production and emission of greenhouse gases and ammonia. Similar effects of the combined compaction and covering of cattle manure heaps have been reported by Sommer (2001) and Chadwick (2005). The solid fraction produced by slurry separation is a fibrous product, which allows air to flow into the heap (Miller et al., 1989). Air inflow, and thereby oxygenation, may be further increased by the upward air flow generated by the heat produced by the composting process (Oenema et al., 2001). During passage through the material in the heap, the O₂ concentration of inflowing air is depleted by the microbial organisms degrading the readily digestible components of the manure (Miller et al., 1989). With depletion of the readily available digestible components, the aerobic zone moves deeper and deeper inside the heap over the course of the storage period, as illustrated in Fig. 1. For the first two months of storage, the

### Table 2
Concentration before and after storage, and storage loss of dry matter (DM) and ammonium nitrogen (NH₄–N) of the covered and the uncovered manure

<table>
<thead>
<tr>
<th>Units</th>
<th>DM</th>
<th>NH₄–N</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Uncover</td>
<td>Cover</td>
</tr>
<tr>
<td>Start of storage kg t⁻¹ (ww)</td>
<td>380 a</td>
<td>377 a</td>
</tr>
<tr>
<td>End of storage kg t⁻¹ (ww)</td>
<td>373 a</td>
<td>373 a</td>
</tr>
<tr>
<td>Loss kg t⁻¹ (initial ww)</td>
<td>45.9 (12.1%)</td>
<td>18.1 (4.8%)</td>
</tr>
</tbody>
</table>

Values in parentheses represent loss in per cent of initial content. Values followed by same letter are not significantly different (P < 0.05%).

### Table 3
Concentration before and after storage, and storage loss of nitrogen (N) and carbon (C) of the covered and the uncovered manure

<table>
<thead>
<tr>
<th>Units</th>
<th>Total N</th>
<th>Total C</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Uncover</td>
<td>Cover</td>
</tr>
<tr>
<td>Start of storage kg t⁻¹ (ww)</td>
<td>13.6 a</td>
<td>13.6 a</td>
</tr>
<tr>
<td>End of storage kg t⁻¹ (ww)</td>
<td>13.0 a</td>
<td>13.4 a</td>
</tr>
<tr>
<td>Loss of initial content kg t⁻¹ (initial ww)</td>
<td>2.0 (14.7%)</td>
<td>0.80 (5.6%)</td>
</tr>
<tr>
<td>NH₃–N or CH₄–C emission kg t⁻¹ (initial ww)</td>
<td>0.042 (0.3%)</td>
<td>0.037 (0.3%)</td>
</tr>
<tr>
<td>N₂O–N or CO₂–C emission kg t⁻¹ (initial ww)</td>
<td>0.65 (4.8%)</td>
<td>0.005 (0.04%)</td>
</tr>
<tr>
<td>Not accounted for kg t⁻¹ (initial ww)</td>
<td>1.3 (9.6%)</td>
<td>0.76 (5.3%)</td>
</tr>
</tbody>
</table>

Values in parentheses represent loss in per cent of initial content. Values followed by same letter are not significantly different (P < 0.05%).
centre of the uncovered heap was thus found to be partly anaerobic, while anaerobic conditions were only observed in the first week 0.1 m below the surface. Anaerobic conditions remained for a much longer period inside the bulk of the covered manure, as covering restricts inflow of air into manure heaps (Sommer, 2001).

4.2. Temperature

Composting has been defined as the degradation of organic matter performed by aerobic micro-organism when oxygen is available (Gray et al., 1971). This catabolic process produces energy that increases the temperature of the organic material, which in turn increases the microbial activity and thus the further degradation of the material. When the inflow of oxygen into the heap is reduced by covering, anaerobic degradation of organic matter takes place (Brock and Madigan, 1988). However, as the energy released by anaerobic degradation is less than that released by aerobic degradation, anaerobic degradation of solid manure does not increase heap temperature, which restricts microbial activity and thus the degradation of organic matter. Loss of total carbon during storage is therefore reduced by covering heaps of organic material.

4.3. Production and emission of methane

While CO₂ is the main end product of the aerobic degradation of organic carbon, the greenhouse gases CH₄ and CO₂ are the main end products of anaerobic carbon degradation (Brock and Madigan, 1988). The production rate of CH₄ is highly dependent on the temperature range (Husted, 1994; Khan et al., 1997). Due to the combination of anaerobic conditions and increasing temperature, concentration of CH₄ in the gas phase in the centre of the uncovered heap increased over the first 30 d of storage. The subsequent decline in CH₄ concentration, after one month of storage, indicates that aerobic conditions had moved toward the centre of the heap. The pattern of aeration and CH₄ production caused the emission of CH₄ to peak 30 d after establishment of the heap. The total emission of CH₄ from the uncovered heap accounted for 1.3% of the initial carbon content. Lower CH₄ emission has been reported by Osada et al. (2001), who found that CH₄ emission during storage of cattle deep litter represented 0.01% of total C; however, this low carbon loss via the CH₄ pathway might partly be explained by the shorter storage period (less than two months) and smaller heap size used, both of which reduced the size of the anaerobic zone inside the heap. Concentration of CH₄ in the bulk of the manure was reduced considerably by covering the heap, despite the fact that anaerobic conditions were observed inside the covered heap for most of the storage period. This indicates that CH₄ production inside the bulk of the covered manure was restricted by the lower temperature observed in covered material. Similar results have been observed by Husted (1994), who found that emission of CH₄ from dung heaps was reduced by between 2.7 and 10.3 times when heap
temperatures were decreased by 10°C. The emission of CH₄ from the covered, cooler heap was therefore reduced to 0.2% of the initial carbon content, which equals only 13% of the CH₄ emitted from the uncovered heap. The imbalance between mass of carbon emitted and the carbon loss determined by the mass balance method is expected to be caused by the inaccuracy of the mass balance method and by emission of carbon monoxide (CO) (Hellebrand and Kalk, 2001b).

4.4. Production and emission of nitrous oxide

In the uncovered heap, N₂O concentrations in the gas phase increased after 30 d of storage in the outer layers and after 60 d of storage at the centre of the heap. This indicates that the N₂O production zone moved toward the centre of the heap, together with the aerobic zone. N₂O production was negligible when air inflow was restricted by covering, which indicates that aeration of the covered material was too low to allow nitrification. Emission loss of N₂O from the uncovered heap peaked after 60 d of storage, declining gradually to low values after 120 d of storage. A similar pattern of N₂O emission has been observed by Hellebrand and Kalk (2001a) and by Sommer and Möller (2000). According to Hellmann et al. (1997), the delay in N₂O emission is caused by the fact that most nitrifying and denitrifying microorganisms are not thermophilic. Production of N₂O by nitrifying and denitrifying processes thus only takes place after heat production has diminished. The total nitrogen loss via the N₂O pathway accounted for almost 5% of the initial nitrogen content of the uncovered heap. Lower relative levels of N₂O loss have been observed by Sommer and Möller (2000) and Osada et al. (2001), who found that nitrogen loss from small heaps of dairy deep litter by the N₂O pathway amounted to 0.2% and 0.8%, respectively, of the initial total N content. Hellebrand (1998) modelled N₂O emission during the composting of green waste, and found that it accounted for approximately 0.5% of the initial total N content. The higher N₂O emission found in the present study may be explained by the higher density of the solids separated from slurry, relative to that of deep litter from dairy cattle production or of green manure. Higher density of solid manure reduces ammonia emission (Sommer, 2001), which increases the amount of ammonium available for nitrification. Thus, N₂O is emitted at a higher rate from solid manure heaps of higher density (Sommer and Möller, 2000). Covering the manure heap nearly eliminated the emission loss of N₂O to account only 0.2% of initial nitrogen content. This indicates that nitrification processes, and thereby denitrification processes, were restricted by low gas-phase oxygen concentrations within the covered material.

4.5. Ammonia emission

The emission of NH₃ from uncovered material was observed to be high immediately after heap establishment; thereafter, the emission rate gradually declined to nearly zero within a few weeks of storage. A lower initial NH₃ emission rate was observed from the covered heap; however, emission continued at a low rate for the first 60 d of storage. Total emission loss of NH₃ was thus only reduced by 12% by covering the heap. A higher reduction has been observed by Chadwick (2005), who found that a combination of plastic covering and compaction of cattle manure heaps reduced NH₃ emission by 90% in one study period, while insignificant reductions were observed during two other study periods. Covering poultry manure heaps with PVC sheets reduced ammonia emission by 90% (Sagoo et al., 2005). The storage loss of NH₃ from uncovered, separated solid manure was found to be lower than the storage loss of NH₃ from other types of solid manure, which, depending on manure type and climatic conditions, has been determined to account for 0.3–34% of the initial nitrogen content (Petersen et al., 1998; Sommer and Dahl, 1999; Möller et al., 2000b; Misselbrook et al., 2001; Webb et al., 2001; Rogstrand et al., 2005). The low ammonia emission loss found by the present study may be due to the high density of the solids separated from pig slurry (Sommer and Möller, 2000; Sommer, 2001). As well, the lower ammonia emission might partly stem from a more readily available carbon source in the solid manure produced by separation; the high straw content of cattle and pig solid manure results in a high content of less-degradable carbon sources, such as lignin and hemicelluloses. A high content of readily available carbon in the separated solid manure triggers high microbial activity and thus the immobilisation of ammonium in the microbial biomass, which restricts NH₃ emission (Fassen and Diek, 1979; Kirchmann, 1985).

4.6. Nutrient value

Covering reduced the total N loss from 15% to 6% of the initial nitrogen content, but the total N
concentration of the manure following storage (kg N t\(^{-1}\) ww) was not influenced by covering. This was due to the higher loss of water and dry matter during uncovered storage. Concentration of NH\(_4\) in uncovered manure was significantly reduced during storage by a combination of NH\(_3\) emission and denitrification processes, which together reduced the NH\(_4\) content by 46%. In contrast, the content of NH\(_4\)–N in the covered manure increased by 20%, indicating that net mineralisation of organically bound nitrogen took place in the covered heap during storage. NH\(_4\) is a readily available plant nutrient, so covering thus increases the nutrient value of separated solids. However, due to the higher pH and NH\(_4\) content of covered material, the benefits of covering may only be realized if the manure is rapidly incorporated following land spreading. Otherwise the ammonium conserved during storage may subsequently be lost as ammonia during land spreading (Hansen, 2004; Sagoo et al., 2005).

5. Summary

The oxygenation level inside heaps of solids separated from pig slurry was found to influence the production and emission of greenhouse gases during the storage period. Covering delayed aeration of the stored manure; this reduced internal heat production, degradation of organic matter, and emission of greenhouse gases and ammonia. Approximately 15% of the initial nitrogen content of the solids separated from pig slurry was found to be lost when the separated solids were stored uncovered. Almost 5% of the initial nitrogen content was found to be lost as N\(_2\)O, while only 0.3% was lost as NH\(_3\). Covering reduced the emission of N\(_2\)O by 99%, and the emission of NH\(_3\) by 12%. Of the initial carbon content, 28% was lost during uncovered storage; the majority of this loss was emitted as carbon dioxide (25%), while 1.3% was emitted as CH\(_4\). Covering was found to reduce CH\(_4\) emission by 88%. As the study was not replicated, no statistical significance can be placed on the quoted reductions in gaseous emissions.

Acknowledgements

This project was financially supported by the Danish Research Centre for Organic Farming and the Greenhouse Gas Emission Foundation. The authors thank Inge Marie Gregersen and Karsten Sørensen for their valuable technical assistance.

References


