Phosphorus Contents in the Profile of an Oxisol With Continuous Application of Liquid Swine Manure

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Received: January 29, 2020      Accepted: March 29, 2020      Online Published: April 15, 2020
doi:10.5539/jas.v12n5p112          URL: https://doi.org/10.5539/jas.v12n5p112

Abstract
This study aimed to evaluate the dynamics of phosphorus in the soil by the determination of phosphorus contents in an Oxisol cultivated with soybean during the 2016/2017 crop season, after 17 years of successive applications of liquid swine manure (LSM). The experiment was conducted in the experimental area of the University of Rio Verde. The treatments were: T1-control; T2-25 m³ ha⁻¹ of LSM; T3-50 m³ ha⁻¹ of LSM; T4-75 m³ ha⁻¹ of LSM; T5-100 m³ ha⁻¹ of LSM and T6-fertilization with mineral fertilizer. The subplots refer to soil depths: 0-10; 10-20 and 20-40 cm. The LSM was applied to the soil surface by spraying, before soybean sowing and the mineral fertilizer was applied to the furrow at the time of planting. After soybean harvesting, soil samples were collected at the studied depths for the determination of the contents of P in the soil. The contents of P in percolated water are within the concentration range considered adequate for human consumption. As for the contents of P in the soil, LSM applications on the soil surface increased the available soil P up to the depth of 20 cm. The highest dose of LSM applied to the soil promoted an increment of P contents that are within the safety range of the critical environmental limit of phosphorus (LCA-P), corresponding to 37% of this limit.

Keywords: environmental contamination, critical limit, organic residues, water quality

1. Introduction
For the agricultural use of Brazilian Cerrado soils, which are naturally acidic and with low fertility (Novais & Smyth, 1999), there is a need to use high doses of phosphorus (P) from phosphate fertilizers or organic sources, such as pig manure, which are abundant in intensive breeding regions. Phosphorus has a strong interaction with the soil, being strongly fixed in the colloids of the soil (Barrow, 1983), causing much of the P supplied in fertilization to be made unavailable in the soil solution due to the phenomenon of P fixation in reactions with the soil components. As a result of fixation, P becomes part of low solubility compounds, being less available for plant absorption. However, the saturation of the surface layer of the soil with this nutrient also favors its release to the solution and, consequently, its loss by surface runoff or leaching via vertical drainage of the soil (Sharpley, 2015; Gatiboni et al., 2015). Excess P contents increase the risk of losses to aquatic systems, an environment where its excess can cause eutrophication, compromising an entire watershed (Sharpley, 2015), which, consequently, can make water unsuitable for consumption.

In Rio Verde, state of Goias, Brazil, there are major production systems of the swine industry, generating large amounts of liquid swine manure (LSM). This residue contains nutrients, mainly nitrogen (N) and phosphorus (P), as well as organic matter (Choudhary, Bailey, & Grant, 1996). The application of this waste to soils via organic fertilization allows the use of its nutrients by crops in total or partial substitution with mineral fertilizers (Scherer, Baldissera, & Nesi, 2007; Lourenzi et al., 2014; Sousa et al., 2014).

An important feature of LSM is the low contents of dry matter, which makes transportation to distant areas difficult due to logistical and economic reasons. Thus, it tends to be applied to areas close to the place of origin of the waste (Basso, Ceretta, Durigon, Poletto, & Giroto, 2005).
The application of LSM follows specific legislations, which may originate from municipal laws, such as the county of Rio Verde-GO, which limits the application of LSM to 180 m³ ha⁻¹ (Menezes et al., 2017). This dosage allows the disposal of high concentrations of manure to the soil regardless of the requirement of the crop. Consequently, the nutrient concentrations in the soil increase, especially when the dose exceeds the demand of the crop. Moreover, the contents of these nutrients might be excessive at the application sites (Mahmood et al., 2017).

The N:P balance of LSM is different from the N:P requirement of the crops. Generally, LSM has an N:P ratio ranging from 2:1 to 6:1, while crops consume between 7:1 to 11:1 (Sadeghpour, Kellerings, Vermeylen, Godwin, & Czymmek, 2016). Therefore, medium to long-term applications of LSM can lead to an imbalance in the relationships between these nutrients, which might be aggravated when losses due to N volatilization occur (Sharpley & Halvorson, 1994; Sharpley, Daniel, Sims, & Pote, 1996). Prolonged and over dose applications can result in changes in the soil nutrient contents due to these imbalances (Scherer et al., 2007; Penha et al., 2015).

Unlike P, which is not lost by volatilization and, as a consequence, accumulates in the soil, much of the N from LSM applied to the soil is lost through volatilization within the first 24 hours after application (Rauber et al., 2017). In tropical soils, such as most of the Brazilian soils represented by Oxisols, P tends to remain in the superficial layers due to the soil mineralogical characteristics, which presents high concentrations of Fe and Al oxides, causing P to remain adsorbed in the system (Novais & Smyth, 1999). Thus, the nutrient has its content increased in the soil, especially in areas where the amount of P applied via LSM is higher than those exported by crops, resulting excessive P build up in the soil (Lewandowiski, Moncrief, & Drewitz, 2006).

According to Emídio (2012), phosphates can be present in water due to natural origins, from the natural result of the decomposition of biological matter and mineral leaching, and also due to anthropogenic actions, as a result of erosion of fertilized agricultural land and failures in wastewater treatment. In natural waters, the phosphorus content should not exceed 1.0 mg L⁻¹ of P₂O₅ (Alves, 2012).

In sloping areas, when there is excess P, there is a potential loss of nutrient due to runoff, configuring the risk to water quality, resulting in eutrophication of water bodies (Gatiboni, Brunetto, Kaminski, & Rheinheimer, 2008). In some European countries and in the United States where swine production is intense, the manure generated by this production system is the main cause of surface water eutrophication (Sharpley & Halvorson, 1994). Thus, regulatory pressures must exist to prevent the application of phosphate mineral waste or fertilizers in areas where there is excess P in the soil, since they exist in some regions of the country, as reported by Gatiboni, Smyth, Scmitt, Cassol & Oliveira (2015) for the state of Santa Catarina, in Brazil, and by Sharpley (2015) in the United States of America.

This study aimed to evaluate the potential risk of environmental contamination of P due to the successive use of liquid swine manure, using the critical environmental phosphorus (LCA-P) method and water percolation.

2. Materials and Methods

2.1 Study Area

The experiment was conducted in an experimental area belonging to the University of Rio Verde, located at the “Fontes do Saber” Farm, in the city of Rio Verde, state of Goias, Brazil. The soil of the area is classified as clayey dystroferric Red Latosol (Oxisol), with 610 g kg⁻¹ of clay and 4% of slope. The experiment has been conducted for 17 years of successive waste applications in soybean and corn crops rotation system.

2.2 Statistical Design

The study was conducted using a randomized block design in split plots, consisting of six fertilizations applied in the main plot and three soil depths in the subplot, with three repetitions. The fertilization treatments applied on the surface were: T1: control (without fertilization); T2: 25 m³ ha⁻¹ of LSM; T3: 50 m³ ha⁻¹ of LSM; T4: 75 m³ ha⁻¹ of LSM; T5: 100 m³ ha⁻¹ LSM and T6: recommended mineral fertilization dose, according to the soil and crop needs (Sousa & Lobato, 2004). The subplots refer to different evaluated soil depths: 0-10 cm; 10-20 cm and 20-40 cm.

The LSM (Table 1) was applied to the soil surface by one-time spraying (T2 to T5) prior to soybean sowing in a monoculture system and the mineral fertilization (T6) was applied to the furrow at the time of planting, at a dose of 380 kg ha⁻¹ of NPK with a formulation of 04.20.18.
Table 1. Average nutrient contents in LSM applied in the experimental area from 2000 to 2017. UniRV, Rio Verde, GO, Brazil

| Crop seasons | N total | P | K | Ca | Mg | S |
|--------------|---------|---|---|----|----|---|
| 2000 to 2017 | 1.1     | 0.3 | 0.9 | 0.48 | 0.2 | 0.09 |

| Crop seasons | OM | MS | Density | pH | Cu | Zn |
|--------------|----|----|---------|----|----|----|
| 2000 to 2017 | 1.0 | 1.24 | 1,006.5 | 7.8 | 2.3 | 8.0 |

The annual applied amounts of P as a function of the manure doses ranged from 21 to 83 kg ha⁻¹ (Table 2), whereas for the mineral fertilization, an equivalent of 33.2 kg ha⁻¹ of P was applied.

Table 2. Contents of P applied to the soil in the 2016/2017 crop season as a function of liquid swine manure (LSM) doses. UniRV, Rio Verde, GO, Brazil

| LSM doses (m³ ha⁻¹) | 25 | 50 | 75 | 100 |
|---------------------|----|----|----|-----|
| P (kg ha⁻¹)         | 21 | 42 | 63 | 83 |

Soybean sowing was carried out after the application of LSM on October 23, 2016, using a cultivar of high productive capacity (NS 6906 IPRO) with plant density of 16 plants m⁻¹, in no-tillage system, with a spacing of 0.5 m between lines. Management and phytosanitary treatments were carried out according to the technical recommendations (Embrapa, 2010) and crop needs.

2.2 Chemical Analysis

After soybean harvest, soil samples were collected from the depths of 0-10, 10-20 and 20-40 cm, using Augers for soil sampling. A composite sample consisting of three simple samples was collected in each plot (15 m × 10 m) in order to determine the soil P, sand, silt and clay contents. The determinations were performed in the Multiuse laboratories of the University of Rio Verde. The contents of P were determined by two methods: Mehlich-1 method (0.05 mol L⁻¹ HCl + 0.0125 mol L⁻¹ H₂SO₄), as described by Mehlich (1953), and the P water method (P-water). The P-water contents were quantified after weighing the equivalent of 1 cm³ wet soil, adding 10 mL of distilled water, and stirring the samples for one hour in an end-over-end shaker at 30 rpm. The samples were then centrifuged at 5,000 rpm (approximately 2,000 g) for 10 min, and the soil solution P was assessed following Murphy and Riley (1962). The contents of clay, silt, and sand were assessed using the pipette method (Embrapa, 1997).

The classification of the soil regarding the risks of P transferring to the soil solution was based on the P contents extracted by Mehlich-1 and the average soil clay content, following the methodology proposed by Gatiboni, Smyth, Scmitt, Cassol & Oliveira (2015), in which P limit and P critical environmental limit (LCA-P) were evaluated. The LCA-P is a tool designed to estimate the relative risk of P loss from the soil. The proposed models to calculate P-Limit (y, P-Mehlich 1 in mg dm⁻³ and x, clay content in%) and LCA-P (y, P-Mehlich 1 in mg dm⁻³ and x, content of clay in%) are obtained from Equations 1 and 2. The average soil clay content in the 0 to 10 cm deep layer was 61%.

\[ y = 54.3 + 1.18x \]  
\[ y = 40 + x \]

2.3 Statistical Analysis

All results were analyzed using analysis of variance (ANOVA). Significantly different means between treatments were separated with the Tukey’s test at the 0.05 probability level. The Scheffé test was applied to compare mineral application versus LSM application and, when relevant, regression analysis was performed, using Sisvar 4.6 software (Build 6.1) (Ferreira, 2019).

3. Results

Table 3 shows the summary of the analysis of variance for the evaluated parameters: soil P content, water P content, limit P content, LCA-P and clay, silt and sand contents.
Table 3. Summary of the analysis of variance of soil P, water P, limit P, LCA-P and clay, silt and sand contents. Rio Verde, GO, 2017

| VS                     | DF | P soil | P water | P limit | LCA-P | Clay  | Silt  | Sand  |
|------------------------|----|--------|---------|---------|-------|-------|-------|-------|
| Blocks                 | 2  | 3.406**| 6.796** | 11.749**| 13.019**| 13.019**| 10.241**| 0.056**|
| Fertilizing            | 5  | 920.833**| 1055.841**| 13.454**| 14.907**| 14.907**| 13.674**| 3.944**|
| Mineral vs Doses       | 1  | 182.537* | 0.448** | 34.098* | 37.781* | 37.781* | 7.170** | 5.633**|
| Doses                  | 4  | 1105.41**| 1319.689**| 8.293** | 9.189** | 9.189** | 15.3** | 3.522**|
| Error                  | 10 | 26.921 | 122.107 | 11.809  | 13.085 | 13.085 | 17.174 | 3.167 |
| Depths                 | 2  | 390.871**| 1344.130**| 3.627** | 4.019** | 4.019** | 9.463** | 11.167**|
| Fertilizing vs Depths  | 10 | 96.835**| 71.707** | 7.899** | 8.752** | 8.752** | 11.73** | 3.344**|
| (Mineral vs Doses) vs Depths | 2 | 26.162** | 247.515** | 12.638** | 14.004** | 14.004** | 21.115** | 1.94** |
| Doses vs Depths        | 8  | 114.503**| 27.756** | 6.714** | 5.189** | 5.189** | 9.383** | 3.372**|
| Error                  | 24 | 12.499 | 91.583  | 3.953  | 4.380  | 4.380  | 5.574 | 4.4814 |

CV (%) - 42.01 44.67 1.96 2.07 3.42 17.66 8.34

Note. VS = Variation Source; DF = Degrees of Freedom; CV = Coefficient of Variation. **, * significant at 1% and 5% probability, respectively by F test. ns = not significant.

For the variable P-water, there was no interaction between fertilization and depths. When the effect of doses was evaluated, the quadratic regression model presented the best fit (Figure 1) regardless of soil depth. The maximum point for P content in water was 0.034 mg L⁻¹ obtained at the dose of 19.84 m³ ha⁻¹, and from that dose there was a decrease of P content in water. Moreover, there were depth effects in relation to fertilization (Figure 2). The P-water contents were 0.015, 0.017 and 0.031 mg L⁻¹ at the depths of 0-10 cm, 10-20 cm and 20-40 cm, respectively.

![Figure 1. Contents of P in water as a function of liquid swine manure doses after application for 17 consecutive years of liquid swine manure (LSM), mineral fertilization and control in a dystroferric Red Latosol (Oxisol). Rio Verde, GO, 2017](image-url)
As for the contents of P in the soil, there was an interaction effect between fertilization and soil depths. When analyzing the soil P content as a function of the applied LSM doses at each depth, a quadratic regression adjustment was observed for the depths 0-10 cm and 10-20 cm. At the depth of 20-40 cm, there was no significant effect (Figure 3). At a depth of 0-10 and 10-20 cm, the soil P contents showed minimum points of 30.4 and 27.8 m\(^3\) ha\(^{-1}\), respectively. At higher doses, there was an increase in the soil P contents. Suitable soil P levels for clay contents ranging from 45 to 60% and higher than 60% are 8 to 12 mg dm\(^{-3}\) and 4.1 to 6.0 mg dm\(^{-3}\), respectively (Sousa & Lobato, 2004).

The application of LSM on the soil surface under the no-tillage system provided higher amounts of P than those exported by crops, especially at the 100 m\(^3\) ha\(^{-1}\) dose of LSM, increasing the available soil P levels to up to the depths of 20 cm. Therefore, there was no model adjustment for the depth of 20-40 cm.

Regarding the contents of soil P, P Limit and LCA-P, a contrast test was performed to compare the effect of mineral application versus LSM application (Table 4). The application of LSM provided higher soil P contents than the mineral fertilization, whereas for P Limit and LCA-P there were no differences.
It should be noted that mineral fertilization is based on crop needs and added nutrients, when soluble, are readily available to the plant, while LSM applications must go through the mineralization process to make them available. This factor might have influenced the higher availability of P in the soil with LSM application, since the P mineralization is only 60% (Ribeiro, Guimaraes, & Alvarez, 1999) in the current application, but the previous applications of LSM increase the P mineralization because of that residual effect.

Table 4. Contrasts of mean P in soil and P Limit under fertilization with doses of liquid swine manure (0; 25; 50; 75; 100 m$^3$ ha$^{-1}$) in relation to mineral fertilization (N-P-K)

| Treatments    | P soil | P Limit | LCA-P |
|---------------|--------|---------|-------|
| Mineral       | 4.30   | 101.87  | 101.44|
| Doses of LSM  | 9.24*  | 101.22ns| 100.76ns|

Note. * Significant by the F test.

Most of the mineral P applied to the soil was not available in the labile fraction in the soil when using the Mehlich-1 extractor. Less than 50% of the total applied P was recovered from the labile fraction to a depth of 40 cm. For doses of LSM up to 75 m$^3$ ha$^{-1}$, most of the available P was found below 10 cm depth. When 100 m$^3$ ha$^{-1}$ of LSM was applied, the effect of P accumulation over the 17 years of application was higher than 100% up to the depth of 20 cm (superficial layer) (Figure 4).

![Figure 4](https://example.com/figure4.png)

Figure 4. Percentage of available soil P (extracted by Mehlich-1) at the depths of 0-10, 10-20 and 20-40 cm as a function of P applied with different fertilizers (mineral, 25, 59, 65 and 100 m$^3$ ha$^{-1}$ of LSM) corresponding to 33.2, 7.5, 15, 22.5 and 30 kg ha$^{-1}$, respectively

4. Discussion

With the long-term application of LSM, there was an increase of P in the soil higher than 30 mg dm$^{-3}$ in the highest applied doses, which is considered high for soils with 61% of clay texture. This represents an increase of approximately 400%. High levels of soil P are often caused by a history of excessive manure application (Lewandowski, Moncrief, & Drewitz, 2006). The results indicate that P is accumulating in the soil, more on the surface, but the transfer to water by leaching is in minimal quantities, but not higher enough to cause potential problems (Figure 1). Although there is P build-up in the soil, it does not reach the problematic level at least in the first 17 years of application. Future research is needed to assess the impacts of manure application on soil in more chemical properties that can affect crop production and the environment.

When 100 m$^3$ ha$^{-1}$ of LSM was applied, the effect of P accumulation over the 17 years of application was greater than 100% up to the depth of 20 cm (superficial layer) (Figure 4). This result indicates that the amount applied at
this dose exceeds soybean P extraction by approximately 5.5 g P kg⁻¹ of harvested grain (Cunha, Casarin, & Prochnow, 2010). Thus, when manure is over applied during the soybean season, there is an excess of P when comparing with the crop P requirement. Ceretta et al. (2010) using 80 m³ ha⁻¹ of LSM broadcasted in the soil for seven years, also found that LSM increase P contents in the soil mainly in labile fractions.

Boitt et al. (2018), in a study of pig-slurry addition for fifteen years, found P accumulation and vertical movement down the soil profile, mainly to the 0-20 cm soil depth, in proportion to application rates when applying 100 and 200 m³ ha⁻¹ y⁻¹. These results are due to the application of manure that was performed superficially without incorporation and with the permanence of crop residues on the soil surface (Adeli, Bolster, Rowe, McLaughlin, & Brink, 2008; Mellek et al., 2010; Lou, Xu, Wang, Sun, & Liang, 2011; Lourenzi et al., 2011). The research carried out by Antonelli et al. (2019) shows the opinion of many other scientists who consider the application of LSM a non-sustainable soil management practice due to its negative effects leading to water pollution. Tabbara (2003) confirmed the potential loss of P only 24 hours after applying the manure, interpreted as a risk of transportation to the land surface through surface runoff and erosion, especially in slopes areas.

In unpolluted natural waters, P concentrations range from 0.010 mg L⁻¹ to 0.050 mg L⁻¹ (Ministry of Health, 2006). Therefore, the values determined in the present study at different depths are lower than the reference value (1.0 mg L⁻¹) for natural waters. In addition, these values are within the range of P levels in drinking water (< 0.050 mg L⁻¹). If long-term LSM inputs can increase quantities of P in soil, there is a potential to enhance the P-water, so critical levels of P in soils must be carefully monitored avoid P transfer to groundwater in the future or some soil conservationist practices should be adopted, as described by Labrière et al. (2015).

For safety reasons, it is recommended that the application of LSM or phosphate fertilizer should not be allowed after the suitable soil P level of soil, 8 to 12 mg dm⁻³ (Sousa & Lobato, 2004) is reached because this indicates high risk of soil becoming a source of P pollution to surface waters. After 17 years of successive applications of LSM in the same place, the dose of 100 m³ ha⁻¹ was considered excessive, because it increases the P contents in the soil even in deeper soil layers, indicating that it would be an unsustainable dose in areas where the application of swine manure is required.

The soil P-limit was 101 mg dm⁻³ (Table 4) from P-Mehlich-1 for soil with 61% clay, following the methodology proposed by Gatiboni, Smyth, Scmitt, Cassol, and Oliveira (2015). Further studies are being carried out in order to define a more accurate P-limit equation for specific soils of the Brazilian Cerrado, such as Oxisols. Sharpley, Mcdowell, Weld, and Kleinmann (2001) chased 75% of P-limit values to be maximum value in soils from the State of Pennsylvania-USA, beyond which measures restricting supplementation with P should be adopted.

5. Conclusions

Successive application of LSM and mineral fertilization does not influence the transfer of P to water after 17 years of successive applications. The contents of P in water at the depths of 0-10 cm, 10-20 and 20-40 cm are within the concentration range considered suitable for human consumption and was not aggressive to the environment.

As for the soil P content, soil surface LSM applications increased the available levels of soil P to a depth of up to 20 cm, especially with the highest dose of LSM.

The highest dose of LSM applied to the soil increased the P content in the soil within the safe range of the LCA-P, corresponding to 37% of this limit.

Acknowledgements

This research was part of the project Monitoring the environmental impact caused by the use of liquid swine manure in agriculture. The authors are grateful to the Fundação de Amparo à Pesquisa do Estado de Goiás (FAPEG), Conselho Nacional de Desenvolvimento Científico e Tecnológico (CNPq), Empresa Brasileira de Pesquisa Agropecuária (EMBRAPA), BRF-Brasil Foods and Universidade de Rio Verde (UniRV), all from Brazil, for financial support and scholarships.

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