Original Research Article

PRODUCTION OF KUNAPAJALA, ITS NUTRITIONAL CONTRIBUTIONS, MICROBIAL AND PESTICIDE EFFECT

Abstract

A study on nutritional and microbial analysis of Kunapajala with different storage time interval was conducted in the Department of Soil Science & Agricultural Chemistry and Department of Plant Pathology, UBKV, Coochbehar-736165, West Bengal during March, 2019. Motive of this work was to estimate the physico-chemical properties, macro and micro nutrient content and various microbial load of Kunapajala with different storage time interval. Kunapajala had highest P, K, Ca, Mg, Fe, Zn, Cu & Mn 40 days after preparation and it had highest N and S 20 days after preparation. It had highest beneficial microbial load of Fungi, *Actinomycetes, Pseudomonus*, Phosphorus Solubilising Bacteria (PSB), *Azotobacter, Azospirillum, Rhizobium* and *Trichoderma* 40 days after preparation. So, continuous foliar and soil application of Kunapajala from 20 days after preparation to 40 days after preparation was beneficial to get maximum utilization. Moreover, Kunapajala can be used as an alternative against chemical fertilizers and pesticides to develop organic farming.

Key words: Kunapajala, Liquid organic manure, organic farming, Organic pesticide, organic fertilizer.

1. Introduction

India faced several famines in its history and these famines claimed millions of life. In the famine of 1943, India lost around four million lives in eastern India alone (Dyson and Maharatna 1991). To solve that situation and to become self-sufficient in food production Govt. Of India launched several scientific ventures. Ultimately in late 1960's India became self-sufficient in food through green revolution. The success of green revolution mainly relied on the heavy use of chemical fertilizers, pesticides, high yielding varieties and modern mechanical agricultural instruments (FAO, 2009). In contrast, modernization of agriculture and dependency on chemical fertilizers and pesticides gradually deteriorates the soil fertility and adversely affects the ecological balance, natural biodiversity and environment (Paull, 2011). Adaptation of organic agriculture is the only way to solve this problem (Manna *et al.*, 2005). The procedure of preparation of Kunapajala was mentioned in Vrikshaurveda written by Surpala. According to verse 101, 102, 103 and 104 of Vrikshaurveda, it could be prepared by mixing excreta, marrow of the bones, flesh, brain and blood of the boar with water. After that the mixture should be boiled and stored in a iron pot after adding sufficient quantity of

husk, sesame oil cake, honey, black gram and ghee. As per availability, the blood, flesh and marrow of fish, goat or other animals could be used for the preparation. The items should be taken at random, no specific proportion is mentioned. Verse 106 of Vrikshaurveda explained that Kunapajala was highly effective for the crop plants. A significant increase in production was observed due to spraying of Kunapajala in several crop plants including mango (Mangifera indica), Soapnut (Sapindus emarginatus), Coconut (Cocos nucifera), kiwi fruit (Actinidia deliciosa) and bringal (Solanum melongana). Spraying of Kunapajala on tea bushes controlled the attack of tea mosquito bug (Helopeltis theivora) and loopers (Biston suppressaria). Narayanan (2006) reported that after spraying Kunapajala (made of rat flesh, Musika kunapa) rats were totally disappeared from tea garden. So it can also be used as an alternative against chemical pesticides and rodenticides (Ayangara, 2004a, 2004b, 2005, 2006a, 2006b) (Narayanan, 2006) (Bhat and Vasanthi 2008). Hence, my motive of this research is to observe the Physical, nutritional and microbial properties of the Kunapajala with different storage time intervals. Due to several microbial interactions, the nutritional status of Kunapajala is continuously changing. Studying the nutritional content and microbial population with different time interval, we can understand the potentiality of Kunapajala in different time intervals. According to that proper spraying schedule of Kunapajala should be recommended to the farmers for maximising crop yield.

2. Materials and Methodology

2.1. Preparation of Kunapajala

Ingredient: Bombay Duck fish (*Harpedon neherus*, cheap, devoid of scales and easily decomposable)(2.5 kg), Powdered sesame oil cake (1 kg), Rice husk (1 kg), Molasses (1 Kg), Jersey cow urine (7.5 litres).

Procedure: All these ingredients were mixed in an earthen pot, closed the container and allowed them to ferment. Stirring twice in a day should be done in both the directions. After 40 days the solution should be filtered and had to be collected (Sarkar *et al.*, 2014).



Kunapajala- Fermentation State and liquid extract after filtering

2.2. Nutritional and microbial analysis of Kunapajala

The physical, nutritional and biological parameters of Kunapajala were analysed on the day of preparation (0 days), 20 days after preparation and 40 days after preparation using

scientifically approved standard procedures. The standard procedures performed for the estimations of these parameters are described in Table-1 and Table-2.

Table-1. Physical and chemical properties of Kunapajala

| Sl. | Parameters | Methods | Reference |
|-----|----------------------|--|----------------|
| No. | | | |
| 1 | Colour | Visual evaluation | |
| 2 | Odour | Sensory evaluation | |
| 3 | Mould Growth | Visual evaluation | |
| 4 | Maggot Population | Visual evaluation | |
| 5 | pН | pH meter method | Jackson (1973) |
| 6 | EC | Conductivity meter method | Jackson (1973) |
| 7 | Organic Carbon (OC) | Walkley and Black wet digestion | Walkley and |
| | | | Black (1934) |
| 8 | Total Nitrogen | Microkjeldhal method | Jackson (1973) |
| 9 | Total Phosphorus | Total Phosphorus Nitric-Perchloric(9:4) digestion and Jack | |
| | | colorimetry using vanado-molybdo | |
| | | phosphoric yellow colour method | |
| 10 | Total Potassium | Nitric-perchloric(9:4) digestion and flame | Jackson (1973) |
| | | photometry | |
| 11 | Total Calcium | Nitric-perchloric(9:4) digestion and AAS | Jackson (1973) |
| 12 | Total Magnesium | Nitric-perchloric(9:4) digestion and AAS | Jackson (1973) |
| 13 | Total Sulphur | Total Sulphur Nitric-perchloric(9:4) digestion and | |
| | | Turbidimetry | Cornfield |
| | | | (1963) |
| 14 | Total Micronutrients | Nitric-perchloric(9:4) digestion and AAS | Jackson (1973) |
| | Fe, Mn, Zn ,Cu | | |

Table-2. Biological properties of Kunapajala

| Sl. | Parameters | Methods | Reference |
|-----|---------------|---------------------------------------|----------------------------------|
| No. | | | |
| 1 | Bacteria | Nutrient Agar medium | Atlas and Parks (1993) |
| 2 | Fungi | Martin's rose Bengal Agar | Martin (1950) |
| 3 | Actinomycetes | Ken knight's Agar medium | Cappuccino |
| | | | and Sheman (1996) |
| 4 | PSB | Pikovskaya's medium | Sundara and |
| | | | Sinha (1963) |
| 5 | Azospirilum | Nitrogen free Bromothymol blue medium | Dobereiner <i>et al.</i> ,(1976) |
| 6 | Azotobacter | Jensen's medium | Jensen (1942) |
| 7 | Trichoderma | Trichoderma specific Medium | Saha and Pan (1997) |
| 8 | Pseudomonus | King's B Agar medium | King et |
| | | | al.,(1954) |

| 9 | Rhizobium | Yeast extract Mannitol Agar with Congo | Fred et |
|---|-----------|--|------------|
| | | red | al.,(1932) |

3. Results and Discussion

Table-3. Physical and Physico-chemical parameters of Kunapajala

| KUNAPAJALA | | | |
|-------------------|------------------------------------|---------------------------|------------------------------|
| Parameters | On the day of preparation (0 days) | 20 days after preparation | 40 days after preparation |
| Colour | Light brownish orange | Brownish orange | Dark brownish orange |
| Odour | Mild alcoholic smell | Foul alcoholic smell | Extreme foul alcoholic smell |
| Mould growth | No mould growth | Heavy mould growth | No mould growth |
| Maggot Population | No maggot found | Heavy maggot growth | No maggot found |
| pН | 6.74 | 3.47 | 8.81 |
| EC (ds/m) | 2.55 | 9.72 | 8.57 |
| Total OC (%) | 1.72 | 2.55 | 4.18 |

The colour of freshly prepared Kunapajala was brownish orange and it became darker from the 20 days onwards. As the storage period progressed, the preparation became darker in colour without much significant change. Through anaerobic respiration, several gases were produced and that cause natural liquids and liquefying tissues. They also caused build-up of pressure combined with the loss of integrity of the skin and ultimately the tissue was ruptured. Ruptures in the skin allowed oxygen to re-enter the tissue and provide more surface area for the development of fly larvae and the activity of aerobic microorganisms. For these activities dark brownish orange colour was developed (Janaway *et al.*, 2009; Carter *et al.*, 2008).

Fresh preparation of Kunapajala possessed a foul alcoholic smell. Extreme foul odour was observed from 20 to 40 days onwards. Foul alcoholic odour was developed due to putrefaction. Anaerobic metabolism took place, leading to the accumulation of gases, such as hydrogen sulphide, carbon dioxide, methane, cadaverine, putrescine and nitrogen. The purging of gases and fluids resulted the strong distinctive odours (Carter *et al.*, 2008; Payne, 1965).

Initially there was no mould growth in Kunapajala whereas it was first observed 5 days after preparation. Mould growth was observed on the liquid surface and also on the sides of the storage vessel from 15 days onwards, the decrease in mould growth was observed in 20 days and was completely absent in 25 days. Fungi consumed energy or food from the decaying tissue and enhanced the decomposition process. Fungi were abundant in the environment. From air or from any other source they might be appeared in the Kunapajala vessel. But when tissues became totally liquefied or almost decomposed their population started declining. It

was due to unavailability of food from that decaying tissue (Hawksworth *et al.*, 2011) (Schwarz *et al.*, 2015) (Hitosugi *et al.*, 2006).

During decomposition, at initial stages Kunapajala attracted flies and these flies laid eggs on it. From those eggs maggots were developed. Young maggots spread throughout the container and took food from the decaying tissue. Due to the activity of maggots the tissue started decomposing faster and the bacterial activity also enhanced. This was the reason behind the heavy development of maggots in Kunapajala after 5 days of its preparation. After 25 days of its preparation due to loss of readily available cadaveric material, maggot population drastically reduced (Anderson, 2000), (Fuller, 1934) (Morovic-Bodac, 1965), (Carter and Tibbett 2008), (Janaway *et al.*, 2009)

On the day of preparation Kunapajala showed of pH (6.74) and after 20 days it became highly acidic in nature (3.47). Then after 40 days it became alkaline in nature (8.81). Animal tissue decomposition initially created an alkaline environment and due to microbial activity it became acidic after 20 days. When decomposition was totally completed, it became alkaline again (Carter, 2005; Hopkins *et al.*, 2000; Rodriguez and Bass 1985) (Gill-King, 1997; Towne, 2000). Similar results were also found by Anandan *et al.*, (2016), Jani *et al.*, (2017) and Ankad *et al.*, (2017) in Kunapajala.

Kunapajala showed highest EC 20 days after preparation (9.72 ds/m) due to high acidic nature of the solution and after that it started declining (8.57 ds/m, 40 days after preparation). On the day of preparation it showed lowest EC (2.55 ds/m) (Carter, 2005; Hopkins *et al.*, 2000; Rodriguez and Bass 1985) (Gill-King, 1997; Towne, 2000). Anandan *et al.*,(2016) and Ankad *et al.*,(2017) also concluded similar trend and results.

Total OC (organic carbon) was highest 40 days after preparation (4.18%) and on the day of preparation it showed minimum value (1.72%) in Kunapajala. In decomposition physical breakdown and biochemical transformation of complex organic molecules occurred, due to that several organic carbon compounds were synthesized (Juma, 1998). This was reason for continuous increase of OC in Kunapajala. Anandan *et al.*, (2016) noticed similar trend of OC and results in his experiment.

Physical and physic-chemical parameters of Kunapajala were mentioned in Table-3.

The highest N content was recorded 20 days after preparation in Kunapajala (7238 mg/dm³) while on the day of preparation it recorded the lowest value (3486 mg/dm³). For the activity of bacteria and maggots, Kunapajala started decomposing faster and due to that N content of the Kunapajala was in an increasing trend. But after 20 days 9-44% of the N was volatized in the form of Ammonia from the solution due to alkalinity of the Kunapajala solution on that moment (Kirchmann and Witter 1989). Ankad *et al.*, (2017) and Jani *et al.*, (2017) also concluded similar trend and results in their experiment.

On the day of preparation Kunapajala recorded the lowest value (208.661 mg/dm³) of P and 40 days after preparation it recorded the highest value (517.717 mg/dm³) of P. Kunapajala contained animal tissue and animal tissues had high P content. According to Tian *et al.*,

(1992), organic matters high in P decompose faster and release P significantly. So, Kunapajala had increasing tread of P content during decomposition. Ankad *et al.*, (2017) and Jani *et al.*, (2017) also analysed the P content of Kunapajala and found similar results.

K content was lowest on the day of preparation (890.396 mg/dm³), after that it was gradually increased and reached the highest value at 40 days after preparation (1873.543 mg/dm³). Activity of fungus and other microorganisms was the reason behind continuous release of K up to 40 days (Carter *et al.*, 2007).

Highest Ca content was observed 40 days after preparation (614 mg/l) and on the day of preparation it was the lowest (376 mg/l). Excessive fungus and microbial activity was the reason for continuous release of Ca up to 40 days (Carter *et al.*, 2007).

On the day of preparation Mg content was the lowest (56 mg/l) and after 40 days Mg content recorded the highest value (88 mg/l). Fungal and microbial activity was the main cause behind gradual release of Mg in Kunapajala (Carter *et al.*, 2007).

S content was lowest on the on the day of preparation (678 mg/l) and after 20 days it recorded the highest value (857 mg/l) but then S content started declining. Due to excessive volatile release of hydrogen sulphide, after 20 days S content started declining (Carter *et al.*, 2007).

Highest Fe content was recorded 40 days after preparation (72 mg/l) and on the day of preparation it was the lowest (55 mg/l). Due to fungal and bacterial activity gradual release of Fe was noticed in Kunapajala (Dent *et al.*, 2004).

On the day of preparation Zn content was minimum (6.78 mg/l) and 40 days after preparation it became maximum (17.75 mg/l). Gradual increase of Zinc content was noticed in Kunapajala due to activity of fungus and bacteria (Hodson *et al.*, 2001, Kearney *et al.*, 2000 and Deydier *et al.*, 2003).

Cu content was maximum 40 days after preparation (8.53 mg/l) and on the day of preparation it recorded lowest value (4.76 mg/l). Continuously increasing trend of Cu content was observed due to activity of several fungal and bacterial species (Hodson *et al.*, 2001, Kearney *et al.*, 2000 and Deydier *et al.*, 2003).

Highest Mn content was noticed 40 days after preparation (2.06 mg/l) and on the day of preparation the Mn content recorded the lowest value (0.58 mg/l). Heavy microbial interaction or activity inside Kunapajala might be the reason behind this trend and result.

The macro and micro nutrient content of Kunapajala was mentioned in Table-4.

Table-4. Macro and micro nutrient content of Kunapajala

| KUNAPAJALA | | | |
|----------------------|------------------------------------|---------------------------|---------------------------|
| Parameters | On the day of preparation (0 days) | 20 days after preparation | 40 days after preparation |
| N mg/dm ³ | 3486 | 7238 | 4690 |
| P mg/dm ³ | 208.661 | 296.260 | 517.717 |

| K mg/dm ³ | 890.396 | 1589.994 | 1873.543 |
|----------------------|---------|----------|----------|
| Ca (mg/l) | 376 | 452 | 614 |
| Mg (mg/l) | 56 | 73 | 88 |
| S (mg/l) | 678 | 857 | 719 |
| Fe (mg/l) | 55 | 67 | 72 |
| Zn (mg/l) | 6.78 | 13.63 | 17.75 |
| Cu (mg/l) | 4.76 | 7.44 | 8.53 |
| Mn (mg/l) | 0.58 | 1.27 | 2.06 |

Fungi population was highest 40 days after preparation (33 x 10^8 cfu/ml) and it was lowest on the day of preparation (4 x 10^4 cfu/ml). This gradual increasing trend was noticed due to enhanced activity of early stage fungi ascomycetes, deuteromycetes and saprophytic basidiomycetes and late stage fungi ectomycorrhizal basidiomycetes in Kunapajala with time (Carter et al., 2003).

On the day of preparation Kunapajala recorded lowest *Actinomycetes* population (3 x 10^3 cfu/ml). After that it increased continuously and reached highest at 40 days after preparation (5 x 10^8 cfu/ml). Continuous decomposition of complex mixture of polymers in dead animal tissues was the prime reason for continuous development of *Actinomycetes* population in Kunapajala (Goodfellow and Williams 1983, McCarthy and Williams 1992, Stach and Bull 2005).

The highest population of *Pseudomonus* was noticed 40 days after preparation (13 x 10^{10} cfu/ml) in Kunapajala and on the day of preparation it recorded the lowest (5 x 10^3 cfu/ml). This type of increasing trend up to 40 days in Kunapajala was also concluded by Ali *et al.*, (2012).

PSB population was highest on the day of preparation $(2 \times 10^5 \text{ cfu/ml})$ and it became maximum at 40 day after preparation $(21 \times 10^{10} \text{ cfu/ml})$ in Kunapajala. Similar trend of population growth was also observed by Ali *et al.*, (2012) in Kunapajala.

On the day of preparation *Azotobacter* population had lowest value $(7 \times 10^4 \text{ cfu/ml})$ in Kunapajala and after 40 days it became the highest $(13 \times 10^{12} \text{ cfu/ml})$. Presence of *Azotobacter* in Kunapajala and this type of growth trend was justified by Ali *et al.*, (2012).

Highest *Azospirilum* population was noticed 40 days after preparation (13 x 10^{10} cfu/ml) and on the day of preparation lowest value was found (11 x 10^3 cfu/ml). Ali *et al.*, (2012) approved the existence of *Azospirilum* in Kunapajala and its growth behaviour in it.

Lowest *Rhizobium* Population was found on the day of preparation $(2 \times 10^3 \text{ cfu/ml})$ and after 40 days highest Rhizobium population $(4 \times 10^{11} \text{ cfu/ml})$ was noticed in Kunapajala. Ali *et al.*, (2012) also concluded similar trend of population growth of *Rhizobium* in Kunapajala.

Trichoderma population was highest 40 days after preparation (21 x 10^8 cfu/ml) in Kunapajala and on the day of preparation it had the lowest population (6 x 10^3 cfu/ml). *Trichoderma* had significant contribution in decomposition and biodegradation of organic

matters and due to that the population of *Trichoderma* in Kunapajala had an continuous increasing trend up to 40 days (Woo *et al.*, 2004).

Microbial population of Kunapajala was mentioned in Table-5.

Table-5. Microbial population of Kunapajala

| KUNAPAJALA | | | |
|-----------------------|------------------------------------|---------------------------|---------------------------|
| Parameters | On the day of preparation (0 days) | 20 days after preparation | 40 days after preparation |
| Fungi (cfu/ml) | 4×10^4 | 16×10^7 | 33 x 10 ⁸ |
| Actinomycetes(cfu/ml) | 3×10^{3} | 6 x 10 ⁴ | 5×10^8 |
| Pseudomonus (cfu/ml) | 5×10^{3} | 8×10^{10} | 13×10^{10} |
| PSB(cfu/ml) | 2×10^{5} | 15 x 10 ¹⁰ | 21×10^{10} |
| Azotobacter (cfu/ml) | 7×10^4 | 9 x 10 ¹² | 13 x 10 ¹² |
| Azospirilum (cfu/ml) | 11×10^3 | 8 x 10 ⁸ | 13×10^{10} |
| Rhizobium (cfu/ml) | 2×10^{3} | 6×10^6 | 4×10^{11} |
| Trichoderma (cfu/ml) | 6×10^{3} | 18×10^8 | 21×10^8 |

4. Conclusion

The study concludes that Kunapajala has high nutrient content and beneficial microbial population. Nutrient content of Kunapajala is highly influenced by its microbial population. Fungi will help to breakdown complex organic compounds and produce simple organic and inorganic compounds useful for plants. Azotobacter, Azospirilum and Rhizobium help to fix more N in crop field. PSB enhance the P solubilisation in crop field. Actinomycetes help to decompose complex organic molecules and antagonistic potential of Pseudomonus and Trichoderma will help to protect the crop from soil-borne diseases. Microbial population is continuously increasing and it become highest after 40 days. So, application of Kunapajala after 40 days is beneficial for crops. But N and S content of Kunapajala is highest 20 days after preparation. So to exploit that spraying of Kunapajala after 20 days is also recommended. Spraying of Kunapajala on the day of preparation is not recommended because on that day microbial population and nutrient content was minimum and most of the organic matter is not properly decomposed, so they will not be highly available for the cop plants. So foliar and soil application of Kunapajala from 20 days of its preparation to 40 days of its preparation is recommended for the crop and soil because we can utilise its total potential. The ingredients required to prepare it is easily available and cheap comparing with chemical fertilizers and pesticides. The crops produced using Kunapajala will be free from any harmful chemical residues. So, it is healthy for the consumer. Moreover use of Kunapajala instead of chemical fertilizer and pesticide is highly useful to increase the crop yield, soil productivity and farmer's income.

5. References

Ali, M. N., 2012. Sustainable Agriculture with Sustainable Agriculture with Sustainable Agriculture with Low Cost Technologies (SALoCT). A project funded by Rural Technilogy action Group – Eastern India (RuTAG-EI), IIT Kharagpur, under DST, Govt. of India.

Anandan, R., Priya, L. and Rajendran, P., 2016. Dynamics of Organic Biofertilizers on *Oryza sativa* ADT-43. *Int. J. Curr. Microbiol. App. Sci*, **5**(4): 902-908.

Anderson, G. S., 2000. Minimum and maximum development rates of some forensically important Calliphoridae (Diptera). *Journal of Forensic Sciences*, **45**:824-832.

Ankad, G. M., Hiremath, J., Patil, R.T and Pramod, H.J., 2017. Nutrient analysis of Kunapajala and Panchagavya and their evaluation on germination of Ashwagandha (*Withania somnifera Dunal.*) and Kalamegha (*Andrographis paniculata Nees*) seeds: a comparative study. Journal of Ayurveda and Integrative Medicine, **xxx**: 1-7.

Atlas, R. M. and Parks, L. C., 1993. Handbook of microbiological media, *CRC Press, Inc.* London, 529p.

Ayangarya, V. S., 2004a. Herbal kunapa. Asian Agri-History, 8:315–317.

Ayangarya, V. S., 2004b. Manujala: A liquid manure. Asian Agri-History 8:319–321.

Ayangarya, V. S., 2005. INDSAFARI – An organic pesticide for tea. *Asian Agri-History*, **9**:317–319.

Ayangarya, V. S., 2006a. Mushika kunapa. Asian Agri-History, 10:157–159.

Ayangarya, V. S., 2006b. Kiwifruit plant treatment on the Himalayas of India: A Vrikshayurveda experience. In: Bridging Gap Between Ancient and Modern Technologies to Increase Agricultural Productivity: Proceedings of the National Conference held from 16-18 December 2005, Central Arid Zone Research Institute, Jodhpur 342 003, Rajasthan, India. (Choudhary, S.L., Saxena, R.C., and Nene, Y.L., eds.). Asian Agri-History Foundation, (AAHF), Secunderabad, India; and Rajasthan Chapter of AAHF, Udaipur, India. 2006. pp. 102-103.

Cappuccino and Sheman., 1996. Microbiology – A laboratory manual (4th ed), *The Benjamin/ Cummings Publishing Company Inc.*, 213p.

Carter, D. O., Yellowlees, D. and Tibbett, M., 2007. Cadaver decomposition in Terrestrial Ecosystems. *Naturwissenschaften*, **94**(1): 12-24.

Carter, D., 2005. Forensic taphonomy: Processes associated with cadaver decomposition in soil. Ph.D. thesis, James Cook University, Townsville, Australia.

Carter, D.O. and Tibbett, M., 2008. Cadaver Decomposition and Soil: Processes". In M. Tibbett; D.O. Carter (eds.). *Soil Analysis in Forensic Taphonomy*, CRC Press. pp. 29–51.

Carter, David, O., Tibbett, Mark., 2003. Taphonomic Mycota: Fungi with forensic potential. *Journal of Forensic Sciences, Blackwell.* **48**(1): 168-171.

Dent, B. B., Forbes, S. L., Stuart, B. H., 2004. Review of human decomposition processes in soil. *Environmental Geology*, **45**(4): 576-585.

Deydier, E., Guilet, R., Sarda, S. and Sharrock, P., 2005. Physical and chemical characterisation of crude meat and bone meal combustion residue: "Waste or raw material?" *Journal of Hazardous Materials*, **121**(1-3): 141–148.

Dobereiner, J., Marriel, I. E. and Nery, M., 1976. Ecological distribution of Spirillum lipoferum, Beijerinck. *Can. J. Microbiol.*, **22**: 1464-1473.

Dyson, T. and Maharatna, A., 1991. Excess mortality during the Bengal famine: a re-evaluation. *Indian Economic and Social History Review*, pp-281-297.

FAO., 2009. Rapid growth of selected Asian economies. Available from http://www.fao.org/docrep/009/ag087e/ AG087E05.htm.

Fred, E. B, Baldwin, I. L. and McCoy, F., 1932. Root Nodule Bacteria and Leguminous Plants. *University of Wisconsin Press*, Madison, Wisconsin.

Fuller, M. E., 1934. The insect inhabitants of carrion: A study in animal ecology. *Council for Scientific and Industrial Research*, Bulletin No.-82, pp-63.

Gill-King, H., 1997. Chemical and ultra-structural aspects of decomposition, in *Forensic Taphonomy: The Postmortem Fate of Human Remains* (W. D. Haglund and M. H. Sorg, Eds.). Boca Raton, FL: CRC Press, 93–108.

Goodfellow, M. and Williams, S. T., 1983. Ecology of Actinomycetes. *Annual Review of Microbiol.* 37: 189-216.

Hawksworth D.L. and Wilthshire, P. E. J., 2011. Forensic mycology: the use of fungi in criminal investigations. *Forensic Sci Int*, **206**:1-11.

Hitosugi, M., Ishii, K., Yaguchi, T., Chigusa, Y., Kurasa, A., Kido, M., Nagai, T., Tokudome, S., 2006. Fungi can be a useful forensic tool. *Leg Med*, **8**:240-242.

Hodson, M. E., Valsami-jones, E., Cotter-howells, J. D., Dubbin, W. E. and Kemp, A. J., 2001. Effect of bone meal (calcium phosphate) amendments on metal release from contaminated soils - a leaching column study. *Environmental Pollution*, **112**:233–243.

Hopkins, D. W., Wiltshire, P. E. J. and Turner, B. D., 2000. Microbial characteristics of soils from graves: An investigation at the interface of soil microbiology and forensic science. *Appl. Soil Ecol.* **14**: 283–288.

Jackson, M. L., 1973. Soil Chemical Analysis. *Prentice Hall of India Pvt. Ltd*, New Delhi, pp.498.

Janaway, R. C., Percival, S. L. and Wilson A.S., 2009. "Decomposition of Human Remains". In Percival, S.L. (ed.). *Microbiology and Aging. Springer Science + Business*. pp. 13–334.

Jani, S., Prajapati, P.K., Harisha, C.R and Patel, B.R., 2017. Kunapajala liquid organic manure: Preparation and its quality Parameters. *World Journal Of Pharmacy and Pharmaceutical Sciences*, Volume **6**:1989-2000.

Jensen, H. L., 1942. Nitrogen fixation in leguminous plants. General characteristics of root nodule bacteria isolated from species of Medicago and Trifolium in Australia. *Proc. Linn. Soc. N.S.W.* **66**:98-108.

Juma, N.G., 1998. The pedosphere and its dynamics: a systems approach to soil science. Volume 1. Edmonton, Canada, *Quality Color Press Inc.* 315 pp.

Kearney, T., 2000. Remediation of Toxic Metal Pollution in Soil Using Bone meal Amendments. *Environment Agency*.

King, E.O., Ward, M.K., Raney, D.E., 1954. Two simple media for the demonstration of payociamin and fluorescein. *J. Lab. Clin. Med.*, 44: 301-307.

Kirchmann, H and Witter, E., 1989. Ammonia Volatilization during aerobic and anaerobic manure decomposition. *Pant and Soil*, **115**:35-41.

Majumdar, G. P., 1935. Upavana-Vinoda (A Sanskrit Treatise on Arbori-Horticulture). *Indian Research Institute*, Calcutta, India. 128 pp.

Manna, M. C., Swarup, A., Wanjari, R. H., Ravankar, H. N., Mishra, B., Saha, M. N., Singh, Y. V., Sahi, D. K., Sarap, P. A., 2005. Long-term effect of fertilizer and manure application on soil organic carbon storage, soil quality and yield sustainability under sub-humid and semi-arid tropical India. *Fields Crop research*, **93**(2-3), 264-280.

Martin, J. P., 1950. Use of acid, rose bengal, and streptomycin in the plate method for estimation soil fungi. *Soil Sci.* **69**: 215-232.

Massoumi, A. and Cornfield, A. H., 1963. A rapid method for determining sulphate in water extracts of soils. *The Analyst.* **88**: 321–322.

McCarthy, A. J. Williams, S. T., 1992. Actinomycetes as agents of biodegradation in the environment- *a review. Gene*, **115**: 189-192.

Morovic-Budak, A., 1965. Experiences in the process of putrefaction in corpses in buried in earth. *Medicine, Science and the Law*, **5**:40-43.

Paull, J., 2011. Nanomaterials in food and agriculture: the big issue of small matter for organic food and farming. In: *3rd Scientific Conference of International Society of Organic Agriculture Research*, Namyangju, Korea, 2:96-99.

Payne, J. A., 1965."A summer carrion study of the baby pig (Sus scrofa Linnaeus)". Ecology, 46 (5): 592–602.

Rodriguez, W. C. and Bass, W. M., 1985. Decomposition of buried bodies and methods that may aid in their location. *J. Forensic Sci.* **30**: 836–852.

Sadhale, Nalini. (Tr.)., 1996. Surapala's Vrikshayurveda (The Science of Plant Life by Surapala). *Agri-History Bulletin No. 1. Asian Agri-History Foundation*, Secunderabad, India. 104 pp.

Saha, D. K. and Pan, S., 1997. Qualitative evaluation of some specific media of Trichoderma and Gliocladium and thier possible modification modications, *J. Mycopathol. Res.* **34**: 7-13.

Sarkar, S., Kundu, S.S. and Ghorai, D., 2014. Validation of ancient liquid organics-Panchagavya and Kunapajala as plant growth promoters. *Indian Journal of Traditional Knowledge*. Vol:**13**(2), pp-398-403.

Schwarz, P., Dannaoui, E., Gehl, A., Felkse-Zech, H., Birngruber, C. G., Dettmeyer, R. B., Verhoff, M. A., 2015. Molecular identification of fungi found on decomposed human bodies in forensic autopsy cases. *Int J Legal Med*, **129**:785-791.

Stach, J. E., Bull, A. T., 2005. Estimating and comparing the diversity of marine Actinobacteria. *Antonie van Leeuwenhoek*, **87**: 3-9.

Sundara Rao, W. V. B., Sinha, M. K., 1963. Phosphate dissolving organisms in the soil and the rhizosphere. *Indian J. Agr. Sci.* **33**: 272±278.

Tian, G., Brussard, L., Kang, T. B., 1995. An index for assessing the quality of plant residues and evaluating their effects on soil and crop in the (sub-) humid tropics. *Applied Soil Ecology*, 2:25-32.

Towne, E. G., 2000. Prairie vegetation and soil nutrient responses to ungulate carcasses. *Oecologia*, **122**: 232–239.

Walkley, A. and Black, I.A., 1934. An examination of the Degtjareff method for determining soil organic matter and a proposed modification of the chromic acid titration method. *Soil Sci.* **39**: 29-38.

Woo, S. L., Ruocco, M., Vinale, F., Nigro, M., Marra, R. and Lombardi, N., 2014. Trichoderma based products and their use in Agriculture. *The open Mycology Journal*, **8**(1): 71-126.