Original Research Article

PROXIMATE NUTRIENT COMPOSITION AND ANTIOXIDANT PROPERTIES OF PLEUROTUS SAPIDUS 969 CULTIVATED ON AGAVE SISALANA SALINE SOLID WASTE

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7 Abstract

8 Effects of pure and mixed substrates of sisal waste, grass (Panicum coloratum) and a 9 combination of the two substrates at 50:50 (w/w) on nutritional composition, minerals and 10 antioxidant potential of sun-dried Pleurotus sapidus 969 were investigated in the present 11 study. To determine the proximate chemical composition and antioxidant properties of the samples, standard analytical procedures were employed. Moisture content, crude protein and 12 crude fibre ranged between 11.09-12.80%, 6.4-6.6% and 18.3-30.5%, respectively. Macro 13 elements Ca, Mg, Na, K, and P were also found in substantial amounts with K being present 14 15 in an exceedingly higher amount (541.3-657.1 mg/100g) than the other macro minerals. The samples from the three substrates contained antioxidant β -carotene (4.6-6.0 mg/100g), 16 lycopene (4.9-5.1mg/100g), Vitamin C (5.2-5.6 mg/100g), phenols (361.0-859.0 mg of GA/g) 17 and flavanoids (33.5-64.0 mg RE/g). Mushroom harvested from mixed substrates contained 18 19 better nutritional qualities than the pure substrate, although the phenolic content in 20 mushrooms cultivated on sisal substrate was higher. The results further showed that, all the extracts exhibited scavenging ability and metal chelating activity. The findings showed that 21 Pleurotus sapidus 969 is rich in nutrients, macro minerals as well as natural antioxidant 22 23 which could be explored for pharmaceutical applications.

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25 Key words: Sisal, antioxidant, free radicals, *Pleurotus*, flavanoids, phenols

27 **1.0 Introduction**

28 Cultivation of the oyster mushroom, *Pleurotus* spp has increased greatly throughout the 29 world during the last few decades and constitute the second largest variety of mushrooms 30 produced in the world (Mshandete, 2011), with China being the primary source. *Pleurotus* 31 cultivation has the advantage of being cultivable in tropical climates, simple to produce, and 32 compatible with organic substrates rich in lignin and cellulose. Their ability to utilize 33 different substrates has made them the subject of broad research that generally mentions their 34 nutritional quality and the effect of substrate variation on the primary metabolites that are 35 directly related to the nutritional quality. Mushrooms have greatly varied and important uses 36 throughout the world (Wan Rosli, 2011). Mushrooms are valuable health foods since they are 37 poor in calories, fat, and essential fatty acids, and rich in proteins, vitamin and minerals (Reis

et al., 2012). Moreover, their medicinal properties have been reported such as anti-tumour
and immunomodulating effects (Ferreira *et al.*, 2010), reduction of blood cholesterol
concentrations, prevention or alleviation of heart disease and reduction of blood glucose
levels (Jeong *et al.*, 2010). These properties of mushrooms have been reported by Ferreira *et al.*, (2009), do be as a result of the bioactive products with antioxidant potential (sterols,
tocopherols, flavonoids, Carotenoids and phenolic compounds) (Ferreira *et al.*, 2009).

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45 Sequences of chemical reactions result in an imbalance between oxidant and antioxidant 46 reactions and are typically referred to as oxidative stress (Poli, et al., 2008). Both classes of 47 substances (oxidants and antioxidants) are generated in an oxidation-reduction (redox) set-up, 48 (Ralser, et al., 2007) and has been implicated as causes of degenerative diseases such as 49 atherosclerosis, cancer, and tissue damage in rheumatoid arthritis (Jang, et al., 2007). 50 Reactive species are commonly identified as substances leading to the oxidation of lipids 51 (epoxidation), glucose (glycation) and proteins (carbonylation). Maintenance of equilibrium 52 between free radicals production and antioxidant defences is an essential condition for normal 53 organism functioning (Valko et al., 2007). Non-controlled production of free radicals has 54 been attributed to various kinds of cancer and diabetes according to Ferreira et al. (2009). 55 Natural products with antioxidant activity, in particular mushrooms, are used to aid the 56 endogenous protective system, increasing interest in the antioxidative role of functional foods 57 or nutraceutical products (Reis et al., 2011). Antioxidants pay an important role in the 58 prevention and treatment of a variety of diseases by removing free radical intermediates and 59 inhibit other oxidation reactions by being oxidized themselves (Sies, 1997). Many studies 60 have found that some species of mushrooms are having therapeutic properties (Oyetayo, 61 2009) due to a wide variety of free radicals or reactive oxygen species scavengers which have 62 made them attractive as nutritionally beneficial foods and as a source for drugs development

63 (Guerra-Dore, 2007). According to Barros *et al.* (2008), mushroom flavonoids can act as free 64 radical scavengers to terminate the radical chain reactions that occur during the oxidation of 65 triglycerides in the food system. Apart from being a delicacy and tasty foods, mushrooms 66 have been reported to have special biochemical compositions, with significant contents of 67 antioxidant compounds, proteins, minerals, vitamins and water, which attract more attention 68 as functional health promoters (Wong and Chye, 2009).

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70 The chemical composition and nutritional quantity of edible mushrooms have been reported 71 previously (Agahar and Subbulakshmi, 2005). Studies have consistently shown an inverse 72 association between consumption of vegetables and fruits and the risk of certain forms of 73 cancer (Liu, 2003). However, the protective effects have been primarily attributed to well-74 known antioxidants, such as ascorbic acid and other related compounds (Soobrattee, et al., 75 2005).Different mushrooms species have been studied for new therapeutic alternatives and 76 the results proved their bioactive properties (Lindquist et al., 2005). Mushrooms are rich 77 sources of nutraceuticals (Elastase al., 2007), which are responsible for their antioxidant 78 content (Lo and Cheung, 2005). Recent investigations revealed that polysaccharides and 79 extracts of mushrooms had strong antioxidant and no synthase activation properties (Acharya 80 and Rai 2013; Patra at el., 2013; Samanta et al., 2013). According to Muhammad Nasir et al., 81 (2006), there are about 5000 different species of mushrooms, of which at least 1220 are 82 reported to be edible. There are about 40 species under *Pleurotus* mushroom, in that 25 83 species are commercially cultivated (Singh, 2011). Most of these cultivated mushrooms are 84 consumed as food or food ingredients in various food preparation and processed food 85 products. This has led to the growing interest in the use of edible mushrooms extracts as 86 dietary supplements based on the facts that they have a lot of bioactive compounds.

88	<i>Pleurotus</i> mushrooms can be grown on various agro-residues (as substrate) as reported by
89	Muthangya et al., (2014). The mushroom cultivation substrate has been reported to influence
90	its growth, yield as well as the functional, organoleptic and chemical composition (Micheal,
91	et al., 2011). This study was therefore designed to investigate the nutritive and antioxidant
92	property of P. sapidus 969 cultivated on Agave Sisalana saline solid waste and on grass
93	(Panicumcoloratum) as well as on a mixture of the two substrates at 50:50 (w/w) as reported
94	in Muthangya <i>et al.</i> ,(2013).

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96 2.0 Method :
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98 2.1 Samples of *Pleurotus sapidus* 969 Mushrooms

P. sapidus 969 mushrooms used in this study were cultivated on pre-treated saline sisal leaf
decortications waste as reported in Muthangya *et al.* (2013).Mushrooms were sundried on a

101 fabricated solar drier for 7 hours on a full sunny day before analysis.

102 **2.2 Determination of moisture, crude fibre and macro element content**

103 The sun-dried *P. sapidus* 969 mushrooms were analysed for moisture and total fibre content 104 using a Near Infrared Reflectance Spectroscopy (NIRS). The NIRS technique uses near-105 infrared light, instead of chemicals as in conventional "wet chemistry" methods. The samples 106 were prepared and analysed as described by Windham *al.*, (1989). The prepared mushrooms 107 samples were analysed for Ca, Mg, Na, K, and P, according to AOAC (2000).

108 **2.3 Crude protein determination**

109 Crude protein in *P. sapidus* 969was determined according to the method previously reported 110 by Tibuhwa *et al.*, (2012a). A known weight of each mushroom sample was taken and 111 digested using the micro Kjeldahl method. After completion of digestion, organic nitrogen 112 was determined calorimetrically using the Indophenol-blue method and NH_4^+ -N as standard. The absorbance was measured at 660 nm. The total crude protein was obtained and calculatedas described in Allen (1989).

115 **2.4 Mushroom crude extracts preparation**

116 The mushroom crude extract was prepared in ethanol according to Tibuhwa, (2012b), with 117 modification, where 1gm of dried whole mushrooms fruiting body was weighed at room 118 temperature (29±3°C). The samples were finely crushed using motor and pestle, and extracted 119 with 250 ml of ethanol as a solvent. The crushed powder was constantly stirred for 48 hrs and 120 thereafter filtered using Whatman number 4, filter paper. The filtrates were evaporated to 121 dryness in a rotary evaporator 90 rpm under reduced pressure and at 40°C. The concentrated 122 extracts obtained were stored in the dark at 4°C until further analysis. The yields of 123 evaporated dried extracts were obtained by the gravimetric method. The percentage yield of 124 the extracts was calculated based on dry weight as:

W₂

- 125 $Yield (\%) = (W_1 X 100)$
- 126
- 127 Where: W_1 = weight of extract after ethanol evaporation
- 128 W_2 = Weight of the ground mushroom powder

129 2.5 Quantitative Antioxidant assay

130 **2.5.1 Determination of total phenolics content (GAE/g)**

The concentration of phenolic compounds in the extract of *P. sapidus* 969 mushroom was measured by Folin-Ciocalteu colourimetric method according to the method previously reported by Tibuhwa (2012b), with modification. A blue colour was developed by reaction of phenolic compounds and Folin-Ciocalteu's reagent. The extract solution (1 ml) was mixed with 1 ml of Folin-Ciocalteau reagent and after 3 min, 0.8 ml of 7.5% (w/v) sodium carbonate was added to the mixture. The reaction was kept in the dark for 30 min with agitation and thereafter centrifuged at 3300 g for 5 min. The absorbance was measured at 765 nm and total phenolic content was expressed as gallic acid equivalent (GAE) to 1 g perextract using gallic acid as a standard.

140 **2.5.2 Determination of total flavonoid**

Determination of total flavanoids was carried out using the aluminium chloride colourimetric method according to Jaita *et al.* (2010), as reported in Tibuhwa (2012b). Each extract (1 ml) was diluted with 4.3 ml of 80 % aqueous ethanol containing 0.1 ml of 10% aluminium nitrate and 0.1 ml of 1M aqueous potassium acetate. The mixture was incubated for 40 minutes at room temperature and the absorbance determined colourimetrically at 415 nm. A standard curve of flavonoids was prepared and concentration of flavonoids in the test samples determined.

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149 **2.5.3** β -carotene and Lycopene contents

150 β -carotene and lycopene were determined according to the method of Nagata and Yamashita, 151 (1992). In brief, 100 ml of mushroom extract (10 mg/ml) was vigorously shaken with 10 ml 152 of acetone-hexane mixture (92:3) for 1 min. and filtered through Whatman number 4 filter 153 paper. The absorbance of the filtrate was measured at 453, 505 and 663 nm. β -carotene and 154 lycopene contents were calculated according to the following equations:

155 Lycopene (mg/100mg) =
$$0.0458 A_{663} + 0.372 A_{505} - 0.0806 A_{453}$$

156
$$\beta$$
-carotene (mg/100mg) = 0.216 A₆₆₃ - 0.304 A₅₀₅ + 0.452 A₄₅₃

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158 **2.5.4 Determination of Vitamin C**

The vitamin C content was determined diametrically using 2, 6 DichlorophenoIndophenol methods according to Plumer (1987).One (1) gram of grounded sample was mixed with 25 ml of 5% metaphosphoric acid solution and shaken for 30 min. The mixture was then filtered through Whatman no. 42 filter paper using suction pump. Ten (10) ml of the filtrate was 163 titrated against 0.025% of 2.6 Dichlorophenol Indophenol reagents. The amount of vitamin C

164 in each extract was calculated from the equation:

165 Ascorbic acid mg/100g = $\underline{A \times I \times V \times 100}$ 166 $V_2 \times W$

167 Whereas A = Quantity of ascorbic acid (mg) reacting with 1ml of 2, 6 Indophenol

168 I = Volume of indophenol (ml) required for the completion of extract titration

169 $V_2 = Total volume of extract$

170 W = Weight of the ground mushroom

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172 **2.6 DPPH free radical scavenging activity**

The scavenging ability on 1,1-diphenyl-2-picrylhydrazyl (DPPH) radicals was determined according to the method of Masuda *et al.*, (2000), and Jaita *et al.*, (2010), as previously reported by Tibuhwa *et al.*, (2012a). Each extract (0.01-0.14 mg/ml) was mixed with 1 ml of methanolic solution containing DPPH radicals (0.4 mM). The mixture was shaken vigorously and left to stand for 30 min in the dark. The absorbance was measured at 515 nm. The percentage of DPPH radical scavenging activity of each extract was determined within the range of dose-response and was calculated as:

180 DPPH radical scavenging activity (%) = $\frac{A_0 - (A_1 - A_s) *}{A_0}$ 100

181 182

Where A_0 = Absorbance of the control solution containing only DPPH

183 A_1 = absorbance in the presence of mushroom extract in DPPH solution

184 $A_s =$ the absorbance of the sample extract solution without DPPH

185 The EC50 value (total antioxidant necessary to decrease the initial DPPH radical 186 concentration by 50%) was determined from a plot of scavenging activity against the 187 concentration of extracts.

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190 **2.7 Chelating effect on ferrous ions**

191	The ability of <i>P. sapidus</i> 969 extracts to chelate ferrous ions was estimated by the method of
192	Dinis et al., (1994). The extract (1 mg/ml) was added to a solution of 2 mM ferrous chloride
193	(0.05 ml). The reaction was initiated by the addition of 5 ferrozine (0.2 ml) and the mixture
194	was then shaken vigorously and left to stand at room temperature (28-30°C) for 10 min. The
195	absorbance of the solution was measured spectrophotometrically at 562 nm. The percentage
196	inhibition of ferrozine-Fe ²⁺ complex formation was calculated as;
197	$\{(A_0 - A_1) \ / \ A_0\} \ imes \ 100$
198	Where A_0 = absorbance of the control
199	A_1 = absorbance in the presence of the mushroom extract
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202	Statistical analysis
203	The experimental results were expressed as mean \pm SD (Standard deviation) of n=3
204	measurements. Statistical analysis of the data was carried out using student's t-test and the
205	results were considered significant when $P = .05$.
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207	3. Results and Discussion
208	3.1 Composition of sun-dried <i>Pleurotus sapidus</i> 969
209	The moisture contents of the dried <i>Pleurotus sapidus</i> 969were found 11.9-12.8% (Table 1)
210	with no significant difference at P = .05 level. The highest moisture content was found in
211	P.sapidus 969cultivated on a mixture of sisal and grass substrate (1:1), followed by grass and
212	the least was in sisal alone.
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216	
217	Table 1. Composition (%) of sun-dried Pleurotus sapidus 969 and crude extract yields,
218	Mean±SD, n=3).

Cultivation	Moisture (%)	Total fibres	Crude	Crude extract yield
substrate		(%)	Proteins (%)	(%)
Sisal	11.9±0.03	6.4±0.1	18.3±0.4	17.0±0.4
Grass	12.2 ± 0.01	6.6±0.2	23.3±0.2	17.9±0.3
Sisal: Grass	12.8±0.04	6.5±0.2	30.5±1.2	13.7±0.4

The fibre content was found highest in *P. said* 969 (6.6g/100g) cultivate on grass. The variation in fibre content between the mushrooms from the three different substrates was not statistically significant at *P*=.05. Comparison of the results of the protein content of the mushroom from the three substrates showed a significant difference at *P*= .05 with the highest crude protein content being recorded from the mushrooms in the combined substrate of sisal and grass (30.5%).

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227 The results of obtained by this study for the dried *Pleurotus sapidus* 969 are within the range 228 of those reported previously for other *Pleurotus* species. Muthangya et al. (2014), working on 229 *Pleurotus* HK 37 from the same substrate reported results, which were within the range of 230 those obtained in this study. Oyetayo and Ariyo, (2013), working on Pleurotus 231 ostreatus reported the moisture content of dried samples to be within 9.00-10.72%. While 232 previously, Chang and Miles(2004), reported the moisture content of dried mushrooms to be 233 in the range 9 - 13%. Sales-Campos et al., (2011), reported a variation in fibre content while 234 working on several *Pleurotus* sp. grown on crushed sugar cane, elephant grass and banana 235 tree leaves, on the other hand, the results obtained on the fibre content was within the range 236 (5.4–30.0%) previously reported by other authors for *Pleurotus* sp. (Kurtzman, 2005) 237 cultivated on different substrates. The protein contents of mushrooms are reported to vary 238 according to genetic structure of species, physical and chemical differences in growing 239 medium (Akyüz, and Kirba'g, 2010), cultivation time and strain (Bernas, et al., 2006; 240 Mshandete and Cuff, 2007), as well as the stage of development and level of nitrogen 241 available (Chang and Miles, 2004). The mushroom protein contents that were found in this

study (Table 1) are in agreement with the range of mushroom protein contents reported in the literature (Bernaś*et al.*, 2006) varying between 17 and 42.5%, but higher in *P.sapidus* 969cultivated on grass and on a combined substrate of grass and sisal than the value (20.28%) reported by Bonatti *et al.*(2004) for *Pleurotus ostreatus*cultivated on cotton waste. The present results showed that protein content of *P.sapidus* 969 was significantly higher when the mushroom was cultivated on a combination of sisal and grass than that obtained for the mushroom grown on separate substrates.

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250 **3.2 Macro-minerals elements**

Pleurotus sapidus 969mushroom samples analysed in this study contained macro-minerals
 including; calcium, magnesium, sodium, potassium and phosphorus (Table 2).

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Table 2. Macro-minerals composition of *P.sapidus* 969(g/100g of dried sample)Mean±SD,
n=3

Cultivation substrate	Macro-minerals (mg/100g)				
	Ca	Mg	Na	K	Р
Sisal	6.1±0.4	16.21±0.6	15.17±0.1	614.5±1.9	117.7±0.9
Grass	7.7±0.3	17.8±0.5	14.2±0.3	541.3±2.2	123.4±0.9
Sisal : Grass	7.6±0.1	18.1±0.4	16.4±0.2	657.1±4.8	131.7±2.0

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258 The highest amount of Ca (7.7 mg/100g) was recorded in the *P. sapidus* 969 samples from 259 grass substrate, followed by sisal: grass (7.6 mg/100g) and lastly sisal (6.1 mg/100g). Mg 260 concentration was the highest in sisal: grass samples (18.1 mg/100g) and the least in samples 261 obtained from sisal substrate (16.21 mg/100g). The value of Na, K and P in the *P.sapidus* 262 969were found to be in the range of 14.2-16.46, 541.2-657.1 and 123.4-131.7 mg per 100g, 263 respectively. Minerals in human diets are essential constituents for metabolic reactions, 264 transmission of nerve impulses, healthy bone formation, regulation of water and salt balance 265 Kalac, and Svoboda, (2000). The mineral contents of P. sapidus 969 from the two different 266 substrates and their combinations in this study did not vary significantly at P=.05. The results 267 of the macro-minerals elements composition of *P.sapidus* 969 are within the range as those 268 reported by Muthangyaet al., (2014), from dried samples of Pleurotus HK 37 cultivated on 269 the same substrates, although slightly higher. The values of calcium in this study are an 270 indication that *P. sapidus* 969 is a valuable food for formation and maintenance of bone and 271 normal function of nerves and muscles in humans and other vertebrates as reported by Wang 272 et al. (2010). Mg, an essential co-factors for certain enzymes in various biochemical pathways 273 was detected in P. sapidus 969and the levels of Mg were quite higher than those reported 274 (1.69-3.57 mg/100g) for *Pleurotus ostreatus* cultivated on different woody substrates(Oyetayo 275 and Ariyo, 2013). Na and K are important in the maintenance of osmotic balance between 276 cells and the interstitial fluid in animal systems (Afiukwa, 2013). These results indicate that 277 these mushrooms could play a role in human health by lowering blood pressure, reducing the 278 risk of osteoporosis and in maintaining bone health (Wang et al., 2010). The results of 279 phosphorus in this study (123.4-131.7mg/100g) compare well with 122.28 mg/100g reported 280 for a wild P. ostreatus (Afiukwa, 2013). The differences in phosphorus contents in mushroom 281 have been attributed previously to substrates what about mushroom species/strain since they 282 differ in substrate utilization/absorption and translocation of biomaterials from substratesused 283 for growing the mushrooms according to Ahmed, (2009). Pleurotus species canprovide a 284 useful source of phosphorus, potassium, calcium, and magnesium. Thus, the inclusion of 285 P.sapidus 969in the diet could be one of the strategies for combating macronutrient 286 deficiencies

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288 **3.3 Antioxidant contents of** *Pleurotus sapidus* **969**

289 **3.3.1 Total Phenol and Flavonoid contents**

290 The total phenolic and flavonoid content in *Pleurotus sapidus* 969analysed in this study are

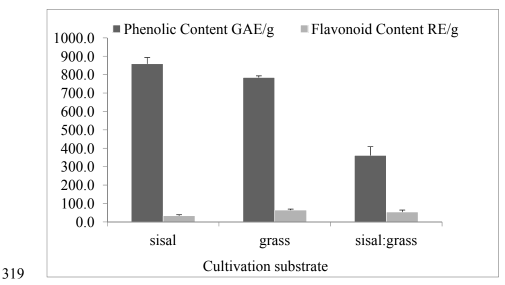
- shown in Figure 1. The total phenolic and flavanoids contents in the mushroom samples were
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859.0, 784.7 and 361.0 mg of GA/g and 33.5, 64.0 and 53.8 mg RE/g in the mushrooms
grown on sisal, grass and sisal: grass substrates, respectively.

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295 The findings of this study is supported by previous findings of Phenolic compounds in 296 mushrooms as reported by Tibuhwa, (2012b)and linked to various biological functions 297 including antioxidant activity Phenolic compounds are well known secondary metabolites 298 commonly found in plants and mushrooms and reported to have vital biological functions 299 including antioxidant activity (Dimitrios, 2006). Knowing the amount of total phenolic 300 compounds in mushrooms is of great importance in their nutritional and functional 301 characterization since the profile of the phenolics has been reported to be species-specific 302 Banerjee et al., (2012). Phenolic compounds have been reported to be of great interest due to 303 their possible use as dietary supplements or food preservatives, Jayakumar et al., (2009). 304 Several species of mushroom have been reported to contain a wide variety of free radicals or 305 reactive oxygen species scavengers, which have made mushrooms attractive as nutritionally 306 beneficial foods and as a source for drugs development (Guerra-Dore, 2007). Barros et al. 307 (2008) reported that mushroom flavonoids can act as free radical scavengers to terminate the 308 radical chain reactions that occur during the oxidation of triglycerides in the food 309 system.Flavonoids have been reported to decrease capillary fragility and exert a cortisone-310 like effect on tissues (Gonzalez-Nunez et al., 2001) and protect against cancer and heart 311 diseases (Filippo set al., 2007). It, therefore, implies that the high flavonoids content in the 312 mushroom extracts might be responsible for the therapeutic effect of some mushroom species 313 earlier reported (Ogbonnia, et al., 2008).

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Fig. 1. Total phenol and flavonoid contents of *P. sapidus* 969, Values are expressed as mean \pm SD mg of Gallic acid equivalent per gram of dry weight (mg GAE/gm).

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Previous studies have shown that food consumption with high phenolic content can reduce the risk of heart disease (Singla *et al.*, 2010). From this study, the high levels of phenols and flavanoids make *P. sapidus* 969 favourable for nutritional and therapeutic application as supported by the findings of Ferreira *et al.*, (2007).

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329 **3.3.2** β-carotene, Lycopene and Vitamin C content

330 Carotenoids are natural colourants, stabilizers and active in the protection process of human 331 body cells, where they balance and offset the destructive effects of free radicals Jayakumar et 332 al., (2009). The quantities of β -carotene, Lycopene and Vitamin C content of P. sapidus 333 969analysed in this study are presented in Figure 2. The content of β -carotene was in the 334 range of 4.6 mg/100g to 6.0 mg/100g, lycopene was in the range of 4.9 mg/100g to 5.1 335 mg/100g, while vitamin C was in the range of 5.2 mg/100g to 5.6 mg/100g in the three 336 substrates. Carotenoids are major antioxidants with known health benefits, while diets high in 337 lycopene; a cyclic isomer of β -carotene has been linked to reduction of prostate cancer and cardiovascular diseases (Raoand Agarwal, 2000); whereas, Ascorbic acid is reported to
directly interact with radicals in plasma, preventing damage to red cell membranes
(Jayakumar *et al.*, 2009).

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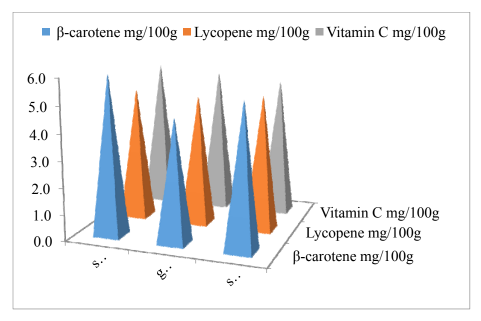


Fig. 2. Total β -carotene, Lycopene and Vitamin C, the content of *P. sapidus* 969 Values are expressed as mean \pm SD mg/100g.

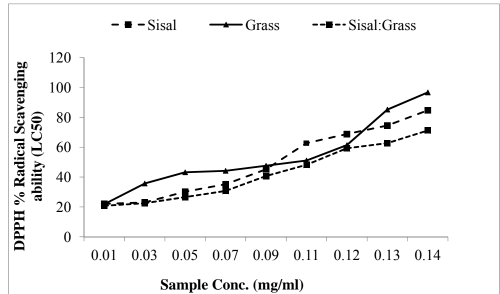
346 The results of β -carotene, lycopene and vitamin C obtained in this study are within the range 347 of those reported previously by Muthangyaet al. (2014), from sun-dried samples of Pleurotus 348 HK 37 cultivated on similar substrates. The presence of these compounds in *P. said* 969 is an 349 indication that these mushrooms are equipped with antioxidant properties. Jayakumar et al., 350 (2009) reported similar findings of carotenoid and ascorbic acid compounds from P. 351 ostreatusmycelium extracts. The quantities of these compounds in various extracts have been 352 suggested to be influenced by the culture medium used for producing the mycelium (Petreet 353 al., 2010), a similar scenario observed in this study where different substrates were used to 354 cultivate P. said 969. These findings support Barros et al., (2007), who reported that the 355 carbon source and especially the nitrogen sources have a direct influence on the quantum of 356 biologically active substances in the extracts.

358 3.4 Antioxidant activities

359 3.4.1 DPPH Free radical scavenging activities

The result from this study showed that the free radical scavenging activity of *P. sapidus* 969extract from the three cultivation substrates increased with increasing concentration of extract indicating the concentration dose dependency of anti-oxidative activities (Figure 3). This observation concurs with that of Banerjee *et al.*, (2012) who also noted a similar trend of anti-oxidative activities dose dependency and associated it with the presence of reductones that are reported to be the terminators of free radical chain reactions.

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Sample Conc. (mg/ml) Fig. 3. DPPH radical scavenging activity (%) of *P. said* 969 (ethanolic extract) cultivated on sisal grass, sisal: grass at 1:1 Values recorded are (mean \pm SD, n=3).

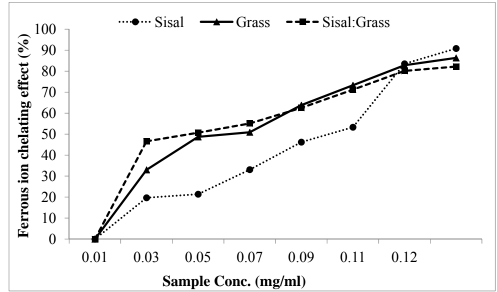
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In this study, the maximum scavenging activity values were at a dilution of 0.14mg/ml. The mushroom extracts from grass substrate showed the highest percentage (96.7%) scavenging power while the extracts from sisal and sisal: grass had 84.7% and 71.2%, respectively.However, the total antioxidant necessary to decrease the initial DPPH radical concentration by 50%, determined from plotted graph of scavenging activityagainst different 376 concentration of the extracts, showed the extract from sisal had the highest ability (EC50 <377 0.09 mg/ml) followed by that from grass (EC50 < 0.11 mg/ml) while the extracts from 378 sisal: grass had the least ability of (EC50 < 0.13 mg/ml), a similar observation reported by 379 Muthangya et al., (2014), working dried samples of Pleurotus HK 37 cultivated on the same 380 substrates. This result shows that the P. sapidus 969mushroom studied have high scavenging 381 ability compared to other mushrooms. Although in this study, mushrooms from sisal: grass 382 had the least ability of (EC50 < 0.13 mg/ml), this value is still better compared to other well 383 appreciated antiradical mushrooms. Filipa al., (2011) established EC50 values in 384 *Paxillusinvolutus* and *Pisolithusarhizus* of (EC50 = 0.61 and EC50 = 0.56 mg/ml), 385 respectively which show them having relatively low free radical scavenging ability compared 386 to mushrooms from sisal: grass with least ability in this study. The higher content of phenolic 387 compounds in mushrooms cultivated on sisal substrate could be the cause of the high total 388 antioxidant necessary to decrease the initial DPPH radical concentration by 50% an 389 observation in line with the findings of Abdullah et al.(2011), working on of Brazilian button 390 mushrooms.

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392 **3.4.2** Chelating ability of ferrous ions

Figure 3 depicts the iron chelating ability of *Pleurotus sapidus* 969 cultivated on the different substrates under investigation in this study. The ferrous ion-chelating effect of all samples increased well with increasing concentrations (Figure 4). *P. sapidus* 969 from sisal substrate had the highest iron chelating ability (90.8% at 0.12 mg/ml), while the weakest metal chelating ability (82.2%) was recorded for samples from a combined substrate of sisal and grass.



400
401 Fig. 4. Ferrous ion chelating effect (%) of *P. said* 969 (ethanolic extract) cultivated on sisal
402 grass, sisal: grass at 1:1.
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Extract from samples cultivated on sisal substrates recorded 86.3% metal chelating ability at the same concentration. It has been observed that metal ion chelating antioxidants would also remove the oxidative damage from other less prominent but equally damaging pro-oxidant metal ions such as Cu (Halliwell, 2001). Thus, the iron chelating capacity of the mushroom species would prevent transition metals to participate in the initiation of oxidative stress.

410

411 Conclusion

412 It was observed that fruiting bodies harvested from different substrates varied in their 413 biochemical analysis. It might be due to the variability of the substrates to provide different 414 nutritional elements to mushroom grown on these substrates. Among the substrates 415 investigated in this study, a combination of Sisal and grass gave the best overall composition 416 of all the nutrients. The nutritional and antioxidant investigations on the mushroom cultivated 417 on the different substrate revealed that all the mushrooms possess high reductive potential 418 and metal chelation activities, with high concentration of macro nutrients, proteins, total 419 phenol and total flavonoids. These bioactive compounds together with the high antioxidant

421	enhance the immune system against oxidative damage, or it may be utilized as a potential			
422	source for drug development.			
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