1	Original Research Article		Comment [u1]: It is very well-marked that this study is acceptable with minor revision
2			and useful for publish in this journal. The discussion section can be developed with
3	Anti-aging activity of <i>Xvlaria striata</i> in		literature data.
4	Drosophila melanogaster		Note: Please, I want to see manuscript again after the corrections.
-			In addition to, please add some papers and you
5			can use them in manuscript.
6			Please see below for papers.
7	ABSTRACT		2. Sevindik M. Investigation of Antioxidant/Oxidant Status and Antimicrobial
8 9	Aims: To evaluate the application potential of <i>Xylaria striata</i> Pat in anti-aging field		Activities of Lentinus tigrinus. Advances in pharmacological sciences 2018
10	Methodology: Extracting the fruit body of <u>Xularia</u> X striata by ultrasonic-assisted		DOI:10.1155/2018/1718025
11	extraction method. In this study, <i>Drosophila melanogaster</i> was used as an anti-aging		antioxidant status of edible mushroom
12	organism material. After feeding with different concentrations of extract, the natural survival time, survival time under oxidative stress and survival time under UV irradiation of		Fungus. 2018; 9(2):165-168
13	Drosophila-D. melanogaster were all measured. In addition, the <i>in vivo</i> activity of SOD.	`	Formatted: Font: Italic
14	CAT and MAD, protein concentration and body weight were determined to evaluate the anti-aging effect of ethanol extract from Xylaria X. striata.		Field Code Changed
15	Results: The results showed that the ethanol extract of Xvlaria -X, striata could extend the		Field Code Changed
10	lifespan of <u>Drosophila D.</u> melanogaster under both irradiation and oxidative stress		Field Code Changed
10	condition. And the ethanol extract could enhance the activity of CAT in <i>Drosophila <u>D</u>.</i> <i>melanogaster</i> , especially at concentration of 50 µM, and the content of MAD in <i>Drosophila</i>		
17	<u>D.</u> melanogaster were decreased significantly.		
18	Conclusion: This study clarified the anti-aging activity of Xylaria X. striata in Drosophila		
19	<u>D.</u> melanogaster and it would provide some theoretical basis for its further development and utilization in anti-aging drugs and health food.		
20			
21	Keywords: Xylaria striata; anti-aging; Drosophila melanogaster; lifespan.		
22			
23 24	1 INTRODUCTION The higher fungi contain compounds with many povel structure, which have variety and		
25	remarkable biological activity ^[1-3] . Because of potential medicinal and economic value, they		
26 27	nave drawn an increasing attention and become a highlight research field of science and industry in recent years.		
28	The genus Xylaria, a big family of mycomycetes, has great value in application according to a		
29 30	large amount of literature reports. There were many kinds of compounds such as terpenoids, sterols, alkaloids, polyketones, polysaccharides, cyclic peptides and carboxylic acids ^[4-6] had		
31	been isolated from the <i>Xylaria</i> genus, which exhibited antioxidant, antimicrobial, antitumor,		
32 33	belong to the family of <i>Xylaria</i> , mainly grows on decayed barks and lived roots of broad-	~	Formatted: Font: Not Italic
34 35	leaved woodland. It was used as folk medicine in China ^[/] . After finding its diverse biological activities in the preliminary screening of large numbers of edible and medicinal mushrooms		
36	our research group started a systematic study on <i>Xylaria striata</i> from cultivation method to		
37 38	cnemical components since 2013 ⁶⁵ . These research results clarified its effect of anti-animal and plant pathogens ^[10] , anti-tumor ^[11] , and promoting sleep ^[12] .		
39	Furthermore, another mushroom from the same genus named <u>Podosordaria nigripes</u>		Formatted: Font: Italic
40 41	(Klotzsch) P.M.D. Martin (Syn: Xylaria nigripes (Klotzsch) Cooke has been widely used to prevent and treat senile diseases in China. It can promote the effect of antidepressants [4413]		Formatted: Font: Italic
•• 1			

42	and alleviate depressive symptoms in patients with epilepsy ^[14] Hence, the main objectives of	
43	this study were to evalute the anti-aging activity of Xylaria X. striata using Drosophila	Formatted: Font: Italic
44	melanogaster as the model organism by assessing the lifespan of flies, the activity of SOD	
45 46	and CAT, the content of MDA and protein and its stress resistant ability under H ₂ O2 and UV irradiation treatment	Field Code Changed
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47		
48	2 MATERIALS	
49		
50	2.1 Biomaterial	Field Code Changed
51	Vularia V. striata was abtained in 2012 from Oingui town, Mianyang, Siahuan province of	
52	<u>Ayidra A.</u> striata was obtained in 2013 from Qingyi town, Mianyang, Sichuan province of China, and identified by professor Xin-sheng He of microbiology. Southwest University of	
54	Science and Technology. The voucher specimen was preserved in the Microbiology	Field Code Changed
55	Laboratory of the same university. Drosophila D. melanogaster, reared under the condition of	Tield Code Changed
56	temperature 25 ± 1 , humidity 60%-80%.	
57	2.2 Reagents	
58		
59	Protein content determination kit, superoxide dismutase (SOD) determination kit, catalase	
60	(CAT) determination kit, and malondialdehyde (MDA) content determination kit are all	
61	purchased from Nanjing Jiancheng bioengineering institute; Corn flour, agar and yeast extract	
62 63	was purchased from Bejing Aobaoing Bio-Tech Co., Ltd.; Ethyl ether, ethanol, 30% hydrogen	
64	reagents are analytical pure grade and pure water is made in the laboratory.	
- -		
65	2.3 Equipments	
67	Biochemical incubator/SPX- 80B, Shanghai Kuntian instrument Co., 1 td.). Thermo Scientific	
68	Microplate Reader(Multiskan Spectrum, Thermo Fisher), Ultraviolet lamp(LP-GDZJ40W,	
69	LONGPRO CO., Ltd.), Ultrasonic cleaner(KQ-200KDE, Kunshan Ultrasonic Instrument Co.,	
70	Ltd.).	
71		
72		
72	3 METHODS	
74	3.1 Sample pretreatment	
75	The cultivated fruiting body ^[15] of Xylaria X. striata was dried at 50 and ethanol-assisted	
76	ultrasonic extraction process was carried out at the ratio of material to liquid (1:30 g:mL) for	
77	three times, 10 min of each. Then, the extract was obtained by vacuum distilling of the filtered	
78	supernatant at 65 and preserved at 4° C untill use.	
79	3.2 Diet preparation and Fly husbandry	
80		
81	The wild-type Drosophila D. melanogaster Canton-S (CS) flies, selected as the experiment	
82	organism for later assay, were reared at temperature 25 ± 1 , humidity 60%-80%. Before the	
83 84	experiment, Drosophila D. melanogasters were mass reared in 500 mL erienmeyer flask containing 100 mL of standard commeal diet ^[16] (13% commeal w/v, 8% sugar w/v, 2.4% Agar	
85	w/v, 1.6% yeast extract w/v and 0.4% acetic acid v/v). To avoid overcrowding. 200 flies of	
86	each bottle was enough. When the progeny drosophila was going to hatch, all adult flies	
87	would be transferred to the new culture bottle. Afterwards, the flies eclosed within 24 h were	Field Code Changed
88	collected and segregated according to their sex.	

3.3 Longevity assay

- Refers to Proshkina *et al*^[17], 10 newly eclosed male flies, reared on a standard cornmeal medium with ethanol extract from <u>Xylaria X.</u> striata. The extract was dissolved in water and mixed with the standard diet at final concentrations of 50, 100 and 200 μ M. Negative control diets contained only water while positive control contained ascorbic acid at final
- 93 94

- 95 $\,$ concentrations of 100 $\mu M.$ During the rearing process, diet was replaced with fresh medium
- 96 every five days and the numbers of alive flies were recorded every two days. The test was
- 97 stopped until all flies were dead. All the treatments were carried out with 3 replicate. Median
- 98 lifespan was calculated using the method reported by Kaplan–Meier^[18] previously.

99 3.4 Oxidative stress resistance

- 100
- 101 The situation of group dividing is the same as Longevity assay. Newly eclosed male flies were
- 102 fed on standard diet with the extract from <u>Xylaria X.</u> striata for 25 days, followed by an
- 103 oxidative challenge of 9% H_2O_2 in 6% glucose solution on filter paper strips^[19]. The number of
- 104 death was recorded every 4 h until all flies were dead. Median lifespan was calculated as the
- 105 same as 3.3.

106 **3.5 Irradiation resistance**

107

113

- 108 The flies were treated at a constant distance of 10 cm from the ultraviolet lamp and irradiation
- 109 time was 20 min^[20]. We strictly controlled the surrounding temperature at 25 to decrease
- 110 heat stroke from the lamp during irradiation. Then, the flies were transferred to vials. Dead
- 111 flies were counted every day until all flies died. Life span was studied in the same way.

112 3.6 SOD, CAT, MDA, Protein content and weight assays

114 After 25 days of feeding, flies were transferred into empty tube for 2 hours. After anesthesia,

- 115 the flies were homogenized in an ice bath at the ratio of weight to normal saline at 1:19 (g:mL).
- 116 The supernatant was obtained after the homogenate was centrifuged in 2500 r · min⁻¹ for 20
- 117 min at 4 . The activity of CAT(catalase) and SOD(superoxide dismutase), the content of
- 118 MDA(malondialdehyde) and protein were determined by the Thermo Scientific Microplate
- 119 Reader according to the instruction described by the kit. Flies were anesthetized and weighed
- 120 every ten flies. For each concentration, 3 replicate were set up.

121 3.7 Statistical Analysis

All the data collected were repeated 3 times, and the data were displayed in the form of

- means±SD. The statistical analysis was performed with SPSS 20.0 software, and the
- difference analysis between groups was performed by One-way ANOVA analysis. Significant
- 126 differences were expressed by: * P < 0.05; ** P < 0.01; *** P < 0.001.

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128 4 RESULTS AND DISCUSSION

130 **4.1 Longevity assay**

131 Drosophila-D. melanogaster has been widely used in anti-aging experiments as a model organism because of its short lifespan and easy reproduction^[21]. The anti-aging ability of the 132 133 134 drugs or its toxicity can be reflected at a certain level by comparing the lifespan of Drosophila 135 D. melanogaster before and after feeding the drugs. As shown in Table 1, the ethanol extract 136 of Xylaria X. striata can significantly shorten the mean lifespan of Drosophila D. melanogaster, and with the increasing of the dosage, the more significant the shorten effect is. In addition, 137 138 the maximum lifespan and half survival time of Drosophila-D. melanogaster were also reduced. Therefore, there may be some biologically toxic substances in the ethanol extracts 139 of Xylaria X. striata. It has been reported by Yuan et al.^[11] that Xylaria X. striata has an 140 inhibitory effect on plant and animal pathogens, which can also prove that it may contain 141 142 some toxic substances.

143

Table 1 The effect of Xylaria-X. striata on lifespan

Treatment	Concentration	Max±SD	H±SD	M±SD
Treatment	(µM)	(day)	(day)	(day)

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Control	-	45.3±1.2	35.7±1.6	30.6±2.1	
N 1 1 N	50	32±0.9*	20±2**	28.2±0.15	
Xylaria X.	100	41.3±6.2	31.3±4.7		Formatted: Font: Italic
Sillala	200	38±4.3	15.3±1.9***	24.2±2.1**	
Ascorbic acid	100	48.9±3.2	39.0±3.4**	37.7±0.7*	

Note: Max, maximum lifespan, days; M, mean lifespan, days; H, half survival lifespan; *p < 0.05, **p < 0.01, ***p<0.001

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145 **4.2 Oxidative stress resistance**

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147 In vivo, hydrogen peroxide reacts with oxygen to produce hydroxyl radicals, which can induce

acute oxidative stress damage and shorten the survival time of <u>*Drosophila*</u> <u>*D*</u>. *melanogaster*^[22].

As shown in Table 2, the ethanol extract of *Xylaria X. striata* could prolong the maximum

150 lifespan, half survival time and mean lifespan of <u>Drosophila D.</u> melanogaster under oxidative

151 stress. When the concentration of extract was 100 μ M, the maximum lifespan and the mean

lifespan were obviously extended, and the prolongation of the mean lifespan was extremely
 significant at the concentration of 200 μ M.

154

Table 2 The effect of oxidative stress on lifespan

Treatment	Concentration (µM)	Max±SD (h)	H±SD (h)	M±SD (h)			
Control	0	23.3±3.1	16.7±1.1	17.2±0.6			
	50	22±2	17±1.3	17±0.6			
Xylaria X. striata	100	29.3±3.3*	18±4	18.3±0.3*			
	200	21.3±2.3	15.3±1.1	18.7±0.3**			
Ascorbic acid	100	26±2	18.7±2.3	18.5±0.5**			
Note: *p < 0.05, **p <	Note: *p < 0.05, **p < 0.01, ***p<0.001						

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157 4.3 UV irradiation resistance

unable repaired the radiation damage.

158 159 Ultraviolet radiation can cause changes in some physiological functions and damages the 160 body. Herein, we determined if the Xylaria-X. striata could repair UV irradiation damage by observing the lifespan changes of Drosophila D. melanogaster under UV irradiation stress. 161 According to a previous report, the sensitivity of *Drosophila_D.* melanogaster to ultraviolet light was closely related to the melanin content^[23]. So, this method could also show the effect of 162 163 the tested agent on melanin production in Drosophila D. melanogaster. As can be shown in 164 Table 3, a low dose of extract could significantly prolong the lifespan of Drosophila D. 165 melanogaster after UV irradiation. Medium, high dose groups could also prolong lifespan, but 166 167 the effect was inferior to the low dose group. This may be due to the variety of substances in the extract. As the increasing of dose, the increasing biotoxicity resulted in a decrease in 168 169 lifespan. In addition, we found ascorbic acid can't prolong the lifespan, which indicated that it

Table 3 Effect of UV irradiation on lifespan

Treatment	Concentration (µM)	Max±SD (day)	H±SD (day)	M±SD (day)	
Control	0	12.5±0.7	6±1.3	5.2±0.4	
	50	17±1.4*	11.5±0.7*	10.4±2.3*	
Xylaria <u>X.</u> striata	100	15±1.4	10±2.8	8.5±1.5	Formatted
	200	14.5±1.5	9.3±2.9	9±2.5	

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¹⁷⁰ 171

	Ascorbic acid	100	12.5±0.7	9.5±0.7	9.4±0.4
172	Note: *p < 0.05, **p <	:0.01, ***p<0.001			

173

175

174 4.4 SOD、CAT、MAD, protein content and weight assay

176After feeding on the extract, the overall living condition can be indirectly reflected by177determining the weight and protein content of *Drosophila-D.* melanogaster. In Table 4, there178were no significant changes in the weight and protein content of flies.

179 Superoxide dismutase (SOD), one kind of free radical enzyme, can scavenge the superoxide 180 anion in body, and prevent the body from being oxidized. Hydrogen peroxide reacts with 181 oxygen can produce hydroxyl radicals, which are harmful to the body. However, hydrogen 182 peroxide decomposition *in vivo* can be catalyzed by catalase (CAT) to reduce the generation 183 of hydroxyl free radicals^[24]. Hence, SOD and CAT are two kinds of representative enzymes in 184 antioxidant system. The activities of these two enzymes can indirectly evaluate the free 185 radical scavenging ability of the organism^[25].

Malondialdehyde (MDA) is a kind of lipid peroxidation product produced by the reaction of
 free radicals and unsaturated fatty acids. MDA residues in body can be further cross-linked
 with proteins and peptides to accelerate the aging of the body ^[26].

189 As can be seen in Table 4, The ethanol extract of <u>Xylaria-X</u> striata has no positive effect on SOD. But the extract at low dose can promote the activity of CAT to reduce the free radicals. Furthermore, the extract can also significantly decrease the content of MDA which is consistent with the content of free radicals. It is implied that the MDA content can also indirectly reflect the content of free radicals and the degree of lipid peroxidation. So, it is conducted that the ethanol extract of <u>Xylaria-X</u> striata keeps body much away from peroxidation and prevent it from aging too quickly.

196 197

Table 4 Effect of Xylaria X. striata on SOD/CAT/MDA

Treatment	Concentration (µM)	Protein content (g/L)	CAT (U/mL)	MDA (nmol/mL)	SOD (U/mg)	Weight (mg/10 flies)
Control	0	1.08±0.16	61.91±5.92	2.65±0.40	43.21±8.5	6.7±0.76
	50	1.32±0.23	87.47±11.34*	0.9±0.27***	34.59±6.71	6.9±0.32
Xylaria <u>X.</u> striata	100	1.37±0.22	69.46±6.94	1.75±0.11**	36.51±6.91	6.6±0.83
othata	200	1.3±0.23	58.49±10.34	1.03±0.26***	29.45±2.91	6.2±0.26
Ascorbic acid	100	1.37±0.23	92.27±13.23*	2.65±0.27	34.11±6.2	5.9±0.07

198 Note: *p < 0.05, **p < 0.01, ***p<0.001

200 5 CONCLUSIONS

201

202 From the previous report that Xylaria X. striata can promote pentobarbital-induced sleep by not only increasing the number of falling asleep and prolonging sleeping time but also reducing sleep latency^[27], this mushroom had potential to be as a functional food used in the 203 204 205 field of geriatrics. Above all, the anti-aging activity of Xylaria-X. striata was evaluated by 206 measuring the survival time under various conditions, the activity of SOD/CAT and the 207 content of MAD/protein of Drosophila-D. melanogaster. The results showed that, although the ethanol extract of Xylaria X. striata could shorten the lifespan of Drosophila D. melanogaster 208 209 under natural conditions which indicated its biologically toxic, it could extend the lifespan of 210 flies in the longevity test of two stress models by repair both ultraviolet radiation damage and 211 hydrogen peroxide oxidative stress damage. The results of enzyme activity and MAD content

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- showed that the ethanol extract of *Xylaria-X_striata* could not only block the source of free
 radicals but also eliminate the reaction products of free radicals.
- 214

215 COMPETING INTERESTS

- 216 Authors have declared that no competing interests exist.
- 217

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