

1                   **APPLICATION OF DIGITAL COLORIMETER FOR PRELIMINARY**  
2                   **CHARACTERIZATION OF GOLD NANOPARTICLE SWARMS PRODUCED BY**  
3                   ***Termitomyces heimii* USING A NOVEL BIOINSPIRED MICROFLUIDICS ASSAY**

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9                   **ABSTRACT**

10                   In our laboratory work extending over several years we have successfully studied the  
11                   biogeochemical role of termite mounds and their occupants the termites and the exosymbiont  
12                   fungus-*Termitomyces*. Fungi appear to be promising for large scale production of nanoparticles  
13                   (NPs) as these are simpler to grow both in laboratory and at industrial scale. This paper reports a  
14                   novel microfluidic based assay system to detect Gold bioreduction capacity of different tissues in  
15                   tissue based and cell free environment. Using sterile microtest wells, different tissues such as  
16                   umbo, pileus, lamellae, stipe context, stipe epicutis, pseudorrhiza context, pseudorrhiza epicutis  
17                   of *Termitomyces heimii* mature fruitbodies were tested with 200µl chloroauric acid (one mM) and  
18                   after an interval of 5, 10, 15, 30, 45, 60, 120 min and 12, 24 and 48 hours. The results in terms  
19                   production of distinct nanoparticles were directly visualized microscopically and using mobile  
20                   based digital colorimeter. Membrane filtered sterile water soluble extracts (SWSE) from the same  
21                   tissues were similarly screened. The results manifested by mono and polydisperse GNPs and

22 microparticles of mixed size groups demonstrated that cell free system can be potentially useful  
23 for bioinspired fabrication of GNPs. Further work in this direction is in progress using several  
24 *termitomyces* pure cultures.

25 Keywords: Microfluidic assay, Gold, Bio reduction, Termitophilic mushrooms

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## 41 **Introduction**

42 Poor and under developed countries like India and resource starved universities find it difficult to  
43 have easy access to expensive instrumentation for characterization of Gold nanoparticles (GNPs).  
44 However now the new Apps such as digital colorimeter from laboratory tools make it possible to  
45 rapidly detect GNP formation and perform quick and simple analysis. Having seen the chemical  
46 creativity of termitophilic mushrooms in our laboratory we aimed to use microfluidic assay for  
47 rapid screening of gold bioreduction system in this species. *Termitomyces heimii* being the most  
48 dominant species and state mushroom of Goa we used this for our study. However, it was not easy  
49 to rapidly detect and characterize swarms of gold micro and nanoparticles. It was at this point that  
50 we came across a mobile based digital colorimeter which was found useful in analysis of the GNP  
51 swarms and obtain spectra in visible band. Swarm are formed by the collective behavior of GNPs.  
52 Inspired by animal interactions, the autonomous movement and collective behavior of synthetic  
53 nanomaterials are of considerable interest as they have implications for the future in  
54 nanomachinery, nanomedicine, and chemical sensing (Kagan et al., 2011). Values such as CIE  
55 LAB, Chroma, Hue°, RGB, color names, real time visible spectra (400 nm to 700 nm) can be  
56 recorded using this digital tool. The CIE LAB color space (also known as CIE L\*a\*b\* or  
57 sometimes abbreviated as simply "Lab" color space) is a color space defined by the International  
58 Commission on Illumination (CIE) in 1976. It expresses color as three values where L\* stands for  
59 the lightness from black to white, a\* from green to red, and b\* from blue to yellow. Chroma  
60 (Saturation) may be defined as the strength or dominance of the hue, the quality of a color's purity,  
61 intensity or saturation (Solomon & Breckon 2011). Hue is common distinction between colors  
62 positioned around a color wheel. On the outer edge of the hue wheel are the intensely saturated  
63 hues whereas towards the center of the color wheel no hue dominates and becomes less and less

64 saturated. The RGB color model is an additive color model in which red, green and blue lights are  
65 added in various ways to reproduce a broad array of colors (Meruga et al., 2014). Finally, it has  
66 been shown that absorbance peaks of GNPs are correlated to their size and we aimed to test the  
67 ability of digital colorimeter to get an idea of size distribution of GNPs in the swarms.

68 Statistical importance of the data was obtained using jvenn, a new JavaScript library  
69 (<http://bioinfo.genotoul.fr/jvenn/example.html>) which processes lists and produces Venn  
70 diagrams. Venn diagrams with more than four lists, are much harder to interpret. To solve this  
71 problem, the classical or Edwards-Venn representation introduces new shapes providing a clearer  
72 view (Philippe Bardou et al., 2014). Jvenn enables to compare up to six lists and updates the  
73 diagram automatically when modifying the lists content.

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## 75 **Methodology**

### 76 **Sample collection**

77 *Termitomyces heimii* being the most dominant species in Goa and state mushroom of Goa we use  
78 this for our study. Fresh, healthy specimens of *Termitomyces heimii* Natarajan (1979) were  
79 collected from fields of Taleigao, Goa during monsoon season, 2019 and taxonomically identified  
80 using standard published *Termitomyces* keys (Heim R. 1942,1977; Natarajan,1979) (**Fig.1**). Dried  
81 herbarium is deposited in Goa university mycological herbarium collection.

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## 85 **Processing of the specimens**

86 *T. heimii* specimens were cleaned with 95% ethanol (v/v) upto 30 seconds and photographed.  
87 Specific processing of each part of the fruitbody ie umbonal tissue, pileus context, lamellae, stipe  
88 context, stipe epicutis, pseudorrhiza context, pseudorrhiza epicutis was carried out. Using sterile  
89 forceps small pieces of the tissues were transferred into a microtest plate (Tarson, Mumbai) with  
90 96 wells having volumetric capacity of 420  $\mu$ l under a laminar air flow bench. Care was taken to  
91 use identical tissue fragments appropriate equivalent to 200  $\mu$ m size. Tissues were tested with 200  
92  $\mu$ l Chloroauric acid (one mM) (**Fig. 2**) and after an interval of 5, 10, 15, 30, 45, 60, 120 minutes  
93 and 12, 24 and 48 hours. Nine replicates of each tissue were used.

## 94 **Preparation of SWSE**

95 Sterile water soluble extracts (SWSE) (**Fig. 3**) were prepared by grinding in sterile mortar with  
96 pestle, centrifuged and membrane filtered (0.22  $\mu$ m pore size, 30 mm diameter-HIMedia  
97 laboratories). The SWSE were stored at refrigerated temperature in sterile test tubes. The extracts  
98 (210  $\mu$ l) and chloroauric acid (210  $\mu$ l) were mixed in equal proportion in the wells of microwell  
99 test plate and checked after interval of 5, 10, 15, 25, 30, 45, 60, 120 minutes and after 12, 24, 48  
100 hours. The assay design was similar to fig.3.

## 101 **Stereomicroscopic visualization of swarms**

102 The microtest plate with the GNP swarms was visualized under stereomicroscope (Olympus SZ51,  
103 model SZ2-ILST, olympus corporation, Tokyo, Japan) (**Fig.4**). Care was taken to bring the swarm  
104 view under uniform illumination in bright light.

## 105 **Use of Digital Colorimeter App**

106 Scanning of the swarms was done with 12 MP plus dual rear (F 1.5/ F 2.4) camera on Samsung  
107 Galaxy Note 9 with colorimeter software (<http://researchlabtools.blogspot.com/>) (Ravindranath et  
108 al, 2018) version 3.5.2, developed by Research Lab Tools, São Paulo, Brazil.

### 109 **Digital color analysis and colorimetric data**

110 The color terminology is used according to color data based of the App. Colorimeter software was  
111 used to record values such as CIE LAB, Chroma, Hue°, RGB, color names, real time visible spectra  
112 (400 nm to 700 nm). The App allows online and offline analysis of samples.

### 113 **Use of Venn diagrams**

114 Venn diagrams are commonly used to display list comparison. However, when the number of input  
115 lists exceeds four, the diagram becomes difficult to read. Alternative layouts and dynamic display  
116 features can improve its use and its readability. The jvenn library accepts three different input  
117 formats “Lists”, “Intersection counts” and “Count lists”. For “Intersection counts”, the lists are  
118 given a label (“A” or “B”) which is used to make the correspondence between the list and its count.  
119 Finally, “Count lists” provide a count number for each element of a list. Hence, with “Count lists”  
120 the figures presented in the diagram correspond to the sums of counts of all elements shared  
121 between lists For “Lists” and “Count lists”, jvenn computes the intersection counts and displays  
122 the chart (<http://bioinformatics.psb.ugent.be/>). Vein diagrams were plotted using tissues and  
123 lambda max values.

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### 125 **Results**

126 Fresh, healthy *Termitomyces heimii* which is dominant species in Goa were successfully obtained  
127 and were taxonomically identified using standard published *Termitomyces* keys (Heim R. 1942,  
128 1977; Natarajan,1979; DeSouza & Kamat, 2018, 2019). GNP swarms were detected in all  
129 treatments and could be visualized easily under stereomicroscope (**Fig.5 & Fig 6**). Umbonal tissue  
130 produced grey GNP swarms and color values as shown in table 1. Overall the color range is from  
131 grey to juniper green. The chromaticity values showed difference and chroma values ranged from  
132 4 to 47 whereas Hue differed from 40 to 199. The R value varied from 131 to 177, G from 102 to  
133 187 whereas B from 27 to 191. Detail treatment of absorbance value is given in table 3. Similarly  
134 the colors and color analysis and absorbance values of other tissues and SWSE are shown in **Table**  
135 **1** and **Table 2**.

136 **Table 3** shows 30 different peaks obtained using each tissue and extract. Absorbance value ranged  
137 from 455 nm to 644 nm. Stipe exhibited most promising results with peaks at 455, 510, 642 nm.  
138 **Table 4** gives approximate GNP size range diameter which ranged from 5 nm to 100 nm. It was  
139 noticed that only extract system was producing GNP swarms at wavelength of 455 nm, whereas  
140 only umbonal tissue produced maximum absorption at 461 nm similar results were obtained in  
141 remaining reaction as shown in table 3. The correlation between the absorbance values verses the  
142 tissue based system and cell free environment is shown (**Fig7a-7f**).

## 143 **Discussion**

144 This paper reports a novel microfluidics based assay system to detect Gold bioreduction capacity  
145 of different tissues in Termitophilic mushrooms (Kalia & Kaur, 2018; DeSouza & Kamat, 2017;  
146 de Souza & Kamat, 2018, 2019) in tissue based and cell free environment. Umbonal tissue, pileus  
147 context, lamellae, stipe context, stipe epicutis, pseudorrhiza context, pseudorrhiza epicutis of  
148 *Termitomyces heimii* mature fruitbodies successfully produced GNPs for the first time. We were

149 successful in producing membrane filtered SWSE from same tissues and also successful in  
150 producing GNPs from the same extracts. Our assay can be useful to carry out large number of  
151 replicates, under sterile forms. Microtest plates can be directly visualized due to its transparent  
152 makeup and swarms can be directly characterized under stereomicroscope. Small amounts of gold  
153 solutions and small amount of SWSE can be tested this microfluidic assay. Using rapid screening  
154 of large number of biological or microbiological gold bioreduction systems.

155 In case of tissues the colour varied from grey, rock blue, ship cove blue, juniper green to  
156 saddle brown where as in case of SWSE it was willow grove, drim grey dark grey, charcoal grey  
157 to brown gramble. For small (~30 nm) monodisperse gold nanoparticles, the surface plasmon  
158 resonance phenomenon causes an absorption of light in the blue-green portion of the spectrum  
159 (~450 nm) while red light (~700 nm) is reflected, yielding a rich red color. As particle size  
160 increases, the wavelength of surface plasmon resonance related absorption shifts to longer, redder  
161 wavelengths (<https://www.sigmaaldrich.com/>). Larger the size, darker is the color and may also  
162 shift to blue in case of colloidal particles (Jana et al., 2001; Haiss et al., 2007; Martinez et al.,  
163 2012). Absorbance reading tells the composition and size of NPs (Doak et al., 2010). SWSE  
164 prepared from same tissue do not produce GNPs of same size or with same concentration. The  
165 GNP swarm population is represented by 18 different size groups ranging from less than 5 nm to  
166 100 nm. The concentration of nanoparticles as function of optical absorbance ranges from 0.12 to  
167 0.8 indicating that some bioreduction systems are much more efficient in production of GNPs this  
168 includes GNPs of the size of less than 5 to 15 nm.

169 It was found that stipe was showing most promising results in case of both tissue and  
170 SWSE. CIE chromaticity values ranged from L (16-75), a (3-33) and b (1-46) thus indicating the  
171 lightness from black to white, green to red and blue to yellow (Cheng et al.,2014). Choma values



172 ranged from 4 to 47 and hue values from 40 to 199 indicating the strength or dominance of the  
173 hue, the quality of a color's purity, intensity or saturation.

174 GNP's produced using tissue showed low absorbance from min 0.10 to max 0.5 whereas  
175 SWSE produced GNP with absorbance ranging from 0.1 to 0.8 thereby indicating a more efficient  
176 system in cell free environment. Low absorbance 0.19 for intact tissue indicating low concentration  
177 of GNP swarms. Mean absorbance produced by treatment with extract 0.40 indicating almost  
178 double the bioreduction efficiency of intact tissue in case of GNP production. Thus, cell free  
179 environment is much better system to produce polydisperse GNP swarms in higher concentration.

180 In all 30 different peaks ranging from 455 to to 644 nm were obtained using each tissue  
181 and extract thus indicating the presence of nanoparticle of size 5 nm to 100 nm  
182 (<https://www.sigmaaldrich.com>). Mushrooms are rich in proteins and have high availability of the  
183 amino acids lysine, tryptophan, glutamic acid and aspartic acid (Hsu et al., 2002). It is also reported  
184 that certain mushroom extract contain polysaccharide/oligosaccharide complex (Cho et al., 2003).  
185 FTIR studies have also shown the possible biomolecules responsible for capping and efficient  
186 stabilization of the metal nanoparticles synthesized using mushroom extract (Philip, 2009). It was  
187 noticed certain SWSE and tissues produced specific wavelength for example umbonal, pileaus  
188 context, pseudorrhizal context extract system was producing GNP swarms at wavelength of 455  
189 nm and only umbonal tissue produced at 461 nm. Similar results were obtained in remaining  
190 reaction as shown in table 3. It was noticed that there is relationship between solubility and swarm  
191 formation and it could be a different molecule based bioreduction system.

192 Preliminary characterization of GNP swarms is important as in high throughput screening  
193 system one cannot differentiate the most promising system and it can be time consuming. Once  
194 you carry out the preliminary results you zero down to the specific system to obtain the promising

195 system and then can go for final characterization of the GNPs. Preliminary results can also help in  
196 standardization of the procedure and also there is no waste of resources. In poor and under  
197 developed countries like India and resource starved universities researchers find it difficult to have  
198 easy access to expensive instrumentation for characterization of GNPs. However now the new  
199 Apps such as digital colorimeter from developers Laboratory tools make it possible to rapidly  
200 detect GNP formation and perform quick analysis.

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### 203 **Conclusions**

204 Our work clearly demonstrates that simple and easy to use mobile digital Colorimeter Apps can  
205 be used for primary optical characterization of swarms of Gold nanoparticles. This is useful in  
206 rapid screening of large number of microbiological gold bioreduction systems. The spectral  
207 absorbance profile detected in visible range also helps in understanding the presumptive size of  
208 GNPs in swarms. For high throughput screening systems we recommend development of more  
209 such mobile based apps. Our present approach has helped us to fabricate a very sensitive Gold  
210 biosensor. Pure mycelial cultures of *T. heimii* also produced identical results. This would be  
211 published separately.

212

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217 Thanks for guidance from Dr. Absar Ahmad director, interdisciplinary center for Nanotechnology  
218 AMU regarding potential of GNPs.

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## 221 **References**

222 Bardou, P., Mariette, J., Escudié, F. et al. jvenn: an interactive Venn diagram viewer. BMC  
223 Bioinformatics. 15: 293 (2014). <https://doi.org/10.1186/1471-2105-15-293>

224 Cheng, X., Dai, D., Yuan, Z., Peng, L., He, Y. and Yeung, E.S., 2014. Color difference  
225 amplification between gold nanoparticles in colorimetric analysis with actively controlled  
226 multiband illumination. Analytical chemistry. 86(15): 7584-7592.

227 Cho, J., Kang, J.S., Long, P.H., Jing, J., Back, Y. and Chung, K.S., 2003. Antioxidant and memory  
228 enhancing effects of purple sweet potato anthocyanin and cordyceps mushroom extract. Archives  
229 of pharmacal research. 26(10): 821-825.

230 de Souza, R.A. and Kamat, N.M., 2018. Evaluation and characterization of pellet morphology of  
231 genus *Termitomyces heim* of a wild tropical edible mushroom. Journal of Pharmaceutical,  
232 Chemical and Biological Sciences. 6(4): 320-328.

233 de Souza, R.A. and Kamat, N.M., 2019. *Termitomyces* holomorph benefits from anomalous  
234 Sulphur content in teleomorph. International Journal of Life sciences Research. 1 (186-192).

235 De Souza, R.A., Kamat, N.M. and Nadkarni, V.S., 2018. Purification and characterisation of a  
236 sulphur rich melanin from edible mushroom *Termitomyces albuminosus* Heim. Mycology. 9(4):  
237 296-306.

238 Doak, J., Gupta, R.K., Manivannan, K., Ghosh, K. and Kahol, P.K., 2010. Effect of particle size  
239 distributions on absorbance spectra of gold nanoparticles. Physica E: Low-dimensional Systems  
240 and Nanostructures. 42(5): 1605-1609.

241 D'Souza, R.A. and Kamat, N.M., 2017. Potential of FTIR spectroscopy in chemical  
242 characterization of *Termitomyces* Pellets. Journal of Applied Biology & Biotechnology. 5(04):  
243 080-084.

244 Haiss, W., Thanh, N.T., Aveyard, J. and Fernig, D.G., 2007. Determination of size and  
245 concentration of gold nanoparticles from UV– Vis spectra. Analytical chemistry.79(11): 4215-  
246 4221.

247 Heim, R., 1942. Nouvelles études descriptives sur les agarics termitophiles d'Afrique tropicale.  
248 Arch. Mus. Natl. Hist. Nat. Paris. 6: 1-133.

249 Heim, R., 1977. Termites et champignons; les champignons termitophiles d'Afrique noire et d'Asie  
250 meridionale.

251 Hsu, T.H., Shiao, L.H., Hsieh, C. and Chang, D.M., 2002. A comparison of the chemical  
252 composition and bioactive ingredients of the Chinese medicinal mushroom Dong Chong Xia Cao,  
253 its counterfeit and mimic, and fermented mycelium of *Cordyceps sinensis*. Food chemistry. 78(4):  
254 463-469.

255 <http://bioinfo.genotoul.fr/jvenn/example.html>

256 <http://bioinformatics.psb.ugent.be/>

257 <http://researchlabtools.blogspot.com>

258 <https://www.sigmaaldrich.com>

259 <https://www.tedpella.com>

260 Jana, N. R., Gearheart, L., & Murphy, C. J. (2001). Seeding growth for size control of 5– 40 nm  
261 diameter gold nanoparticles. *Langmuir*. 17(22): 6782-6786.

262 Kagan, D., Balasubramanian, S., and Wang, J. (2011). Chemically triggered swarming of gold  
263 microparticles. *Angewandte Chemie International Edition*, 50(2): 503-506.

264 Kalia, A. and Kaur, G., 2018. Biosynthesis of Nanoparticles Using Mushrooms. In *Biology of*  
265 *Macrofungi*. 351-360. Springer, Cham.

266 Meruga, Jeevan Manikyarao, Aravind Baride, William Cross, Jon J. Kellar, and P. Stanley May.  
267 "Red-green-blue printing using luminescence-up conversion inks." *Journal of Materials*  
268 *Chemistry C* 2, no. 12 (2014): 2221-2227.

269 Natarajan, K., 1979. South Indian Agaricales V: *Termitomyces heimii*. *Mycologia*. 71(4): 853-855.

270 Philip, D., 2009. Biosynthesis of Au, Ag and Au–Ag nanoparticles using edible mushroom extract.  
271 *Spectrochimica Acta Part A: Molecular and Biomolecular Spectroscopy*. 73(2): 374-381.

272 Sen, I.K., Maity, K. and Islam, S.S., 2013. Green synthesis of gold nanoparticles using a glucan of  
273 an edible mushroom and study of catalytic activity. *Carbohydrate polymers*. 91(2): 518-528.

274 Solomon, C. and Breckon, T., 2011. *Fundamentals of Digital Image Processing: A practical*  
275 *approach with examples in Matlab*. John Wiley & Sons.

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293 **Figures and Tables**

294 **Fig.1.** Scheme for specific processing of different fruitbody parts

295 **Fig.2.** Design of Microfluidics based assay using Microtest plate

296 **Fig.3.** Homogenized aqueous extracts from different tissues

297 **Fig.4.** Direct Acquisition of images for stereomicroscopic characterization

298 **Fig.5.** Positive bioreduction obtained with homogenized tissues as indicated by color changes

299 **Fig.6.** Positive bioreduction with cell free membrane filtered SWSE indicated by change in colour,  
300 yellow is control

301 **Fig.7 (a-f).** 7(a-c): The shape corresponding to the lists involved in the intersection are highlighted  
302 in case of tissue, (a-455-463, b-510-540, c-632-644  $\lambda_{max}$ ). 7(d-f): The shape corresponding to the  
303 lists involved in the intersection are highlighted in case of extracts, (a-455-463  $\lambda_{max}$ , b-510-540,  
304 c-632-644  $\lambda_{max}$ ).

305

## 306 **Tables**

307 Table 1 & 2: Colour analysis and Colorimetric absorption characterization of presumptive GNP  
308 swarms

309 Table 3: Visible spectral characteristics of GNP swarms using *T. heimii* tissue sample and SWSE

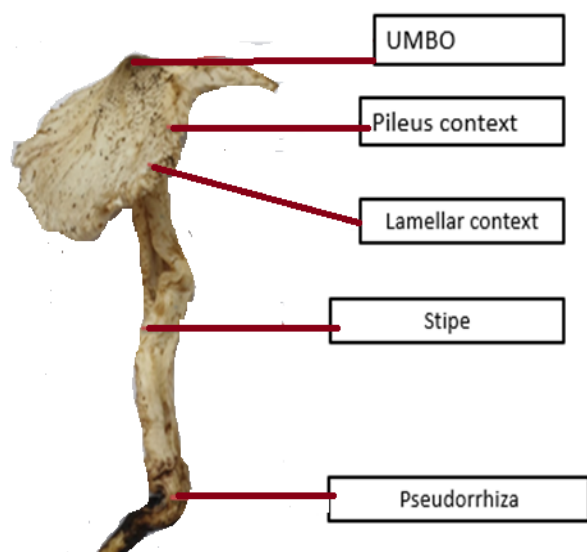
310 Table 4: Approximate size range of GNP swarms produced using *T. Heimii*

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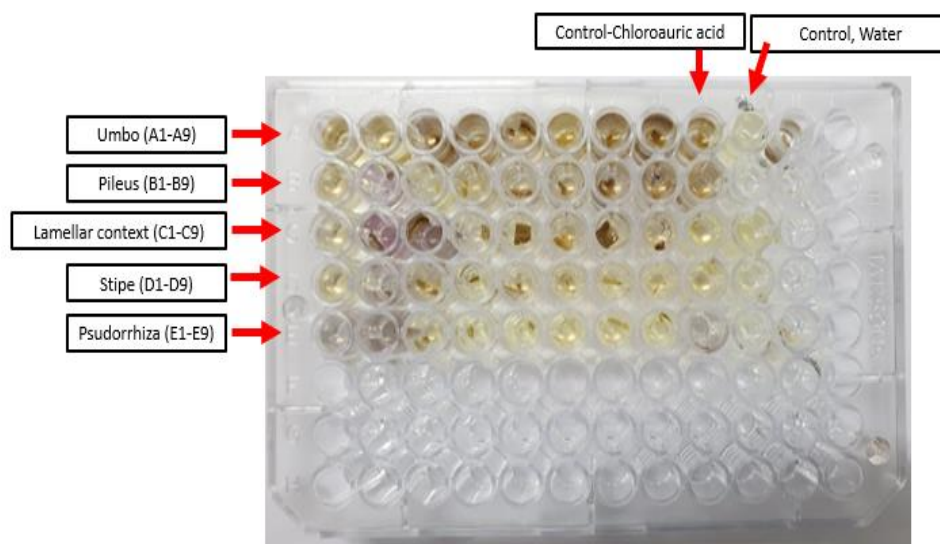
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Fig.1. Scheme for specific processing of different fruitbody parts



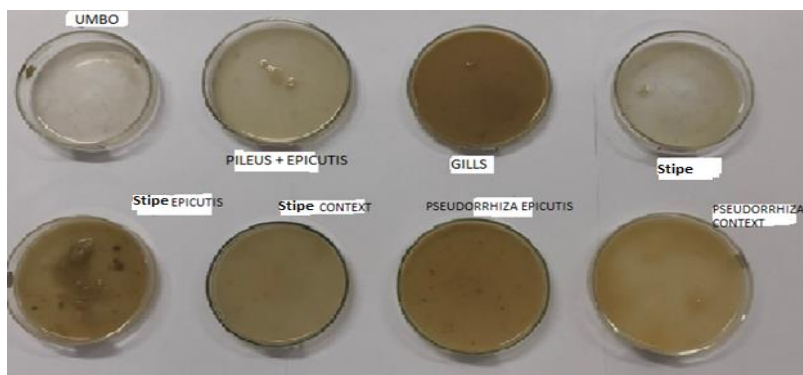
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Fig.2. Design of Microfluidics based assay using Microtest plate

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Fig.3. Homogenized aqueous extracts from different *T. heimii* fruit body tissue

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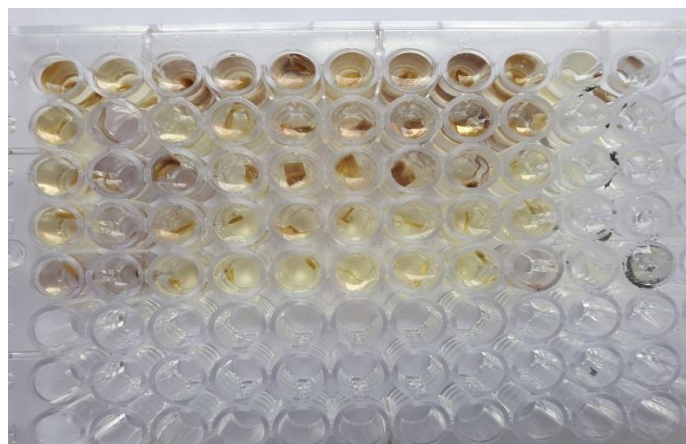


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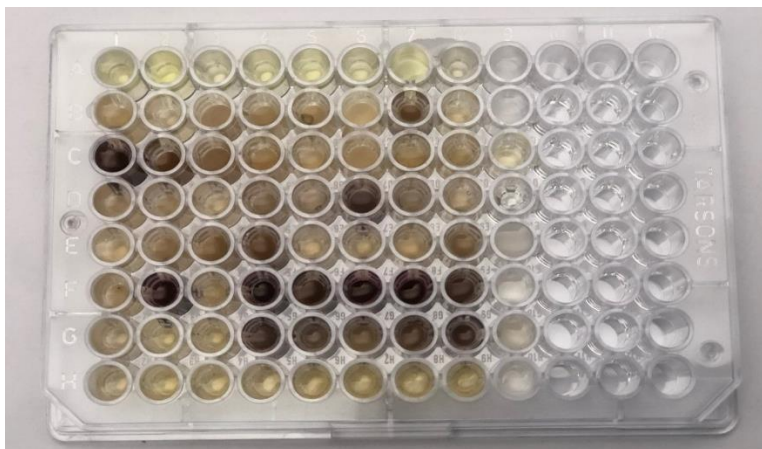
Fig.4. Direct Acquisition of images for stereomicroscopic characterization

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327 **Fig.5.** Positive bioreduction obtained with homogenized tissues as indicated by color changes



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329 **Fig.6.** Positive bioreduction with cell free membrane filtered SWSE indicated by change in  
330 colour, yellow is control

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Fig.7a

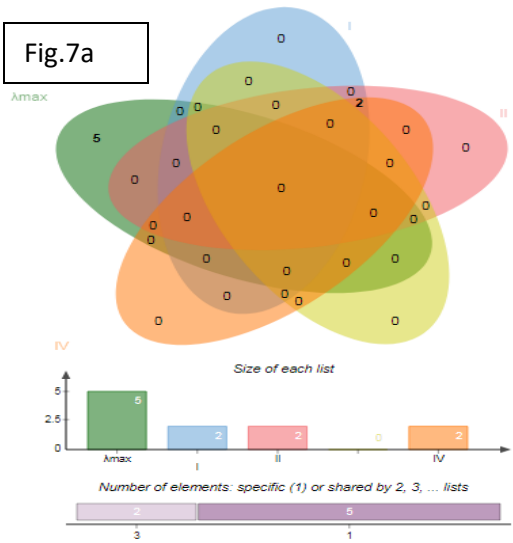


Fig.7b

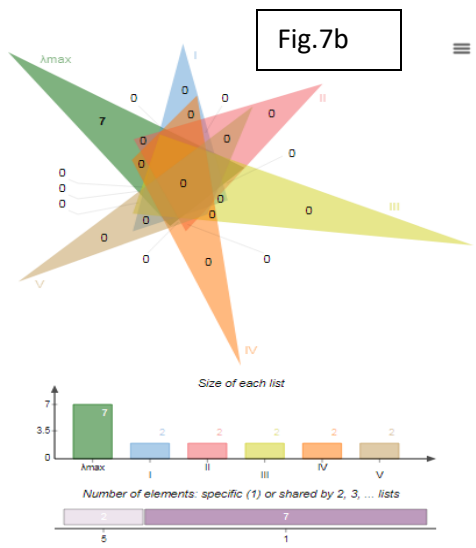


Fig.7c

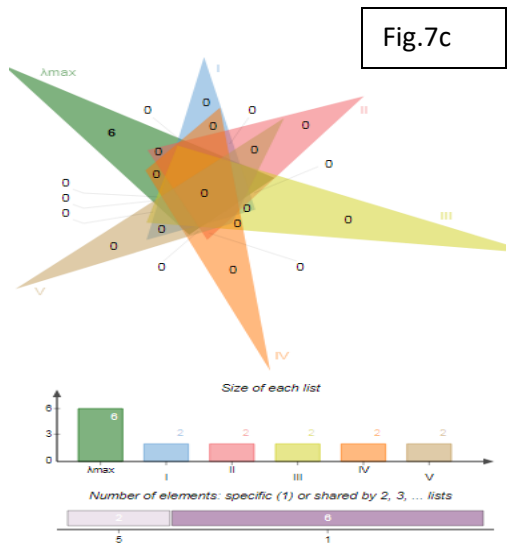


Fig.7d

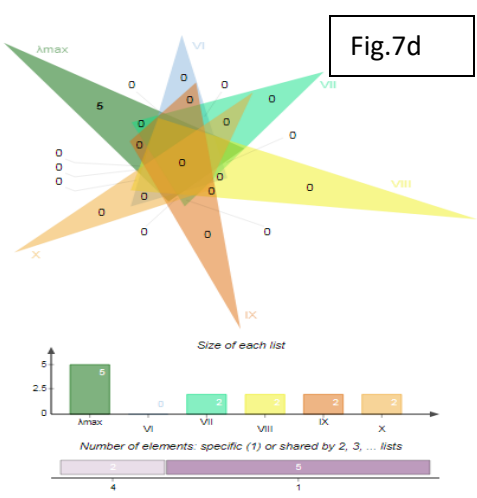


Fig.7e

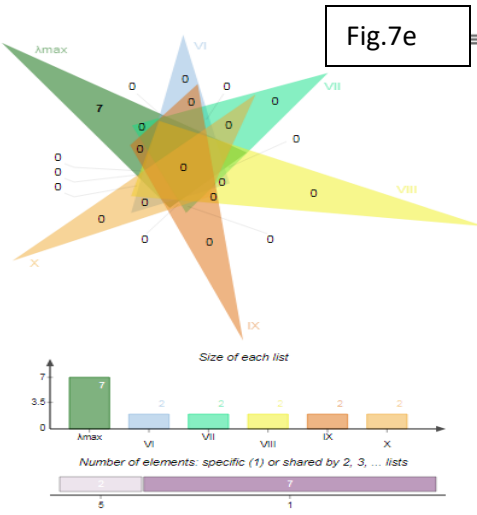
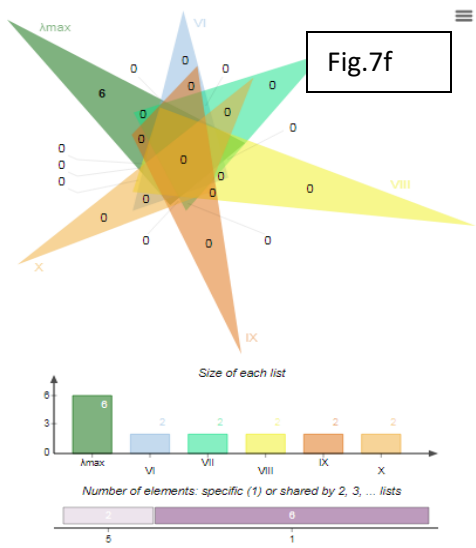

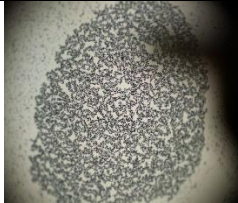
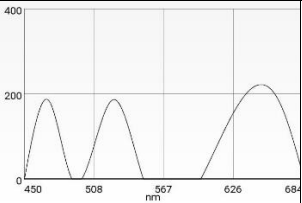


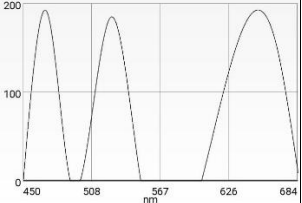

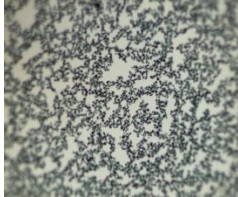
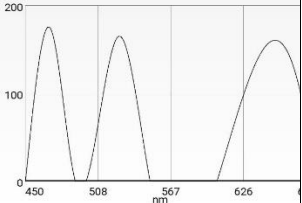


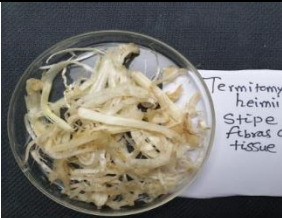

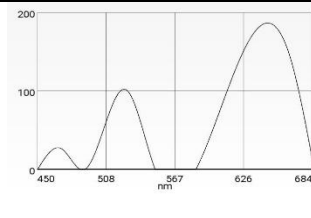


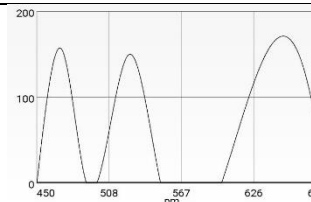
Fig.7f



**Fig.7 (a-f):** 7(a-c)- The shape corresponding to the lists involved in the intersection are highlighted in case of tissue, (a-455-463  $\lambda_{\max}$ , b-510-540,c-632-644). 7(d-f)- The shape corresponding to the lists involved in the intersection are highlighted in case of extracts (a-455-463  $\lambda_{\max}$ ,b-510-540,c-632-644).


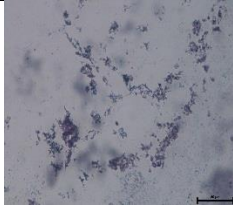
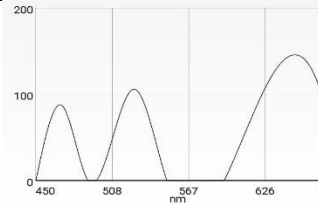

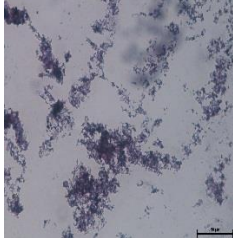
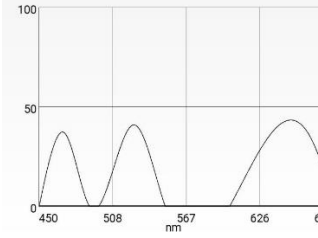

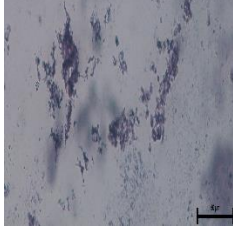
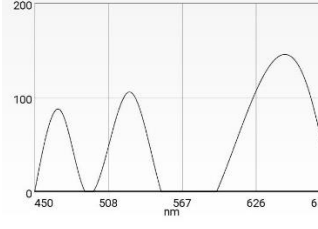
**Table 1:** Colour analysis and Colorimetric absorption characterization of presumptive GNP swarms

Mushroom Tissues	GNPs swarms (X 6000)	Colour	CIE Chromaticity values	Chroma	Hue	R, G, B value	Absorbance (nm)	Absorbance values
		Grey	L=75 a=4 b=1	4	174	177, 187, 186		463 (0.18), 520 (0.18), 641(0.22)
		Rock Blue	L=73 a=9 b=6	11	190	156, 185, 191		462(0.19), 522(0.18), 642(0.19)
		Ship Cove Blue	L=66 a=10 b=9	13	193	131, 165, 174		462(0.17), 520(0.16), 641(0.15)

		Saddle Brown	L=46 a=9 b=46	47	40	141, 102, 27		460(0.5), 520(0.10), 643(0.18)
		Juniper Green	L=61 a=3 b=5	6	199	137, 150, 156		540 (0.15), 519 (0.14), 642(0.17)

**Table 2:** Colour analysis and Colorimetric absorption characterization of presumptive GNP swarms

Mushroom Tissues	GNPs swarms (X 6000)	Color	CIE Chromaticity values	Chroma	Hue	R, G,	Absorbance (nm)	Absorbance values

						B value		
		Willow Grove	L=45, b=12	12	42	114, 106, 87		461(0.8), 519 (0.11), 640 (0.15)
		Dark Grey	L=16, b=2	4	140	35, 41, 37		455(0.2), 512(0.3), 632 (0.4)
		Charcoal grey	L=33 a=8 b=7	11	111	67, 79, 65		455(0.8), 511(0.11), 632(0.15)


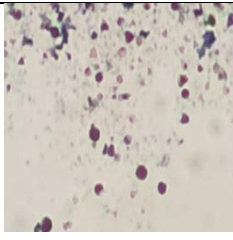
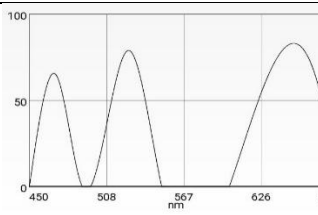

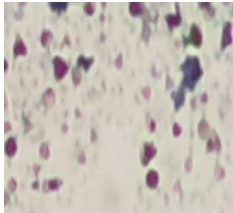
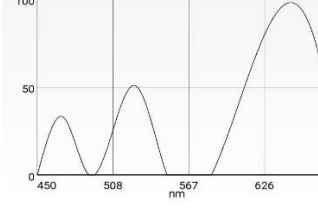
 <p>Stipe context</p>		<p>Brown Bramble</p>	<p>L=24 a=8 b=16</p>	<p>18</p>	<p>27</p>	<p>74, 51, 33</p>		<p>460(0.6), 522(0.8), 644(0.7)</p>
 <p>Pseudo context</p>		<p>Dim grey</p>	<p>L=42 a=5 b=5</p>	<p>7</p>	<p>96</p>	<p>94, 100, 90</p>		<p>455 (0.2), 510(0.5), 642(0.10)</p>



Table 3: Visible spectral characteristics of GNP swarms using *T. heimii* tissue sample and SWSE

Absorbance wavelength (nm)	Tissue					SWSE				
	I Umbonal tissue	II Pileus context	III Lamellae	IV Stipe context	V Pseudorrhiza context	VI Umbonal context	VII Pileus context	VIII Lamellae	IX Stipe context	X Pseudorrhiza context
455	-	-	-	-	-	-	+	+	-	+
460	-	-	-	+	-	-	-	-	+	-
461	+	-	-	-	-	-	-	-	-	-
462	+	+	-	-	-	-	-	-	-	-
463	+	-	-	-	-	-	-	-	-	-
510	-	-	-	-	-	-	-	-	-	+
511	-	-	-	-	-	-	-	+	-	-
512	-	-	-	-	-	-	+	-	-	-
519	-	-	-	-	+	+	-	-	-	-
520	+	-	+	+	-	-	-	-	-	-

522	-	+	-	-	-	-	-	-	+	-
540	-	-	-	-	+	-	-	-	-	-
632	-	-	-	-	-	-	+	+	-	-
640	-	-	-	-	-	+	-	-	-	-
641	+	-	+	-	-	-	-	-	-	-
642	-	+	-	-	+	-	-	-	-	+
643	-	-	-	+	-	-	-	-	-	-
644	-	-	-	-	-	-	-	-	+	-

Table 4: Approximate size range of GNP swarms produced using *T. Heimii*

<b>Wavelength reported in present assay (nm)</b>	<b>Approximate GNP size range (nm)</b>	<b>Reference</b>
455, 460, 461, 462, 463	< 5	Ted Pella Inc.
510, 511	5	Ted Pella Inc.
512, 519	10	Sigma-Aldrich
520, 522	15	Sigma-Aldrich, Ted Pella Inc.
540	10-30	Radtsig et al., 2016
632	50-80	Ted Pella Inc.
640	80-100	Sigma-Aldrich
641	100	Sigma-Aldrich
642, 643, 644	100	Sigma-Aldrich, Ted Pella Inc.