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3 **Total Phenols, Antioxidant Capacity and**

4 **Antibacterial Activity of Manuka Honey Extract**

5

6 **ABSTRACT**

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Aims: To evaluate total phenols content (TPC), antioxidant capacity (TAC) and antibacterial activity of Manuka honey extract (MHE) and to compare such properties with those for unfractionated Manuka honey.

Study design: *In vitro* study

Place and Duration of Study: School of Biomedical Sciences, Ulster University, Coleraine, UK. Between September 2016 and September 2017.

Methodology: MHE was prepared by solvent extraction using ethyl acetate. TPC was determined by Folin-Ciocalteu assay. The iron (III) reducing antioxidant capacity (IRAC) method was used to determine TAC. Antibacterial activity was evaluated using disc diffusion assay and 96-well microtiter plate method with absorbance measured at 600 nm..

Results: The TPC for MHE was 30-fold higher than the value for Manuka honey (33420 ± 1685 mg vs. 1018 ± 78 mg GAE/kg) while TAC values were ~100-times greater ($83,198 \pm 7064$ vs. 793 ± 104 TEAC, respectively). Antibacterial activity assessed by disc diffusion for Manuka honey (18.5mm on *S. aureus* and 20mm on *E. coli*) was two times greater than for MHE (9mm for both *S. aureus* and *E. coli*). The 96-well microtiter plate assay confirmed the greater antibacterial activity for Manuka honey compared to equal concentrations MHE.

Conclusion: A polyphenol-rich Manuka honey extract with a high total antioxidant capacity, showed little or no antibacterial activity against *E. coli* and *S. aureus* in contrast with unfractionated Manuka honey.

8 *Keywords: Manuka honey, Manuka extract, total phenols, antioxidant, antibacterial,*

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11 **1. INTRODUCTION**

12 Infectious diseases continue to pose a threat to human health worldwide. Wound infections

13 are commonly caused by bacterial pathogens [1-3]. Owing to the rising occurrence of

14 antibiotic-resistant bacterial strains, alternative ancient remedies and plant-based products

15 such as honey are being evaluated for therapeutic use. The medicinal importance of honey

16 has been widely documented in the world's medical literature; standardized active Manuka

17 honey has been registered as a wound care product with appropriate medical regulatory

18 bodies [1-3]. Sherlock et al [4] demonstrated antibacterial activity for Chilean Honey (Ulmo

19 90 honey) and New Zealand Manuka honey (UMF® 25+) against 5 MRSA strains. Ahmed

20 and Othman [5] found that Tualang honey and Manuka honey could inhibit growth of Gram-

21 positive MRSA strains including *S. aureus* and *S. pyogenes* and Gram-negative strains like *P.*

22 *aeruginosa*, *E. coli* and *Enterobacter cloacae* [5].

23

24 Kwakman and Zaat [6] reported the antibacterial activity for Revamil™ honey and Manuka

25 honey, the two leading medicinal grade honeys, arose from different mechanisms involving,

26 hydrogen peroxide, bee defensin, methylglyoxal, and unidentified components. Several

27 studies proposed that phenolic compounds may contribute to the non-peroxide antibacterial

28 activity of Manuka honey [1, 6]. However, the specific contribution of phenolic components to

29 the antibacterial action of Manuka honey has not been well investigated. The general aims

30 of this project were to evaluate, the total phenols content (TPC), total antioxidant capacity
31 (TAC), and antibacterial activity of an ethyl acetate extract from Manuka honey (Manuka
32 Honey Extract; MHE) and to compare these characteristics with unfractionated Manuka
33 honey.

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36 **2. MATERIAL AND METHODS**

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38 **2.1 Materials**

39 Manuka honey rated Unique Manuka Factor (UMF) 10+, 15+ and 18+ were purchased from
40 Comvita Ltd (UK). Ethyl acetate, Folin & Ciocalteu reagent, sodium carbonate, gallic acid,
41 Trizma base, ferrozine (3-(2-Pyridyl)-5,6-diphenyl-1,2,4-triazine-4',4''-disulfonic acid sodium
42 salt), ammonium iron (III) sulfate dodecahydrate, methanol and other chemicals were
43 purchased from Sigma-Aldrich Ltd (Gillingham, UK). Nutrient broth (Oxoid), nutrient agar
44 (Oxoid), and penicillin-streptomycin mixture (Pen-strep) were purchased from ThermoFisher
45 Scientific (UK). Bacteria strains (*Staphylococcus aureus*, *Escherichia coli*) were obtained
46 from School of Biomedical Sciences, Ulster University (UK).

47

48 **2.2 Preparation of Manuka honey extract (MHE)**

49 Honey extract was prepared using ethyl acetate as solvent as described by Tan et al [7]
50 with modification. Manuka honey (UMF 10+, 20g) was dispersed in 80ml of distilled-
51 deionized water and 100ml ethyl acetate. The mixture was stirred using a magnetic stirrer for
52 24 hours. The emulsion formed was transferred to glass centrifuge tubes and centrifuged at
53 2,000 RPM for 15 minutes. The non-aqueous ethyl acetate phase was air-dried and the
54 residue formed was re-dissolved in methanol solvent, filtered through 0.2 µm, and then
55 stored at -18°C until used. The solids content of MHE extract was determined by drying 50
56 µl of MHE and weighing the residue.

57

58 **2.3 Determination of total phenols content**

59 The total phenols content was determined using Folin-Ciocalteu method adapted for
60 microplate analysis as described previously [8, 9].

61

62 **2.4 Determination of antioxidant capacity**

63 Antioxidant capacity was determined using the iron (III) reducing antioxidant capacity
64 (IRAC) as described recently [8]. The IRAC reagent was prepared by dissolving 20 mg
65 ferrozine dye in 9ml of Tris buffer and adding ferric (III) ammonium sulphate (4mg in 1ml
66 water). For TAC determinations, samples of honey (20µl) were mixed with 280 µl of
67 ferrozine solution and incubating for 30 min at 37 °C. TAC assay were calibrated using
68 trolox (0-1000µM) as antioxidant standard. A microplate reader (VersaMax, Molecular
69 Devices, Sunnyvale, California, USA) was used for absorbance measurements at 562nm.

70

71 **2.5 Antibacterial activity screening**

72 **2.5.1. Antibacterial screening by disc diffusion assay**

73 The disc diffusion assay for antibacterial activity was carried out as described previously
74 with minor modification [10] using two bacteria strains, one Gram-positive bacteria- *S. aureus*
75 and one Gram-negative bacteria- *E. coli*. Working in laminar flow hood, bacterial inoculated
76 broth (200 µl) was transferred to blank nutrient agar plates (each bacteria x2), and allowed to
77 dry. Thereafter, 6 blank paper discs were transferred onto agar plates using tweezers and
78 ensuring equal spacing between each. Samples (20µl) of Manuka honey (UMF10+ Manuka
79 honey extract, 25% UMF10+, 15+, 18+ Manuka honey), MHE and controls (Pen-strep.) were
80 slowly added to the blank disc and were left to dry briefly. Plates were incubated upside
81 down overnight at 37°C. The diameter of zones of inhibitions was measured after 24 hours
82 in mm.

83

84 **2.5.2 Micro-plate assay for antibacterial activity**

85 Samples of a 24-h grown liquid culture (50µL) were transferred to 96-well microtiter plate
86 (x3), and 50µL of sterile honey sample (25% w/v in water), MHE or antibiotic (Pen-strep)
87 was added. The 96-well microtiter plates were then incubated at 37°C for 24h with gentle
88 shaking and absorbances were read at 600nm. The antibacterial effect (%) was determined
89 from the expression, $100 \cdot (1 - (A_H/A_0))$ where A_H and A_0 are absorbance readings with and
90 without honey treatment..

91

92 **2.6 Data analysis**

93 Data analysis was performed using Microsoft excel and IBM SPSS Statistics Version 24.
94 Correlation was obtained by Pearson correlation and significance was assessed in two-
95 tailed at level of significance of $P= 0.01$.

96

97

98 **3. RESULTS AND DISCUSSION**

99

100 **3.1 Total phenols content and antioxidant capacity of Manuka honey extract**

101 The total phenols content (TPC) for MHE and unfractionated honey were expressed in mg-
102 Gallic acid equivalent (GAE) per kg and are shown in Table 1. The TPC for Manuka honey
103 UMF10+ was 1018.32 ± 78.84 mg GAE/kg honey (Table 1) compared with values in the
104 range of 430 - 2706 mg GAE/kg reported previously [11]. The TPC for honey is known to
105 vary with various factors, including monoflorality of honey, age and geographic origins of
106 honey samples [12]. Compared to the original honey the, MHE had a 32.8-fold increase
107 TPC. The TAC for honey evaluated in terms of IRAC is shown in Table 1 for ethyl acetate
108 honey extract, and original honey (Table 1). The TAC for MHE was 105-fold
109 increase compared to unfractionated Manuka honey. We reported that the TPC for Manuka
110 honey was strongly correlated with antioxidant capacity and UMF rating (UMF5+, 10+, 15+
111 and UMF18+) [9]. but there was no specific reason for working with UMF10+ on this
112 occasion.

113 Moniruzzaman et al [12], Alvarez-Suarez et al [13] and Can et al. [14] reported that phenols
114 play an important role in the antioxidant capacity of honey. The dominating phenolic
115 components identified in Manuka honey were phenyllactic acid and a group of methoxylated
116 benzoic acids [7, 9]. Other components in Manuka honey that contribute to the antioxidant
117 capacity include flavonoids such as chrysin, quercetin, isorhamnetin and luteolin, phenolic
118 acid like gallic acid, caffeic acid and syringic acid [13]

119 **Table 1. Total phenols content and total antioxidant capacity of Manuka honey**
120 **(UMF10+) and the Manuka honey Extract**

121

Sample	TPC (mg GAE/kg)	TAC(mg TE/kg)
Honey (UMF10+)	1018 ± 79	793 ± 104.4
MHE	33420 ± 1685	83198 ± 7064

122 *The total phenols content (TPC) and total antioxidant capacity (TAC) for Manuka honey (UMF10+)
123 and ethyl acetate extract from Manuka honey (MHE) as mg Gallic acid Equivalent (GAE)/ kg or mg
124 Trolox Equivalent (TE)/ kg sample.

125

126

127 **3.2 Antibacterial activity**

128 Table 2 shows the antibacterial activity using the disc diffusion assay for *S. aureus* and *E.*
129 *coli*. The MHE was adjusted to deliver similar quantities GAE per paper disc as was used for
130 Manuka honey. For example, paper discs were loaded with 20-µl Manuka honey UMF10+

135 (250g/l) and hence the GAE loading per disc was $(250\text{g/l} \cdot 20 \times 10^{-6}\text{l} \cdot 1.108 \times 10^{-3}\text{g GAE/g}) =$
 136 $5.1 \times 10^{-6}\text{ gGAE}$. After pre-diluting MHE by 33-fold, the GAE loading per disc was
 137 $(7.5\text{g/l} \cdot 2 \times 10^{-6}\text{l} \cdot 33.420 \times 10^{-3}\text{g GAE/g}) = 5.0 \times 10^{-6}\text{ gGAE}$. The results show clearly that the honey
 138 had antibacterial activity while MHE showed little or no antibacterial activity. (Fig. 1).
 136
 137

Table 2. Measurement of zone of inhibition (mm) by disc diffusion assay

Bacteria	Zone of inhibition (mm)					
	Controls		Honey Samples			
	Pen-Step	Methanol	UMF10+	UMF15+	UMF18+	MHE
<i>S. aureus</i>	27.0	9.0	18.5	18.5	20.5	9.0
<i>E. coli</i>	26.0	9.5	20.0	21.0	22.5	9.0

139 *MH= Manuka honey, Tests involve 20 µl of 25% (w/v) added to paper discs. MHE = Manuka honey
 140 extract from UMF10+ honey.
 141

149 Antibacterial activity testing using microplate/ spectrophotometric method also showed that
 150 the honey extract had no antibacterial activity when tested at concentrations similar to those
 151 in honey (Figure 1). For these tests, the exposure concentrations for 10+ Manuka honey and
 152 MHE were 0.76 mM and 0.74 mM GAE, respectively. The microorganism used for testing (*S.*
 153 *aureus* and *E. coli*) are known to be sensitive to Manuka honey [5, 16]. Therefore, it was
 154 expected that screening (by disc diffusion assay and spectroscopic analysis) showed
 155 antibacterial activity with Manuka honey. Interestingly, MHE showed little or no antibacterial
 156 activity when tested at a similar concentration as honey. Indeed, *S. aureus* measurements
 157 were consistent with increased growth after exposure to MHE (Fig. 1).
 150

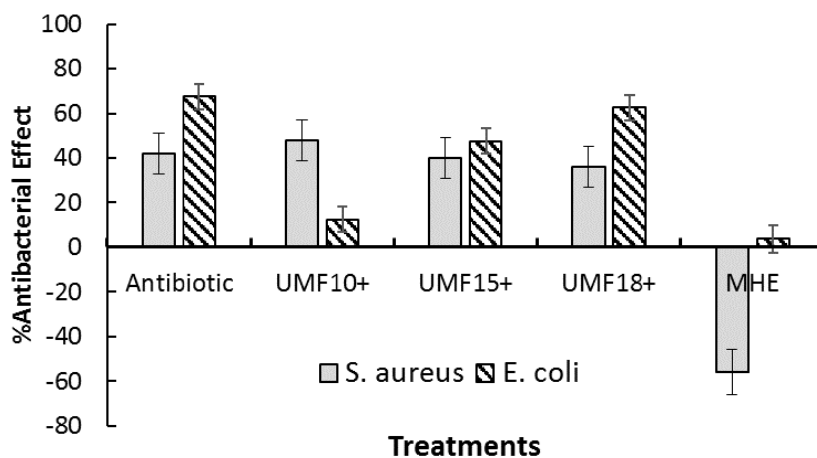


Fig.1. Antibacterial effect for Manuka honey and Manuka honey extract (MHE)

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 152
 155 Tested with 96-well microtiter plates with *E. coli* or *S. aureus*. Penicillin-streptomycin was
 156 used as +ve control. Honey samples were 12.5% (rated UMF 10+ -UMF18+). MHE is
 157 Manuka Honey Extract See text for details.
 156

164 Based on current results, the polyphenols from Manuka honey may not be a major factor
 165 contributing to the antibacterial activity. The findings agree with previous reports, which
 166 noted that Manuka honey polyphenols (benzoic acids, cinnamic acids and flavonoids) could
 167 not account for entirely for the observed antibacterial activity [17, 18]. Alternatively, the
 168 concentration of phenolic compounds from honey maybe too low to contribute to
 169 antibacterial activity [6]. By contrast, there is considerable evidence showing a correlation
 170 between methylglyoxal content and antibacterial activity of Manuka honey [19] while only
 171 slight antibacterial activity was ascribed to the high sugar content and acidity [6]. The low

164 antibacterial activity ascribed to MHE reported in the current paper, is different from the
165 results obtained extracts from 30 local honeys from Saudi Arabia which showed that
166 antibacterial activity was strongly correlated with total phenols content and antioxidant
167 power [15] but there are differences in the approaches adopted for these studies. Manuka
168 honey contains comparable levels of polyphenols and methylglyoxal [20], further research is
169 underway to determine if such components interact and if this has possible consequences on
170 antibacterial activity. The behavior of extracted phenols may also be different from
171 antibacterial effects observed in the presence of the honey components. Further
172 investigations are underway to consider the chemical constituents of Manuka honey that
173 contribute to its antibacterial activity using the pathogenic strains used in this study.
174

175 **4. CONCLUSION**

176 Manuka honey organic extract, containing a high total phenols content and total antioxidant
177 capacity showed little or no antibacterial activity. Further in-depth research is needed to
178 understand the composition and characteristics of Manuka honey extracts.
179

180 **CONSENT**

181 No consent required

182

183 **ETHICAL APPROVAL (WHEREEVER APPLICABLE)**

184 No ethical approval required

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187

188 **REFERENCES**

189

- 190 1. McLoone P, Warnock M, Fyfe L. Honey: A realistic antimicrobial for disorders of the
191 skin. *J Microbiol Immunol Infect.* 2016 49(2):161-7.
- 192 2. Carter DA, Blair SE, Cokcetin NN, Bouzo D, Brooks P, Therapeutic Manuka honey: no
193 longer so alternative. *Front Microbiol.* 2016 7: 569.
- 194 3. Oryan A, Alemzadeh E, Moshiri A. Biological properties and therapeutic activities of
195 honey in wound healing: a narrative review and meta-analysis. *J Tissue*
196 *Viability.* 2016;25(2):98-118.
- 197 4. Sherlock O, Dolan A, Athman R, Power A, Gethin G, Cowman S, Humphreys H.
198 Comparison of the antimicrobial activity of Ulmo honey from Chile and Manuka honey
199 against methicillin-resistant *Staphylococcus aureus*, *Escherichia coli* and
200 *Pseudomonas aeruginosa*. *BMC Compl Alt Med.* 2010;2:10(1):47.
- 201 5. Ahmed S, Othman NH. Review of the medicinal effects of Tualang honey and a
202 comparison with Manuka honey. *MJMS.* 2013;20(3):6.-13.
- 203 6. Kwakman, PHS, Zaat, SA.J. Antibacterial components of honey. *lubmb Life.* 2012 64(1):
204 48-55.
- 205 7. Tan ST, Holland PT, Wilkins AL, Molan PC. Extractives from New Zealand honeys. 1.
206 White clover, Manuka and Kanuka unifloral honeys. *J Agric Food Chem.* 1988.36(3):
207 453-460.

- 208 8. Owusu-Apenten R, Mohd Yusof H I, Nigam PS. Cryptic total antioxidant capacity of
209 Manuka honey measured by anew microplate assay for iron (III) reducing antioxidant
210 capacity (IRAC)and other methods. Food Chem. (2017): Submitted
- 211 9. Portokalakis I, Mohd Yusof HI, Ghanotakis D, Nigam P, Owusu-Apenten R , Manuka
212 Honey-induced cytotoxicity against MCF7 breast cancer cells is correlated to total
213 phenol content and antioxidant power.JABB.2016. 8(2): p. 1-10.
- 214 10. Barlow R, Barnes D, Campbell A, Nigam PS, Owusu-Apenten R. Antioxidant,
215 anticancer and antimicrobial, effects of Rubia cordifolia aqueous root extract. JABB.
216 2015;5(1):6-14.
- 217 11. Stephens JM, Schlothauer RC, Morris BD, Yang D, Fearnley L, Greenwood DR,
218 Loomes KM. Phenolic compounds and methylglyoxal in some New Zealand Manuka
219 and Kanuka honeys. Food Chem.2010;120(1): 78-86.
- 220 12. Moniruzzaman M, Sulaiman SA, Khalil MI, Gan SH. Evaluation of physicochemical and
221 antioxidant properties of sourwood and other Malaysian honeys: a comparison with
222 Manuka honey. Chem Cent J. 2013;7(1):138.
- 223 13. Alvarez-Suarez JM, Gasparrini M, Forbes-Hernández TY, Mazzoni L, Giampieri F. The
224 composition and biological activity of honey: A focus on Manuka honey.Foods.
225 2014;3(3):420-432.
- 226 14. Can Z, Yildiz O, Sahin H, Turumtay EA, Silici S, Kolayli S. An investigation of Turkish
227 honeys: their physico-chemical properties, antioxidant capacities and phenolic profiles.
228 Food Chemistry. 2015 Aug 1;180:133-41.
- 229 15. Al-Hindi RR, Bin-Masalam MS, El-Shahawi MS. Antioxidant and antibacterial
230 characteristics of phenolic extracts of locally produced honey in Saudi Arabia. Int J
231 Food Sci Nutr. 2011;62(5):513-7.
- 232 16. Alzahrani HA, Alsabehi R, Boukraâ L, Abdellah F, Bellik Y, Bakhotmah BA.
233 Antibacterial and antioxidant potency of floral honeys from different botanical and
234 geographical origins.Molecules. 2012;17(9):10540-9.
- 235 17. Weston RJ, Mitchell KR, Allen KL. Antibacterial phenolic components of New Zealand
236 Manuka honey. Food Chem. 1999;64(3):295-301.
- 237 18. Weston RJ, Brocklebank LK, Lu Y. Identification and quantitative levels of antibacterial
238 components of some New Zealand honeys. Food Chem. 2000; 70(4):427-435.
- 239 19. Atrott J, Henle T. Methylglyoxal in Manuka honey—correlation with antibacterial
240 properties. Czech J Food Sci.2009;27(Spec.):S163-S165.
- 241 20. Kwok TH, Kirkpatrick G, Mohd Yusof HI, Portokalakis I, Nigam P, Owusu-Apenten R.
242 Rapid colorimetric determination of methylglyoxal equivalents for Manuka honey.
243 JABB. 2016;7(1):1-6.
- 244