Upregulation of heat-shock proteins in larvae, but not adults, of the flesh fly during hot summer days

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- 3 Up-regulation of heat-shock proteins in larvae, but not adults, of the flesh fly during hot
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15 **Abstract** Heat-shock proteins (HSPs) are highly expressed when organisms are exposed to 16 thermal stresses. The HSPs are considered to play significant roles in thermal adaptation 17 because they function as molecular chaperones facilitating proper protein synthesis. The 18 expression of HSPs under field conditions, however, has not been evaluated much, and their 19 importance, based on the ecological contexts in nature, is still unclear. We investigated this 20 aspect in the larvae and adults of the flesh fly, Sarcophaga similis. These larvae spend their 21 larval life in the carrion or faeces of vertebrates; therefore, they are less mobile and are 22 occasionally exposed to high temperature. In contrast, the adults of this species can fly and, 23 therefore, they are highly mobile. Massive transcription of Hsps was detected both in the larvae 24 and adults in a laboratory heat shock experiment. The larvae in the field showed no or less Hsp 25 production on thermally mild days, whereas considerable upregulation of Hsp expression was 26 detected on days with high temperature. The adults can also be exposed to thermal stress as high 27 as 40°C or higher in the field. However, most of the flies showed no or less Hsp expression. The 28 observations in the experimental cage under field conditions revealed behavioural 29 thermoregulation of adults through microhabitat selection. The present study demonstrates 30 ontogenetic alteration of the strategy to overcome thermal stress in an insect; in the field, less 31 mobile larvae use physiological protection against heat (HSP production), whereas highly 32 mobile adults avoid the stress behaviourally (through microhabitat selection).

33

34 Key words Behavioural thermoregulation • Flesh fly• Heat-shock protein • Mobility •
35 Natural thermal stress

36

37 Introduction

38 The expression of heat-shock proteins (HSPs) is highly induced when organisms, including 39 insects, are exposed to environmental stresses, such as extremes of temperatures. HSPs function 40 as molecular chaperones, facilitating proper synthesis and preventing aggregation and 41 denaturation of other proteins. Thus, in many organisms, HSPs have been considered to play an 42 important role in survival under thermal stress (Parsell and Lindquist 1993; Yiangou et al. 1997; 43 Feder and Hofmann 1999; King and MacRae 2015). In insects, four major families of HSPs are 44 recognized, namely the small heat shock proteins (sHSPs), HSP60, HSP70, and HSP90. The 45 sHSPs, acting independently of ATP, are the first line of cell defence, preventing irreversible 46 denaturation of substrate proteins and facilitating subsequent refolding by ATP-dependent 47 chaperones. The remaining HSPs interact with proteins and promote protein folding, 48 degradation, disaggregation, and localization in an ATP-dependent manner (Basha et al. 2012; 49 Clare and Saibil 2013; King and MacRae 2015). The primary function of HSP70 is to bind to 50 unfolded or partially unfolded proteins to prevent their aggregation and to release them for 51 folding. HSP90 also prevents aggregation of non-native proteins, but it appears to be more 52 selective for substrates compared to other general chaperones (Clare and Saibil 2013). 53 Laboratory studies have demonstrated a clear relationship between HSP production and

acquisition of heat tolerance. For example, when larvae of the fruit fly *Drosophila melanogaster* are exposed to severe heat shock, the majority of animals die. If, however, a mild heat shock, which induces the production of HSPs, is applied immediately before a severe heat shock, approximately 50% of the animals survive, because they have become thermotolerant or protected (Mitchell et al. 1979). Similar results were also observed in other organisms (Patriarca and Maresca 1990; Denlinger et al. 1991; Gehring and Wehner 1995). Adaptive changes in HSP expression over seasons (Fader et al. 1994; Hofmann and Somero 1995; Minier et al. 2000;

61 Dieterich et al. 2013) also support the ecological significance of HSPs in natural populations. 62 Engineering of gene expression further verified the significance of HSPs in thermotolerance in 63 insects (Feder et al. 1996; Feder and Krebs 1998; Rinehart et al. 2007; Colinet and Hoffmann 64 2010; Koštál and Tallarová-Borovanská 2009; Lü and Wan 2011). Thus, HSPs are the 65 candidates for playing a significant role in adaptation in natural populations. However, despite 66 HSPs being extremely important for survival following heat shock under laboratory conditions, 67 their ecological relevance and adaptive importance under field conditions has only been rarely 68 investigated directly and is less clear (Sørensen 2010). Especially, some insects with high 69 mobility possibly regulate their body temperature to avoid "overheating" by microhabitat 70 selection (May 1979; Chown and Nicolson 2004). This implies that less mobile insects, which 71 are unable to escape from detrimental environments, may rely on physiological mechanisms, 72 such as HSP production to protect themselves from heat damage. On the contrary, highly 73 mobile insects, which can avoid overheating behaviourally, may not rely on such physiological 74protection, because HSP production is costly and sometimes deleterious (Huang et al. 2007; 75 Krebs and Feder 1997; Hoekstra and Montooth 2013). However, this issue has been paid little 76 attention (Sørensen 2010, but see Feder et al. 1997, 2000).

77 Here, we investigated relationships between insects' mobility and HSP production in 78 larvae and adults of the flesh fly, Sarcophaga similis, in the laboratory as well as in the field. 79 Adult males of this species occur from late April to the end of November in Osaka, Japan 80 (Tachibana and Numata 2006). This species is the most dominant sarcophagid species in our 81 research field (the campus of Osaka City University; Tachibana and Numata 2006). It is 82 important to note that Osaka City is now one of the hottest cities in Japan, and August is the 83 hottest month in Osaka (Japan Meteorological Agency, 2017; see Fig. S1). The adult males of S. 84 similis establish temporal territories on sunlit places, wait for a female to pass, and frequently 85 fly to pursue females and begin copulation in flight, as observed in other sarcophagid species.

86	After copulation, S. similis females carries the egg until hatching and then deposits the newly
87	hatched 1st instar larvae (ovoviviparous) on carrion or animal faeces. The entire feeding phase
88	of larval development is spent within the carrion or faeces, and when feeding is completed, the
89	larvae leave the food (wandering stage), burrow into the soil, and pupariate and pupate. In the
90	present study, we investigated the levels of expression of Hsp70, Hsp23, and Hsp83 in the
91	larvae and adults in the laboratory. Thereafter, we investigated their expression in the field-
92	collected larvae and adults. We also examined the behavioural thermoregulation of adults in the
93	experimental cage in the field.
94	
95	Materials and methods
96	
97	Insect rearing
98	
99	The laboratory colony of S. similis originally collected on the campus of Osaka City University,
100	Osaka, Japan (34.6°N, 135.5°E), in 2013, was maintained at 25°C under long-day conditions
101	(LD 16:8 h). The adult flies were provisioned with water and sugar, and fed on a piece of beef
102	liver 2 days after the emergence of adults. Ten days later the females larvipoisted on the liver,
103	and the larvae were reared on a new piece of liver under the same environmental conditions.
104	The larvae used for the experiments were not sexed, whereas only males were used for
105	experiments on adult flies.
106	

107 Hsp23 sequence

109	The nucleotide sequences of S. similis Hsp70 and Hsp90 were found in the database
110	(DDBJ/GenBank/EMBL accession no. LC176075 and AB196477, respectively), but that of
111	Hsp23 was not found. Therefore, we cloned the gene and identified its sequence. A DNA
112	fragment of S. similis Hsp23 was obtained by RT-PCR. An adult male was exposed to 41°C in a
113	water bath for 15 min, and total RNA was isolated from the whole body with Trizol Reagent
114	(Invitrogen, Carlsbad, CA, USA). The cDNA was synthesized from the RNA with M-MLV
115	reverse transcriptase and oligo $(dT)_{12-13}$ primers (Invitrogen). PCR was performed using GoTaq
116	DNA Polymerase (Promega, Madison, WI, USA). The primers, which were designed from the
117	conserved Hsp23 nucleotide sequence among several dipteran species, were as follows;
118	hsp23cl-F, 5'-CGT CAT CGA GGG CAA GCA YGA RGA RMG-3' and hsp23cl-R, 5'-GGC
119	GGG GCC GGY YTG YTG DAT YT-3'. The amplified fragment, approximately 300 bp in
120	length, was purified with Wizard Plus SV Minipreps DNA Purification System (Promega) and
121	sequenced on 3130 Genetic Analyzer (Applied Biosystems, Foster City, CA, USA) with
122	BigDye Terminator v3.1 Cycle Sequence Kit (Applied Biosystems). The nucleotide sequence of
123	S. similis Hsp23 was deposited in the database (DDBJ/GenBank/EMBL accession no.
124	LC150013).
105	

126 Laboratory heat shock

128 The feeding 3rd (final) instar larvae (3 days after larviposition) and adult males (5 days after 129 adult emergence) were exposed to various temperatures for 15 min in water baths in the 130 laboratory.

131

132 Field observations of larvae

133

134	Approximately 100 feeding 3rd instar larvae in 100–200 g of beef liver were placed on sunlit
135	ground on 14-, 15-, 19-, and 20-Aug-2014. The probes ($\phi 6 \text{ mm} \times 3 \text{ cm}$) of the data logger (TR-
136	71wf, T and D, Matsumoto, Japan) were inserted approximately 2 mm beneath the top surface
137	and 2 mm above the bottom surface of the liver to measure the temperature where larvae reside.
138	Since larvae stayed in the liver during the experiments, the body temperatures of larvae were
139	considered to fall within the range of these two temperatures. The temperatures in these two
140	positions and air temperature near the liver were recorded at intervals of 15, 30, and 60 min
141	from around 10:30 to around 17:30 h (JST).
142	Two or three larvae were removed from the liver every 15, 30, and 60 min, and total
143	RNA was isolated from single larvae to assess gene expression.
144	
145	Field observation of the adults
146	
147	From 16-Jul-2013 to 25-Sep-2013 and from 22-Jul-2014 to 9-Oct-2014, the body-surface
148	temperatures (surface temperature of the thorax) of the S. similis adult males attracted by a piece
149	of beef liver in the field were measured using an infrared thermographic camera (i3, FLIR,
	7

150	Wilsonville, OR, USA). The air temperatures around the flies were also measured using the
151	temperature data logger (TR-71wf). We adopted body-surface temperature in this experiment,
152	because of the difficulty in the measurement of immediate body temperatures of the flies in the
153	field. It was difficult to identify the species of the flying sarcophagid males in the field, but it
154	was reported that S. similis was the most dominant sarcophagid species on this field area
155	(Tachibana and Numata 2006). To confirm this, we collected 95 males after body-surface
156	temperature measurements and observed their genitalia under the stereoscopic microscope in
157	our laboratory. We confirmed that all of the males that we captured were S. similis. Thus, we
158	measured the body-surface temperature and isolated RNAs by immediate homogenization from
159	28 flies without species identification. The observed data were fitted to a non-linear regression
160	line with the nonlinear least squares curve fitting
161	(http://www.colby.edu/chemistry/PChem/scripts/lsfitpl.html).
162	
163	Body-surface temperature and body temperature of adults
164	
165	To see the difference between the body-surface temperature and body temperature (inner
166	temperature of the thorax), we simultaneously monitored these temperatures in the male flies in

- 167 the laboratory. A type K thermocouple (φ 0.127 mm; ST-55K-TC-1.2M, Graphtec, Yokohama,
- 168 Japan) was inserted into the thorax of a fly. To simulate the direct sunlight exposure, the flies
- 169 were placed under a reflector lamp (500W) at a distance of 10–15 cm and their body-surface
- 170 and body temperatures were monitored simultaneously with a data logger (midi Logger GL220,
- 171 Graphtec) and a thermographic camera.

- **qPCR**

175	Total RNA was isolated with Trizol Reagent. After treatment of the total RNA with
176	deoxyribonuclease I (amplification grade; Invitrogen), cDNA was synthesized from each total
177	RNA (0.3–2.0 µg) by using High Capacity cDNA Reverse Transcription Kit (Applied
178	Biosystems). For quantitative real-time PCR analysis with a 7500 Real-Time PCR system
179	(Applied Biosystems), 1 μ L of cDNA was used in a final volume of 1X Go Taq qPCR Master
180	Mix (Promega) and 0.2 μ M of each primer, according to the supplier's instructions. Each
181	reaction was performed in duplicate. As a control gene, 28SrRNA was used for normalization
182	(Rinehart et al. 2006; de Boer et al. 2009; Udaka et al. 2010). The primers used were: hsp70qrt-
183	F (5'-TCT TGG TTG GCG GTT CTA-3') and hsp70qrt-R (5'-CCA TAG GCA ACT GCT TCA
184	TC-3') for Hsp70, hsp83qrt-F (5'-TGA GCC CAA GAT TGA AGA TG-3') and hsp83qrt-R (5'-
185	CTT GTG TGT AGG CAA CCT TA-3') for Hsp83, hsp23qrt-F (5'-GGT TAT ATC TCA CGG
186	CAC TTT-3') and hsp23qrt-R (5'-GGA ACA CTT ACG GTG AGA AC-3') for Hsp23 and
187	28Sqrt-F (5'-CCG ATG AAC CTG AAT ATC CAT T-3') and 28qSrt-R (5'-AGG TTT TGA
188	TAC CCA ATA ACT TGC-3') for 28SrRNA. The amplified fragments were approximately 100
189	bp in length. In all the reactions, the generation of only a single expected amplicon was
190	confirmed by performing the melting-curve analysis. Quantification was performed by the
191	standard curve methodology with known amounts of the DNA fragments amplified by PCR.
192	The primers used to amplify the fragment for the standard curve analysis were hsp70rt-F (5'-
193	CCC GTT TCG AAG AAT TGT GT-3') and hsp70rt-R (5'-ACC GCC TGC TGT TTC AAT
194	AC-3') for hsp70, hsp83rt-F (5'-GAA CGC GAC CAA GAG GTT AG-3') and hsp83rt-R (5'-
195	CAC CAT ATT CGG CTT GTG TG-3') for hsp83, hsp23rt-F (5'-ATG AGG AAC GCG AAG
196	ACG-3') and hsp23rt-R (5'-CTG GCG CTC ATT GGA TTT-3') for hsp23, and 28Srt-F (5'-

197 CCG ATG AAC CTG AAT ATC CAT T-3') and 28Srt-R (5'-GGT TTT GAT ACC CAA TAA
198 CTT GC-3') for 28SrRNA.

199

200 Sun-shade preference of the adults

201

202	On 29-Aug-2014 and 6- and 9-Sep-2014, we investigated the sun-shade preference of S. similis
203	males on the campus of Osaka City University. Five-day-old males were individually placed in
204	a framed cage (40-cm width, 20-cm depth, 15-cm height) with all its faces covered with a nylon
205	mesh to allow air circulation into the cage from outside. One half of the cage was shaded using
206	aluminium foil to make a shaded part. The cage was set 1-m above the sunlit ground. The
207	position of the male, i.e., in the sunlit or shaded part of the cage, was observed every 2 min, and
208	simultaneously, the body-surface temperature was measured with an infrared thermographic
209	camera. The air temperature in the cage was also recorded with the temperature data logger.
210	Eighteen flies were used in this experiment.
211	
212	Results

213

214 Larval Hsp expression in the laboratory

215

The larvae were exposed to a variety of high temperatures for 15 min in the laboratory and the relative abundance of *Hsp70*, *Hsp23*, and *Hsp83* transcripts were measured (Fig. 1). *Hsp70* mRNA was almost undetectable at the normal rearing temperature, but its level was

219	considerably increased at higher temperatures. Massive transcription started at 32°C (a 17-fold
220	increase was observed with the increase in temperature from 25 to 32°C). The highest transcript
221	level was observed at 40°C and the level in this case was 479-fold higher than the level prior to
222	the heat shock (25°C). <i>Hsp23</i> also responded to the heat shock and much mRNA accumulation
223	was observed at 36-44°C. Because of weak expression at the rearing temperature, the maximum
224	level of expression was 15-fold higher than the level prior to the heat shock. <i>Hsp83</i> was
225	constitutively expressed at the rearing temperature and also responded to the heat shock, and
226	thus, only a 2.4-fold change in expression was noted.

228 Field observation of the larvae

229

230 We placed the larvae on a piece of beef liver on sunlit ground and measured the air 231 temperature and the temperature inside the liver in August 2014 (Fig. 2). Although both the air 232 temperature and temperature inside the liver gradually increased toward mid-day and decreased 233 toward evening in all these days, the exact temperatures were greatly variable among the 234 different days. On cloudy days of 14- and 15-Aug, the maximum air temperature never reached 235 40°C, and the top surface temperature of the liver was 1–5°C lower than the air temperature and 236 was as high as 38°C. In contrast, it was bright on 19- and 20-Aug. On these days, the maximum 237 air temperatures were much higher than that on 14- and 15-Aug. The top surface temperature of 238 the liver was nearly identical to the air temperature on 19-Aug and reached 45°C during mid-239 day. The highest temperature of the bottom of the liver was as high as 40°C. Air temperature 240 change on 20-Aug was similar to that on 19-Aug, but the top surface of the liver reached 50°C. 241 Among each observation, we found 2-5 dead larvae at the end of the experiments on 19- and 242 20-Aug, but no such larvae was found on 14- and 15-Aug.

243	The larvae inside the liver were collected every 15, 30, and 60 min and the expression
244	of Hsp was investigated (Fig. 2). On the mild days of 14- and 15- Aug, Hsps70 expression
245	remained at low levels and never reached the highest mean level observed in the laboratory heat
246	shock experiment. Similarly, Hsp23 and Hsp83 expression remained at low levels on these
247	days, although some individuals showed slightly higher expression levels during mid-day. In
248	contrast, on the severe days of 19- and 20-Aug, considerable upregulation of these Hsp genes
249	was noted during mid-day. The highest levels of Hsp70, Hsp23, and Hsp83 mRNA were 4.8-,
250	36.0-, and 14.6-times higher than those observed in the laboratory heat shock experiment. The
251	larvae in the field suffered extensive heat shock on severe days.
252	
253	Body temperature and body-surface temperature of adults
254	
255	We measured the body temperature (temperature inside the thorax) and body-surface
256	temperature (surface temperature of the thorax) of the adult flies simultaneously under a
257	reflector lamp in the laboratory, to see how these temperatures were different (Fig. 3). Upon
258	increasing the body-surface temperature, the body temperature rose linearly. The body surface
259	temperature was found to be 2.3 ± 1.7 °C higher than the body temperature in this experimental
260	setup.
261	We measured the air temperature and the body-surface temperatures of free-living adult
262	males in the field (Fig. 4). When air temperatures were higher, the body-surface temperatures
263	were also higher, in general. However, when air temperature was higher than approximately
264	37°C the increment rate of body-surface temperatures became smaller. Thus, free living flies
201	57 C, the merement rate of body-surface temperatures became smaller. Thus, net-inving files

265 seem to maintain their body-surface temperature as high as 40° C even under severe thermal

266	conditions. The highest air temperature and the highest body-surface temperature that we
267	observed were 48.1 and 44.5°C, respectively.

269	Adult Hsp	expression in	the laboratory
10/	munt mp	capi coolon m	the habor atory

270

271 The adults were exposed to various temperatures for 15 min in the laboratory and relative

amounts of *Hsp70*, *Hsp23*, and *Hsp83* transcripts were measured (Fig. 5, left panels). The

273 expression patterns of all the three *Hsps* were mostly similar. The highest expression levels of

274 *Hsp70*, *Hsp23*, and *Hsp90* were detected at 40, 42, and 42°C, respectively, and they were 58-,

275 44-, and 3.7-fold higher than the levels prior to heat shock (25 °C).

276

277 Field observation of the adults

278

279	We captured flies in the field just after measurement of their body-surface temperatures by an
280	infrared thermographic camera, and investigated their Hsp70, Hsp23, and Hsp83 expression
281	(Fig. 5, right panels). The body-surface temperatures ranged from 29.3°C to 44.5°C and the
282	interquartile range were $36.5-41.2^{\circ}C$ (median = $39.9^{\circ}C$). Despite the higher body-surface
283	temperatures, strong expression of <i>Hsps</i> was not observed; only a few individuals showed the
284	expression comparable to that in the laboratory heat shock experiment and others showed
285	almost undetectable levels of <i>Hsp</i> mRNAs. Such low expression levels in the adults in the field
286	were greatly different from the expression in the larvae in the field (Fig. 2).

287

288 Sun-shade preference of the adults

289

290	We released an adult male into the experimental cage, half of which was covered with
291	aluminium foil to form a shaded part, and the cage was placed on sunlit ground. We observed
292	the position of the fly in the cage and measured air temperature and body-surface temperature of
293	the fly every 2 min, to clarify the relationship among these factors. The total number of flies that
294	were in a sunlit or shaded part are shown in Figure 6A. Flies preferred to stay in the shaded part
295	when the air temperature was higher, whereas they preferred to stay in the sunlit part when the
296	air temperature was lower (Mann-Whitney U test, $P < 0.05$). It is also important to note that no
297	significant difference was detected when the body-surface temperatures of the flies in sunlit and
298	shaded parts were compared (Mann-Whitney U test, $P > 0.05$; Fig. 6B).

299

300 Discussion

301

Our laboratory experiments showed that *Hsp70* and *Hsp23* are heat-inducible in *S. similis*similar to their expression in other dipteran species (Goto and Kimura 1998; Yocum et al. 1998;
Rinehart et al. 2000; Tachibana et al. 2005; Lopez-Martinez and Denlinger 2008; Concha et al.
2012). *Hsp90* is constitutively expressed but it also responds to heat shock. Such expression
pattern is also reported in other fly species (Rinehart and Denlinger 2000; Chen et al. 2005;
Tachibana et al. 2005).

308 Although the expression of Hsps has been examined in various organisms, most of the 309 studies focus on the physiological mechanisms of the heat-shock response or on the comparison 310 of the response among species to find the significance of HSPs in their thermotolerance 311 (Hoffmann et al. 2003; King and MacRae 2015). We still lack an ecological context on whether 312 Hsp expression is fine-tuned to naturally-encountered heat shock (Sørensen 2010). In the 313 present study, we found considerable upregulation of these genes in S. similis larvae in the field 314 on severe days (19- and 20-Aug-2014), when air and liver temperatures exceeded 40–45°C. The 315 environmental conditions on these days were so severe that the larvae protected themselves 316 against thermal stress by expressing large amounts of Hsps. Feder et al. (1997) reported similar 317 results in D. melanogaster. The larvae collected from the necrotic fruit that were on the sunlit 318 ground in the field accumulated large amounts of HSP70 protein and the levels were high in 319 comparison to the levels after standard heat shock in the laboratory. Lopez-Martinez and 320 Denlinger (2008) also investigated heat-shock response of the apple maggot *Rhagoletis* 321 pomonella in the field. This species lays eggs on an apple and the larvae spend their larval life 322 inside it. The larvae are frequently exposed to summertime apple temperatures that exceed 323 40°C. The field temperature cycles ranging from 16 to 47°C elicit strong Hsp70 and Hsp83 324 expression. Thus, the natural heat shock is sufficiently intense to induce HSP expression in 325 these dipteran larvae.

326 The expression of all the Hsps was also greatly enhanced by laboratory heat shock in 327 the S. similis adults, as observed in the larvae. However, drastic induction was scarcely detected 328 in adults in the field, despite their high body-surface temperatures. These results indicate that 329 the free-living adults in the field do not rely on HSP expression to survive in the field in contrast 330 to larvae, and further imply that the adult flies were scarcely damaged by hyperthermia in the 331 field. Feder et al. (2000) investigated the heat-shock response in the field-captured D. 332 melanogaster adults. The levels of HSP70 in most of the flies captured on warm days were very 333 low and comparable to the levels previously reported in unstressed flies in the laboratory. Flies 334 showed frequent responses only when they were caged and placed in direct sunlight. These

335 results indicate that even on warm days most of the flies avoid thermal stress, presumably 336 through microhabitat selection. Behavioural thermoregulation to avoid overheating of body 337 temperature has been reported in various ectotherms (Kleckova et al. 2014; Sunday et al. 2014). 338 Thermoregulation of S. similis adults was also evident in the present study. The increment in the 339 body-surface temperature was suppressed at higher environmental temperatures in the field and 340 flies maintained their body-surface temperatures as high as 40°C. Based on the laboratory 341 experiments, a 15-min exposure to 40° C induced considerable upregulation of *Hsp* mRNAs, but 342 such increased expression was not observed in the field. These results indicate that the body 343 temperature is far lower than the temperatures that induce the *Hsp* expression. Furthermore, the 344 present study clarified that the adult flies frequently fly from shade to sunlight or vice versa and 345 this behaviour maintains their body-surface temperatures at a certain level. Although adult 346 males of S. similis are also exposed to sun heat for many hours in the field, they may regulate 347 their body temperatures so as not to be affected with the extreme thermal stress by frequent 348 movement between sunlit and shade places, as suggested in *D. melanogaster* (Feder et al. 2000). 349 Here, we clarified that, in the field, the flesh fly larvae with low ability to escape from

their habitat protect themselves from thermal damage by expressing *Hsps*. On the other hand, adults with high ability to escape from heat stress by flying can regulate their body temperatures and, therefore, they do not have the necessity of the expression of *Hsps*. The flesh flies appear change their strategy, during ontogeny, from physiological protection against heat damage to behavioural protection against heat exposure.

355

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359

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480 **Figure legends**

481

482 Fig. 1 Relative amounts of *Hsp70* (top), *Hsp23* (middle), and *Hsp83* (bottom) mRNAs in

- 483 Sarcophaga similis larvae that were exposed to various temperatures for 15 min in the
- 484 laboratory. *n*= 3–4 for each temperature. The highest mean levels of *Hsp70*, *Hsp23*, and *Hsp83*
- 485 mRNA at 40, 38, and 44°C, respectively, were set at 1.0. Lines were drawn through the means
 486 of biological replicates

487

488 Fig. 2 Air temperature, temperatures of a piece of beef liver in which Sarcophaga similis larvae 489 reside, and relative amounts of Hsp70, Hsp23, and Hsp83 mRNA in the larvae, on 14-, 15-, 19-, 490 and 20-Aug-2014. A piece of beef liver in which the larvae reside were placed on sunlit ground 491 on each day and air temperature (closed squares) and temperatures of top (open triangles) and 492 bottom (closed triangles) surfaces of the beef liver were recorded (top panels). The larvae were 493 taken out from the liver and the relative amounts of Hsp70, Hsp23, and Hsp83 mRNAs were 494 measured. The highest mean expression levels of each gene in the laboratory heat shock 495 experiment (40, 38, and 44°C for Hsp70, Hsp23, and Hsp83, respectively; Fig. 1) were set at 1.0 496 and are shown in the horizontal grey bars. The expression levels in 21–37 larvae were plotted 497

- 498 **Fig. 3** Body-surface and body temperatures of *Sarcophaga similis* adults under a reflector lamp
- 499 in the laboratory. A linear regression line is also shown. The data of 248 temperature
- 500 measurements from 6 flies were plotted

501

502	Fig. 4 Body-surface temperatures of Sarcophaga similis adults in the field and air temperatures
503	at which the body-surface temperatures were measured. A non-linear regression line (solid line)
504	is also shown. Circles and diamonds are the data from flies captured in 2013 and 2014,
505	respectively, and open and closed marks are the data of flies, the species of which were
506	identified and unidentified, respectively. Data from 151 flies were plotted
507	

508	Fig. 5 Relative amounts of <i>Hsp/0</i> , <i>Hsp23</i> , and <i>Hsp83</i> mRNAs of <i>Sarcophaga similis</i> adult flies
509	that were exposed to various temperatures in the laboratory (left panels) and were captured in
510	the field (right panels). The body-surface temperatures of the adults were also recorded. The
511	highest mean levels of Hsp70, Hsp23, and Hsp83 mRNAs at 40, 42, and 42°C in the laboratory
512	heat shock experiment were set at 1.0. $n = 3-4$ for each temperature in the laboratory heat shock
513	experiment. $n = 27-28$ for the expression of <i>Hsps</i> in adults in the field

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Fig. 6 Sun-shade preference of *Sarcophaga similis* adults. An adult fly was released into the experimental cage, half of which was covered with aluminium foil to form a shaded part. The cage was placed on sunlit ground. The air temperature, body-surface temperature, and the position of the fly (sunlit or shaded part) were recorded continuously. (A) air temperature and the total number of flies that were in a sunlit or shaded part. (B) body-surface temperatures of the flies in a sunlit or shaded part. Median, interquartile, maximum, and minimum values are shown. The data from 275 observations in 18 flies are shown

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Time-of-day (JST)

Fig. 2













