

Leeches *Baicalobdella* sp. feed on hemolymph but do not affect the cellular immune response of amphipod *Eulimnogammarus verrucosus* (Amphipoda, Crustacea) from Lake Baikal

Anna Nazarova¹, Andrei Mutin¹, Denis Skafar^{2,3}, Nadezhda Bolbat¹, Sofya Sedova¹, Polina Chupalova¹, Polina Drozdova^{1,4}, Anton Gurkov^{1,4}, Maxim Timofeyev^{Corresp. 1}

¹ Institute of Biology, Irkutsk State University, Irkutsk, Russia

² Faculty of Biology, Department of Aquatic Bioresources and aquaculture, Kuban State University, Krasnodar, Russia

³ Krasnodar department, Azov estuaries sector, Azov-Black Sea Branch of the Russian Federal Research Institute of Fisheries and Oceanography, Rostov-on-Don, Russia

⁴ Baikal Research Centre, Irkutsk, Russia

Corresponding Author: Maxim Timofeyev
Email address: m.a.timofeyev@gmail.com

Lake Baikal is one of the largest and oldest freshwater reservoirs on the planet with a huge endemic diversity of amphipods (Amphipoda, Crustacea). These crustaceans have various symbiotic relationships, including the rarely described phenomenon of leech parasitism on amphipods. It is known that leeches feeding on the hemolymph of crustacean hosts can influence their physiological status, especially under stressful conditions. Here we show that leeches *Baicalobdella* sp. found on the gills of the amphipod *Eulimnogammarus verrucosus*, one of the most abundant amphipods in the Baikal littoral zone, indeed feed on the hemolymph of their host. However, the leech infection had no effect on such immune parameters as hemocyte concentration and phenoloxidase activity, as well as glycogen content. The intensity of hemocyte reaction to foreign bodies in a primary culture was identical between leech-free and leech-infected animals. Artificial infection with leeches also had almost no modulating effect on bacterial influence on the hemocyte concentration and composition in hemolymph of amphipods after the injection modeling the microbial outburst. Thus, our study shows that the influence of a few leeches on *E. verrucosus* is probably negligible, and leech-infected amphipods can be used at least for some types of ecophysiological experiments.

1 **Leeches *Baicalobdella* sp. feed on hemolymph but do not**
2 **affect the cellular immune response of amphipod**
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4 **Lake Baikal**

5

6 Anna Nazarova ¹, Andrei Mutin ¹, Denis Skafar ^{2,3}, Nadezhda Bolbat ¹, Sofya Sedova ¹, Polina
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8

9 ¹ Institute of Biology, Irkutsk State University, Irkutsk, Russia

10 ² Faculty of Biology, Department of Aquatic Bioresources and aquaculture, Kuban State
11 University, Krasnodar, Russia

12 ³ Krasnodar department, Azov estuaries sector, Azov-Black Sea Branch of the Russian Federal
13 Research Institute of Fisheries and Oceanography, Rostov-on-Don, Russia

14 ⁴ Baikal Research Centre, Irkutsk, Russia

15

16 Corresponding Author:

17 Maxim Timofeyev ¹

18 Lenin street, 3, Irkutsk, 336003, Russia

19 Email address: m.a.timofeyev@gmail.com

20

21 **Abstract**

22 Lake Baikal is one of the largest and oldest freshwater reservoirs on the planet with a huge
23 endemic diversity of amphipods (Amphipoda, Crustacea). These crustaceans have various
24 symbiotic relationships, including the rarely described phenomenon of leech parasitism on
25 amphipods. It is known that leeches feeding on the hemolymph of crustacean hosts can influence
26 their physiological status, especially under stressful conditions. Here we show that leeches
27 *Baicalobdella* sp. found on the gills of the amphipod *Eulimnogammarus verrucosus*, one of the
28 most abundant amphipods in the Baikal littoral zone, indeed feed on the hemolymph of their
29 host. However, the leech infection had no effect on such immune parameters as hemocyte
30 concentration and phenoloxidase activity, as well as glycogen content. The intensity of hemocyte
31 reaction to foreign bodies in a primary culture was identical between leech-free and leech-
32 infected animals. Artificial infection with leeches also had almost no modulating effect on
33 bacterial influence on the hemocyte concentration and composition in hemolymph of amphipods
34 after the injection modeling the microbial outburst. Thus, our study shows that the influence of a
35 few leeches on *E. verrucosus* is probably negligible, and leech-infected amphipods can be used
36 at least for some types of ecophysiological experiments.

37

38 **Introduction**

39 Various parasites are now considered as a significant environmental factor influencing survival
40 of aquatic animals under stressful conditions and sometimes acting synergistically with such
41 factors as pollution [1,2]. In particular, some parasites have been shown to manipulate behavior,
42 distort sex ratio, modify energy budget and compromise the immune defense in amphipods
43 (Amphipoda, Crustacea), one of the most important groups of freshwater invertebrates [3].
44 Leeches are annelid worms (Hirudinea, Annelida), many species of which parasitize various
45 animals and feed on the host blood or hemolymph. Importantly, saliva components of these
46 parasites can have anticoagulant, anti-inflammatory and other roles, but such bioactive
47 components and their effects are mostly studied in medically important species [4,5,6]. Leeches
48 and crustaceans can exist in different types of ecological relationships. For example, leeches of
49 the species *Myzobdella lugubris* Leidy, 1851 are parasites of crabs *Callinectes bocourti* Milne-
50 Edwards, 1879 feeding on their hemolymph and laying eggs on the surface of the crab body [7].
51 The South African leech *Marsupiobdella africana* is a facultative ectoparasite of the amphibian
52 *Xenopus laevis* and has a phoretic relationship (i.e. promoting spreading of the attached phoront)
53 with the freshwater crab *Potamonautes perlatus* Milne-Edwards, 1837. The sex of the host crab
54 has been shown to be important in leech infestation. In addition, the period of residence of the
55 leeches on crabs corresponds to the development of leech eggs, which may indicate additional
56 benefits of these relationships for leeches [8]. The crayfish *Orconectes rusticus* (Girard, 1852)
57 has the cleaning leech-like symbiont *Cambarincola fallax* Hoffman, 1963 that removes fouling
58 organisms and thus improves growth rates of the host [9,10,11]. The fish leech *Johanssonia*
59 *arctica* (Johansson, 1898) is also an epibiont of the red king crab *Paralithodes camtschaticus*
60 (Tilesius, 1815) [12,13].

61 Lake Baikal is among the largest and most ancient freshwater reservoirs on the planet and also
62 the birthplace of outstanding endemic diversity of amphipods playing various roles in the lake
63 ecosystem [14]. Over 350 morphological species and subspecies of amphipods have been
64 described from Baikal, constituting about 19% of all known freshwater species and
65 demonstrating tremendous morphological variety [15,16]. Yet, symbionts and parasites of Baikal
66 amphipods and their potential influence on physiology of these crustaceans are understudied. It is
67 known that the hemolymph of the amphipods can contain various bacteria [17] and DNA of
68 microsporidians [18]. Baikal endemic amphipods are also known to be intermediate hosts for
69 acanthocephalans, but the fraction of infected individuals is generally low [19].

70 However, the parasites that can be most easily found on amphipods in Lake Baikal are leeches.
71 According to our observations, leeches are mostly attached to the gills of the largest
72 morphological species in the Baikal littoral zone such as *Eulimnogammarus verrucosus*
73 (Gerstfeldt, 1858) or *Pallasea cancellus* (Pallas, 1772) and much less often to a smaller *E.*
74 *vittatus* (Dybowsky, 1874). The hypothesis that the parasites prefer larger species as hosts is also
75 supported by observations of leeches on even larger deep-water Baikal amphipods [20]. Again,
76 according to our previous observations in *E. verrucosus*, leeches can infect a substantial
77 proportion of the population on the order of dozens of percents at least in some seasons. These
78 parasites of *E. verrucosus* belong to the genus *Baicalobdella* containing at least two species,

79 *B. cottidarum* Dogiel, 1957 and *B. torquata* (Grube, 1871) [21,22,23]. *E. verrucosus* is a
80 widespread and abundant morphological species in the littoral zone of Lake Baikal [24] and yet
81 influence of leeches on its physiology is fairly unstudied. Moreover, the whole phenomenon of
82 leeches infecting amphipods seems to be very rare if not unique to Lake Baikal, which might be
83 related to the larger size of many Baikal endemics in comparison to most freshwater amphipods.
84 Literature search gave us no other examples of such a phenomenon, and a recent review
85 categorizing parasites of amphipods do not mention leeches at all [25].
86 If leeches indeed feed on hemolymph of amphipods in Lake Baikal, the infection may directly
87 impair their immune defense and indirectly lower the available energy resources besides the
88 potential effects of leech saliva. The crustacean immune system relies on hemolymph
89 components such as hemocytes (i.e. circulating cells) and the phenoloxidase system. Hemocytes
90 perform phagocytosis and encapsulation of foreign bodies, while phenoloxidase is responsible
91 for the melanization process, which is also a part of foreign body encapsulation and hemolymph
92 clotting after injury [26]. So, in this study we mostly concentrated on testing the effects of leech
93 infection on these immune factors in *E. verrucosus* from Lake Baikal.

94

95 **Materials & Methods**

96 **Animal sampling and handling**

97 All experimental procedures were conducted in accordance with the EU Directive 2010/63/EU
98 for animal experiments and the Declaration of Helsinki; the protocol of the study was registered
99 and approved before the start of the experiments by the Animal Subjects Research Committee of
100 the Institute of Biology at Irkutsk State University (protocol #2022/11). Leech-free and leech-
101 infected amphipods *Eulimnogammarus verrucosus* (Gerstfeldt, 1858) were collected by kick
102 sampling with a hand net in Baikal littoral zone near the Listvyanka village (51°52'05.5"N
103 104°49'47.1"E) at depths 0-1.2m (the animals belong to the W barcoding species [27]).
104 Amphipods were acclimated to the laboratory conditions in well aerated 3 L plastic aquaria at
105 6°C in MIR-254 incubators (Sanyo, Osaka, Japan) for at least 3 days prior to any experiments.
106 Typically, leeches were attached to the gills of amphipods (Figure 1A,B).

107

108 **Identification of leech species**

109 After samplings in October 2022, February and April 2023 all clearly visible leeches were
110 detached from amphipods in and fixed in 96% ethanol for further species identification.
111 Morphological analysis of fixed specimens was performed on a stereo microscope SPM0880
112 (Altami, Russia) according to the standard keys (Bauer, 1987; Lukin, 1976). DNA extraction
113 from the posterior sucker of leeches was performed using the S-sorb kit (Syntol, EX-516,
114 Russia). PCR amplification of the cytochrome oxidase subunit I (COI) gene fragment was
115 performed with a 5× Screen Mix (Evrogen, Russia), the Folmer primers (LCO1490/HCO2198
116 [28]) and the following program: 94°C for 1 min, 30 cycles of 94°C for 20 s, 43°C for 2 min, and
117 72°C for 1 min.

118 The sequencing reactions were performed in both directions using BigDye Terminator v3.1
119 Cycle Sequencing kit (Life Technologies, USA) and analyzed with a Nanophor-05 Sanger
120 sequencer (Syntol, Russia). Sequencing reads were basecalled with the programs Mutation
121 Surveyor v5.1 and Chromas v2.6.6. Consensus sequences were compiled with UGENE v41.0
122 [29] using the sequence from *Baicalobdella* sp. (NCBI Genbank accession MN854834) as the
123 reference COI fragment. Sequences were aligned with the MAFFT algorithm [30] and trimmed
124 to 559 bp in the UGENE program. The sequences were deposited to the NCBI GenBank
125 database with accession numbers OR077511-OR077525. Similar sequences were searched with
126 BLAST (<https://blast.ncbi.nlm.nih.gov/Blast.cgi>) in the nt database. Phylogenetic tree was built
127 with the IQ-Tree web server (<http://www.iqtree.org/>) using automatic model selection with
128 Model Finder [31] and ultrafast bootstrap for assessment of the branch support values [32]. The
129 resulting phylogeny was visualized with iTOL (<https://itol.embl.de/>) [33].

130

131 **Injection of fluorescent latex beads into amphipods and further visualization**

132 After sampling in June 2023, we analyzed the ability of leeches *Baicalobdella* sp. to consume
133 amphipod hemolymph. For this, leeches were detached from amphipod gills with tweezers and
134 kept in aquaria separately from hosts for ~24 h. Next, 10 non-infected individuals of *E.*
135 *verrucosus* were injected with 1 μ L of suspension containing about 3×10^6 latex microbeads
136 (L3030, Sigma-Aldrich) using a IM-9B microinjector (Narishige, Tokyo, Japan). Right after the
137 injection, the amphipods were placed in aquaria with free leeches, which attached to the new
138 hosts during 30 min.

139 Four hours post injection we anesthetized the amphipods in clove oil suspension (50 μ L of clove
140 oil per 50 mL of Baikal water) and detached leeches and two pieces of gills from each individual
141 for further observation under an inverted fluorescent microscope Celena S (Logos Biosystems,
142 Republic of Korea). Prior to the visualization, the leeches were placed into sterile 1.5-mL
143 microtubes and homogenized with 50 μ L of phosphate buffered saline using a plastic pestle.

144

145 **Hemolymph extraction of characterization of hemocytes**

146 Before the hemolymph extraction, the dorsal side of the amphipod pereon surface was always
147 sterilized with 70% ethanol. The central hemolymph vessel was punctured with a sterile needle,
148 and hemolymph was collected with a sterile glass capillary. The amphipod hemolymph was
149 mixed (1:1) with the isotonic anticoagulant solution (150 mM NaCl, 5 mM Na₂HPO₄, 30 mM
150 sodium citrate, 10 mM EDTA, pH 8.0; filtered through a 0.45 μ m syringe filter) on ice to avoid
151 degranulation of granulocytes [17]. Amphipod hemolymph was always extracted before
152 detachment of leeches. Hemocytes were visualized using the Celena S inverted microscope
153 (Logos Biosystems, Republic of Korea) or the Mikmed-2 microscope (LOMO, Russia). Total
154 hemocyte count (THC) and granulocyte percentage was estimated in disposable hemocytometers
155 (Aptaca, Italy).

156 Characterization of hemocyte types was performed with a CytoFLEX flow cytometer (Beckman
157 Coulter, USA, CA). Hemolymph of 8 non-infected amphipods *E. verrucosus* was extracted and

158 measured for forward (allows for the discrimination of cells by size) and side scatter (gives the
159 information about cell complexity).

160

161 **Biochemical measurements of phenoloxidase activity and glycogen content**

162 Along with estimation of THC, part of infected and non-infected animals collected in October
163 2022, February or April 2023 were used for measurements of phenoloxidase activity and
164 glycogen content. For phenoloxidase measurements, hemolymph was collected between the 7th
165 and 8th segments of mesosome as described above, mixed 1:1 with a buffer solution (150 mM
166 NaCl, 10 mM Na₂HPO₄, 7 mg/mL phenylmethanesulfonyl fluoride, pH 8.0) and frozen at -80
167 °C. The samples were melted at 4°C and centrifuged for 10 min at 500 g and 4°C to precipitate
168 the cellular pellets. 10 µl of hemolymph extract were mixed with 40 µl of buffer solution, 280
169 µL of distilled water, and 40 µl of 4 mg/ml 3,4-dihydroxy-L-phenylalanine. Measurements were
170 performed with the CLARIOstar Plus microplate reader (BMG Labtech, Germany) at 490 nm
171 (absorbance) for 40 min. Activity of phenoloxidase was assessed in arbitrary units as the slope of
172 the reaction curve during the linear phase [17].

173 Glycogen extraction was performed as described previously [34] with modifications. Frozen
174 amphipod tissues (after hemolymph extraction) were ground into a powder, mixed with the
175 solution (0.5 mL per 100 mg of wet weight) containing 0.6 M HClO₄, and further homogenized
176 in a Potter-Elvehjem tissue grinder until no visible particles remained. Next, 20 µL of the
177 homogenate was mixed with 75 µl of 1% amyloglucosidase (10115-5G-F, Sigma-Aldrich,
178 Germany; 5250 U/µL) in a 0.2 M acetic acid buffer (acetic acid/sodium acetate; pH 4.8). The
179 mix was incubated at 40°C for two hours and then 62.5 µl of 0.6 M HClO₄ and 100 µl of 1 M
180 KHCO₃ were added. The supernatant was centrifuged at 13000 rpm for 15 minutes. Glycogen
181 concentration was measured with the kit "Glucose-Vital" (Vital Development, Russia), 40 µl of
182 experimental sample was added to 190 µl of "Glucose-Vital" monoreagent and incubated at
183 25 °C for 15 minutes. Light absorption was measured at 510 nm with a CLARIOstar plus
184 microplate reader (BMG Labtech, Germany).

185

186 **Assessing encapsulation of Sephadex beads by amphipod hemocytes in primary culture**

187 Sephadex microbeads (G100120-50G, Sigma-Aldrich, United States) were washed with 5 mg/ml
188 streptomycin and 5000 U/mL penicillin solution (1.3.18, Biolot, Russia). The bead suspension
189 was pipetted into a sterile 96-well plate (GT204-0096DV, Minimed, Russia) in a laminar flow
190 box. Then, 100 µL of complete medium 1X L-15 (Leibovitz medium with L-glutamine, L4386-
191 10X1L, Sigma-Aldrich, United States) containing 15% fetal bovine serum (FBS-HI-11A,
192 Capricorn Scientific, Germany) was added, and the beads were imaged under Mikmed-2
193 microscope (LOMO, Russia) with attached EOS 1200D camera (Canon, Taiwan) and counted
194 using the Count Things application (CountThings.com). Since the amount of microbeads per
195 well varied substantially, for further tests we used only the wells with approximately the same
196 number of beads (on average 230±70 beads per well).

197 Hemolymph was extracted from 10 leech-free and 12 leech-infected amphipods (collected in
198 April 2023) as described above and was pooled within each group. 10- μ L aliquotes were
199 collected from the pools to estimate the hemocyte concentrations. Then, each pool of
200 hemolymph was divided to the selected wells with microbeads controlling for the equal amounts
201 of hemocytes for leech-free (15 wells) and leech-infected (11 wells) groups (on average,
202 $1 \pm 0.4 \times 10^5$ cells per well).

203 After cell sedimentation to the well bottom, the upper layer of the suspension was collected, and
204 100 μ L of fresh medium was added. The hemocyte response to the Sephadex microbeads was
205 analyzed after 24 hours of incubation, and the number of microbeads with hemocyte aggregates
206 was counted under the Celena S inverted fluorescent microscope (Logos Biosystems, Republic
207 of Korea). We categorized 4 stages of the encapsulation reaction: no reaction, low reaction,
208 medium reaction, the stage showing partially encapsulated beads, and the intense reaction
209 showing fully covered beads (Figure 1C,D). The hemocyte nuclei were stained with 10 μ g/mL
210 4',6-diamidino-2-phenylindole (DAPI, A4099, AppliChem, Germany) to visually contrast the
211 encapsulation reaction (Figure 1D). Cell viability was assessed by staining with 1 μ g/mL
212 propidium iodide (81845-100MG, Sigma-Adrich, Germany).

213

214 **Artificial infection of amphipods with leeches and bacteria**

215 In this research, we used the bacterial strain *Pseudomonas* sp. H5-2 (belongs to the *P.*
216 *fluorescens* species group) that was previously extracted from the hemolymph of *E. verrucosus*
217 collected in the same location [17]. For the cultivation, we used the tryptic soy broth (TSB)
218 medium (casein peptone, dipotassium hydrogen phosphate, glucose, NaCl, soy peptone) as
219 suggested previously [35,36]. For injection into amphipods, *Pseudomonas* sp. cells were washed
220 by centrifugation and resuspended in physiological solution (150 mM NaCl, 10 mM Na_2HPO_4).
221 2.5 μ L of physiological solution with 10^5 *Pseudomonas* sp. cells per 1 μ L was injected into the
222 central hemolymph vessel of amphipods between 5th and 6th segments with an IM-9B
223 microinjector (Narishige, Tokyo, Japan).

224 After ~15-30 min, the amphipods with and without the bacterial injection were infected by
225 leeches as described above with 1:1 parasite to host ratio. After 1.5 hours, 1 and 3 days
226 hemolymph was extracted from amphipods and mixed 1:1 with anticoagulant solution (150 mM
227 NaCl, 5 mM Na_2HPO_4 , 30 mM sodium citrate, 10 mM EDTA, 50 mM EDTA- Na_2 , pH 8.0). This
228 adjusted anticoagulant solution allows to fix hemocytes in the state when nuclei and granules are
229 visible more clearly [37] and was applied to later visually distinguish granulocytes among all
230 hemocytes. THC and granulocyte proportion were estimated under the Mikmed-2 microscope
231 (LOMO, Russia) in a glass hemocytometer.

232 All experimental groups for 1.5 h and 1 day timepoints included 10 animals per group and
233 showed no mortality both with and without bacterial infection (since hemolymph samplings
234 always failed for part of animals, the number of analyzed hemolymph samples could be reduced
235 down to 7 for some groups). The first round for 3-day timepoint also included 10 animals per
236 each experimental group but showed high mortality specifically for animals injected with

237 bacteria (60% for leech-free and 50% for leech-infected) with no mortality for amphipods
238 without injection. Since this high mortality could be an artifact of specific injection procedure,
239 we performed the second round of the experiment with bacterial injection into 9 animals per
240 experimental group. Both leech-free and leech-infected animals showed no mortality during 3
241 days post injection, and their hemolymph was used for the tests along with hemolymph of the
242 animals from the first round.

243

244 **Statistical analysis**

245 Statistically significant differences between experimental groups were always estimated using
246 Mann-Whitney U test with the Holm's correction for multiple comparisons. Mann-Whitney tests
247 were performed in the program Past 4.03 [38], and Holm's corrections were applied in R v.4.3.1
248 [39]. The differences were considered statistically significant with $p < 0.05$.

249 Specifically for the experiment with artificial infections with bacteria and leeches we applied a
250 generalized linear model (GLM) for analysis of factor effects in R v.4.3.1 [39]. The model was
251 fitted using the `glm()` function with gaussian distribution to three independent factors (time as
252 numeric variable, absence or presence of leech and injected bacteria) and all of their interactions.
253 The assumptions for GLM were mostly met for the dataset: the outcome with time was
254 acceptably linear (slightly violated specifically for THC), the residuals were always
255 homoscedastic and the normality assumption was slightly violated only for THC.

256

257 **Results**

258 **Infection rate and identification of leeches**

259 We collected amphipods *E. verrucosus* infected with leeches at the same location in Lake Baikal
260 but in different seasons. The infection rate was not estimated precisely, but it clearly varied
261 greatly. In October 2022, ~80 % of individuals were infected, while in February and April 2023
262 the rate was substantially lower on the order of ~5 %. There were from one to nine leeches per
263 animal with 2-4 parasites found on most individuals.

264 Morphological analyses showed us that all 42 leeches sampled in these seasons belong to one
265 genus *Baicalobdella* with prevalence of morphospecies *B. torquata*. Several leeches were
266 identified as potentially belonging to morphospecies *B. cottidarum* but recent data [40] indicate
267 that *B. torquata* may have significant morphological variability, so identification of these
268 specimens remained uncertain. In order to clarify the diversity of the leeches, we performed
269 sequencing of the COI gene fragment in 15 specimens in total; all samples with ambiguous
270 morphological identification were included in the analysis, and the rest identified as *B. torquata*
271 were chosen randomly from all three samplings.

272 The phylogenetic tree clearly showed that all 15 leeches belong to the same species (Figure 2).
273 They were found to be approximately 90% similar to the available sequence of *Baicalobdella* sp.
274 (#MN854837.1). However, the sequence of *B. torquata* (#MN854834) belongs to a different
275 group. Previously, it was found that morphological species *B. torquata* may in fact be a complex
276 of cryptic species [41], and the leeches found in our study could belong to one of them but we

277 have no means to identify the species more precisely. However, since *E. verrucosus* was found to
278 be infected with only one species of *Baicalobdella* locally, we had the possibility to test the
279 physiological influence of these leeches onto the amphipods.

280

281 **Leeches *Baicalobdella* sp. consume amphipod hemolymph**

282 The assumption that the leeches attached to gills of amphipods also feed on their hemolymph is
283 obvious, but these ectosymbionts may simply be in phoretic relationships with specifically these
284 hosts. In order to test this assumption, we injected fluorescent microbeads into the hemolymph of
285 *E. verrucosus* and tracked their distribution. Five hours post injection the microbeads were easily
286 observable in amphipod gills and also inside some ciliates that were found to be attached to gills
287 (Figure 3A,B). The homogenate of 5 out of 10 tested leech bodies also contained these
288 fluorescent microbeads (Figure 3C). Since the leech oral apparatus is not suitable for
289 consumption of ciliates [22], our data unambiguously confirms that the leeches *Baicalobdella* sp.
290 indeed can feed on hemolymph of amphipods *E. verrucosus*.

291

292 **Characterization of amphipod hemocytes**

293 Since hemocytes are an important component of the crustacean immune system, before further
294 analysis we investigated their possible subdivision into populations. Flow cytometry clearly
295 differentiated hemocytes of *E. verrucosus* into two main groups, one with smaller cell size and
296 lower internal complexity and the other with larger cell size and higher internal complexity
297 (Figure 4). The groups are usually called hyalinocytes and granulocytes respectively [42] and can
298 also be observed with common phase contrast microscopy. In particular, larger size of
299 granulocytes is evident right after the sample is placed under the microscope, while higher
300 amount of vesicular structures in granulocytes is better visualized after attachment to the surface
301 (Figure 4). Additionally, we observed hemocytes with intermediate internal complexity and size
302 between granulocytes and hyalinocytes, i.e. semi-granulocytes, but their proportion was only
303 ~10%.

304

305 **Influence of leeches on hemocyte concentration and other parameters of amphipods in dif-** 306 **ferent seasons**

307 We used the amphipods collected in October 2022, February and April 2023 for discriminating
308 the effects of leech infection on several physiological parameters of *E. verrucosus* in different
309 seasons. The consumption of hemolymph by leeches may directly reduce the hemocyte
310 concentration and phenoloxidase content in the hemolymph and indirectly reduce the available
311 glycogen due to the compensation of the tissue loss.

312 Total hemocyte count (THC) varied with season, but leech-infected and non-infected amphipods
313 never had a statistically significant difference in this parameter (Figure 5A). Moreover, median
314 THC in February 2023 was even slightly higher in infected than in non-infected animals despite
315 the fact the size of leeches was the largest (Figure 5B,C) and hemolymph consumption would be
316 expected to be the highest. It could be hypothesized that hemolymph consumption, on the

317 contrary, lowers specifically for the largest leeches that are ready to switch their host.
318 Interestingly, there was indeed a statistically significant difference between infected amphipods
319 in October and February (Figure 5A), which partially supports this hypothesis. Since most
320 amphipods were infected with 2-4 leeches, we could not check the correlation between THC and
321 leech size directly, but the dependence between THC and summarized leech width per host was
322 absent or even indicated slightly higher THC in amphipods with higher biomass of leeches
323 (Figure 5D). In general, these data show that influence of leeches on THC is either negligible or
324 the hemocyte loss is compensated by the host.
325 Similarly, phenoloxidase activity of *E. verrucosus* hemolymph (analyzed only in April 2023)
326 was not significantly different between infected and non-infected amphipods (Figure 5E).
327 Finally, glycogen content was also almost identical between the groups, which indicates no
328 prominent energetic burden of the infection.
329 Overall, our data suggest that infection with leeches does not have substantial deleterious effects
330 for amphipods *E. verrucosus* at least in the analyzed seasons.

331

332 **Cellular immune response of infected and non-infected amphipods estimated *in vitro***

333 Despite the fact that leeches did not substantially affect the amounts of immune components in
334 amphipod hemolymph, they might modulate intensity of the immune response through bioactive
335 components in their saliva. For preliminary testing of this hypothesis, we chose the primary
336 culture of amphipod hemocytes as a convenient model system and Sephadex microbeads
337 (consisting of specifically processed dextran) as model foreign bodies. The primary hemocyte
338 culture allows for observing the behavior of these immune cells and quantitative estimation of
339 their reactions such as aggregation and further encapsulation of foreign bodies.

340 In particular, we measured the fraction of Sephadex beads encapsulated by hemocytes originally
341 extracted from leech-infected and non-infected amphipods 24 h after contact with the beads. This
342 time point was previously shown to be enough for development of strong immune reaction even
343 to artificial non-microbial foreign bodies [43]. We found no difference in the intensity of the
344 immune reaction between the experimental groups since the proportions of fully encapsulated
345 (~10%) and partially encapsulated microbeads (~85%) were equal for hemocytes from infected
346 and non-infected amphipods (Figure 6A). Some of the beads were not encapsulated at all, and
347 there was a high mortality of hemocytes around Sephadex microbeads in contrast to free
348 hemocytes, as indicated by propidium iodide staining (Figure 6B).

349 However, the humoral components of hemolymph were mostly removed during hemocyte
350 transfer into culture medium (dilution was ~12x), which could alleviate the potential influence of
351 leech infection during the experiment. Thus, here we could not fully exclude the possible minor
352 effects of *Baicalobdella* sp. saliva on the intensity of cellular immune response in amphipods of
353 Lake Baikal.

354

355 **Changes in hemocyte concentration and composition after injection of bacteria and** 356 **artificial leech infection**

357 Finally, in order to evaluate the potential synergistic effects of leeches and other immunity-
358 related factors we experimentally tested the influence of leeches on the ability of amphipods to
359 deal with bacterial infection. In this study we performed (i) an artificial infection of leech-free
360 *E. verrucosus* with the *Pseudomonas* sp. strain originally extracted from hemolymph of the same
361 species and (ii) an artificial infection with leeches. The amount of injected bacterial cells was
362 comparable to the number of circulating hemocytes in the animal hemolymph to model a
363 significant microbial outburst. Some of the amphipods were then infected with one leech per
364 animal, and such parameters as hemocyte concentration in the hemolymph and the fraction of
365 granulocytes were evaluated for three days (Figure 7; Table 1).

366 The mortality during the three-day experiment was mostly low, and it never was higher for
367 leech-infected animals than for leech-free ones. The generalized linear model indicated that
368 infection with leeches itself and time after the infections had no statistically significant effects on
369 both the concentration of hemocytes in amphipod hemolymph and the fraction of granulocytes
370 among them, while the injection of bacteria clearly leads to a statistically significant decrease in
371 hemocyte concentration by ~2800 cells per μl on average and an increase in granulocyte
372 proportion by ~16% (Table 1). Interestingly, the interaction between bacterial injection and
373 infection with leeches, oppositely, led to a statistically significant decrease in the fraction of
374 granulocytes by 12% but caused no statistically significant changes in hemocyte concentration
375 (Table 1). Other interactions between factors even being statistically significant in the case of
376 granulocytes percentage did not exceed 1% by module in the estimated effect. However, pair-
377 wise comparisons between leech-free and leech-infected animals gave no statistically significant
378 differences between any experimental groups not only in hemocyte concentration but also in
379 granulocytes fraction during the whole experiment (all adjusted $p > 0.05$; Figure 7). Thus, the
380 effect of leech infection partially compensating the effect of bacterial infection specifically for
381 hemocyte composition deserves further attention, but overall our results clearly demonstrated no
382 synergistic interaction between these two factors.

383

384 Discussion

385 Our research group focuses on environmental physiology of the amphipods endemic to Lake
386 Baikal, and almost exclusively the previously published experiments were made with amphipods
387 without visible leech infection [44,45,46,47] since the infected individuals were considered as
388 potentially weakened. Here we questioned this assumption.

389 Sequencing of leeches from *E. verrucosus* collected in three independent sampling campaigns
390 clearly demonstrated that the parasites in the used sampling location belong to the same species,
391 and thus their influence on these amphipods can be studied without preliminary species
392 identification. Since the phylogeny and diversity of leeches in Lake Baikal are still being revised
393 [48,20,49,50], we could not identify the species precisely but it belongs to the genus
394 Baicalobdella (Figure 2). Our tests also showed that the leeches can consume amphipod
395 hemolymph (Figure 3), and thus the effects of the infection on amphipod physiology are worth
396 studying. However, within the 4-hour experiment, only a half of the artificially attached leeches

397 consumed hemolymph, and according to our observations, in nature the leeches were mostly
398 attached to gills with their posterior sucker, which indicates that these parasites do not consume
399 hemolymph constantly.

400 The studies investigating the host-symbiont relationships of amphipods commonly use such
401 techniques as histological analysis, spectrophotometry, metagenomics, PCR and microscopy,
402 while such an important component of the immune system as hemocytes is rarely mentioned
403 [25]. Rigaut and Moret studied phenoloxidase activity of *Gammarus pulex* (Linnaeus, 1758) and
404 *G. roeselii* and found a correlation between infection by acanthocephalans and a decrease in the
405 enzyme activity [51]. Another freshwater amphipod *G. fossarum* (Koch, 1835) was used as a test
406 organism to investigate potential pollutant-parasite interactions for infection with larvae of the
407 acanthocephalan species *Polymorphus minutus* (Zeder, 1800). Phenoloxidase activity, glycogen,
408 and lipid concentrations showed a significant increase in infected gammarids [52]. In the other
409 case acanthocephalan infection was associated with reduction of the phenoloxidase activity and
410 the hemocyte concentration [53].

411 Here we were looking for any substantial effects of leech infection on hemocytes and other
412 related parameters of amphipods. Most studies on crustacean hemocytes have been performed
413 for decapods and revealed three main types of these immune cells with different morphology and
414 functions: hyalinocytes (hyaline cell), semi-granulocytes (semi-granular cells) and granulocytes
415 (granular cells) [42]. Such information for amphipods is less abundant. Using light and electron
416 microscopy the following hemocyte types were found in the body of the amphipod *G. setosus*
417 (Dementieva, 1931): granulocytes, adipohemocytes, plasmatocytes, and rare prohemocytes [54].
418 In the hemolymph of the amphipod *G. pulex* four types of circulating cells were identified with
419 microscopy and histochemical staining: hyalocytes I (cells with a transparent cytoplasm),
420 hyalocytes II (cells with a slightly basophilic cytoplasm), granulocytes, and adipohemocytes
421 (with large nucleus surrounded by granules) [55]. In the case of *Parhyale hawaiiensis* (Dana,
422 1853), it was shown that hemolymph contained tree typical type of hemocytes: granulocytes,
423 semi-granulocytes and hyalinocytes with semi-granulocytes being rare [56]. Our research on
424 *E. verrucosus* seems to be the first or among the first studies checking amphipod hemocyte
425 diversity with flow cytometry, which demonstrated the prevalence of two types of hemocytes,
426 granulocytes and hyalinocytes, while the intermediate semi-granulocytes were found to be
427 relatively rare (Figure 4).

428 We used leech-free and leech-infected amphipods from the same samplings and of similar size in
429 order to compare the concentration of hemocytes in hemolymph, phenoloxidase activity and
430 glycogen content and found no influence of leeches on these parameters (Figure 5). Hemocyte
431 concentration varied greatly with sampling campaigns and could be partially influenced by the
432 reproduction season, which starts in autumn for *E. verrucosus*. Interestingly, the infection rates
433 were also very different in different months (dropped from ~80% to ~5% from October to April),
434 which indirectly indicates that the same individual of this species can be infected with different
435 leeches multiple times during their lifespan of about five years. A previous transcriptomic study
436 indicated that even *E. verrucosus* without visible leech infection sometimes can bear the

437 parasites, so the mentioned values might be an underestimate [44]. However, the infection seems
438 to substantially affect neither immune defense, nor energy budget of the animal. This discovery
439 sheds some light into the host-symbiont relationships of *E. verrucosus* and *Baicalobdella* sp.
440 showing that the infection with leeches is probably more natural for large amphipods of Lake
441 Baikal than we would assume.

442 Next, we checked for potential influence of leech saliva on reaction intensity of hemocytes to
443 artificial foreign bodies in the primary cell culture. Hemocytes from leech-free and leech-
444 infected animals demonstrated the same results (Figure 6), but since humoral components of
445 hemolymph were diluted for ~12 times during extraction into the primary culture, we can only
446 exclude a very intense influence of the saliva components. Finally, we checked for potential
447 synergistic interaction, of leeches with artificial bacterial infection and found no or even a slight
448 antagonistic interaction, as indicated by the estimates of granulocyte fraction among all
449 hemocytes (Figure 7; Table 1). Artificial infection with leech did not influence hemocyte
450 concentration or granulocyte percentage in the amphipod hemolymph at all, while injection of
451 bacteria clearly decreased the first and increased the second (Figure 7; Table 1). The decrease in
452 THC was expected from a number of studies [57,58,59,60]. The increase in fraction of
453 granulocytes among all hemocytes probably reflects high mortality of hyalinocytes during the
454 immune response to bacteria but possible discharge of granulocytes from some tissues also
455 cannot be excluded. The antagonistic interaction of leech infection with bacterial injection
456 specifically in the case of the granulocyte fraction among all hemocytes might be speculatively
457 explained by a potential decrease in the concentration of bacteria due to hemolymph
458 consumption by the leech, but this effect clearly demands further exploration.

459 An unexpected finding of our research was the discovery of numerous parasitic ciliates on the
460 gills of *E. verrucosus* that clearly consumed amphipod hemolymph (Figure 3B). It is known that
461 ciliates of the genus *Lagenophryidae* can seat on gills of amphipods *E. verrucosus* [61].
462 However, their potential influence on the amphipods is a subject for a separate research.
463 Overall, our study indicated no substantial influence of leeches on the amphipods *E. verrucosus*
464 from Lake Baikal. Therefore, the individuals infected with *Baicalobdella* sp. can or sometimes
465 even should be included into ecophysiological experiments for performing them on the
466 representative part of the population.

467

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675

Figure 1

Representative photos of the research objects.

(a) Photo of amphipod *Eulimnogammarus verrucosus* with a leech attached to its gills indicated by the black arrow. (b) Microscopic photo of a leech with amphipod gill after detachment. (c,d) Hemocytes of *E. verrucosus* and stages of their encapsulation reaction to Sephadex beads after DAPI staining. c, bright field channel; d, DAPI channel of the fluorescent microscope. Pink arrows, no response; green arrows, low response; dark blue arrows, intermediate response; red arrows, intense response.

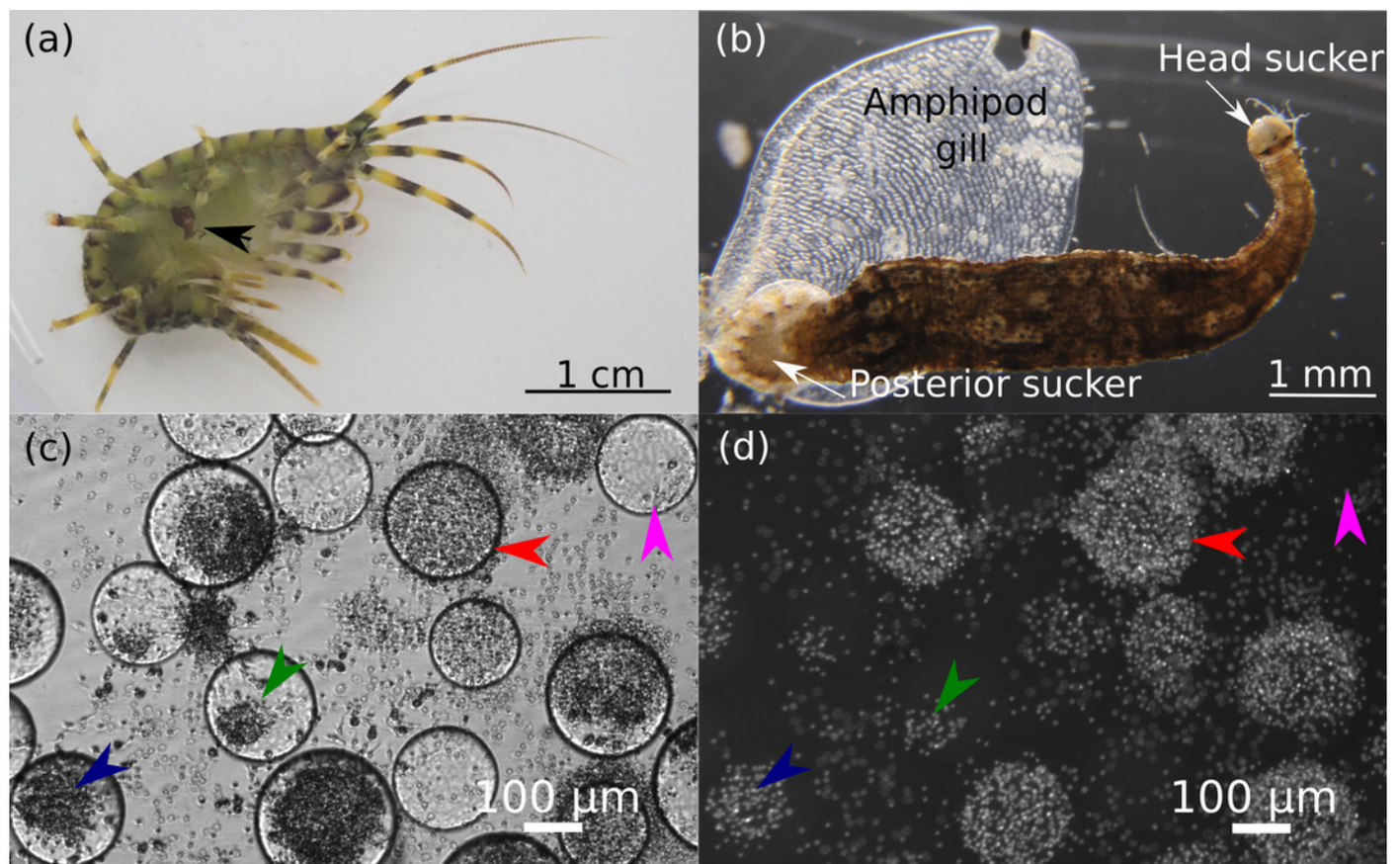


Figure 2

Phylogenetic tree of partial COI gene sequences of leech samples detached from amphipods *E. verrucosus* collected in Baikal littoral zone nearby Listvyanka village (highlighted in blue) and sequences of other closely related leeches.

The numbers next to the nodes mean percent of their ultrafast bootstrap support.

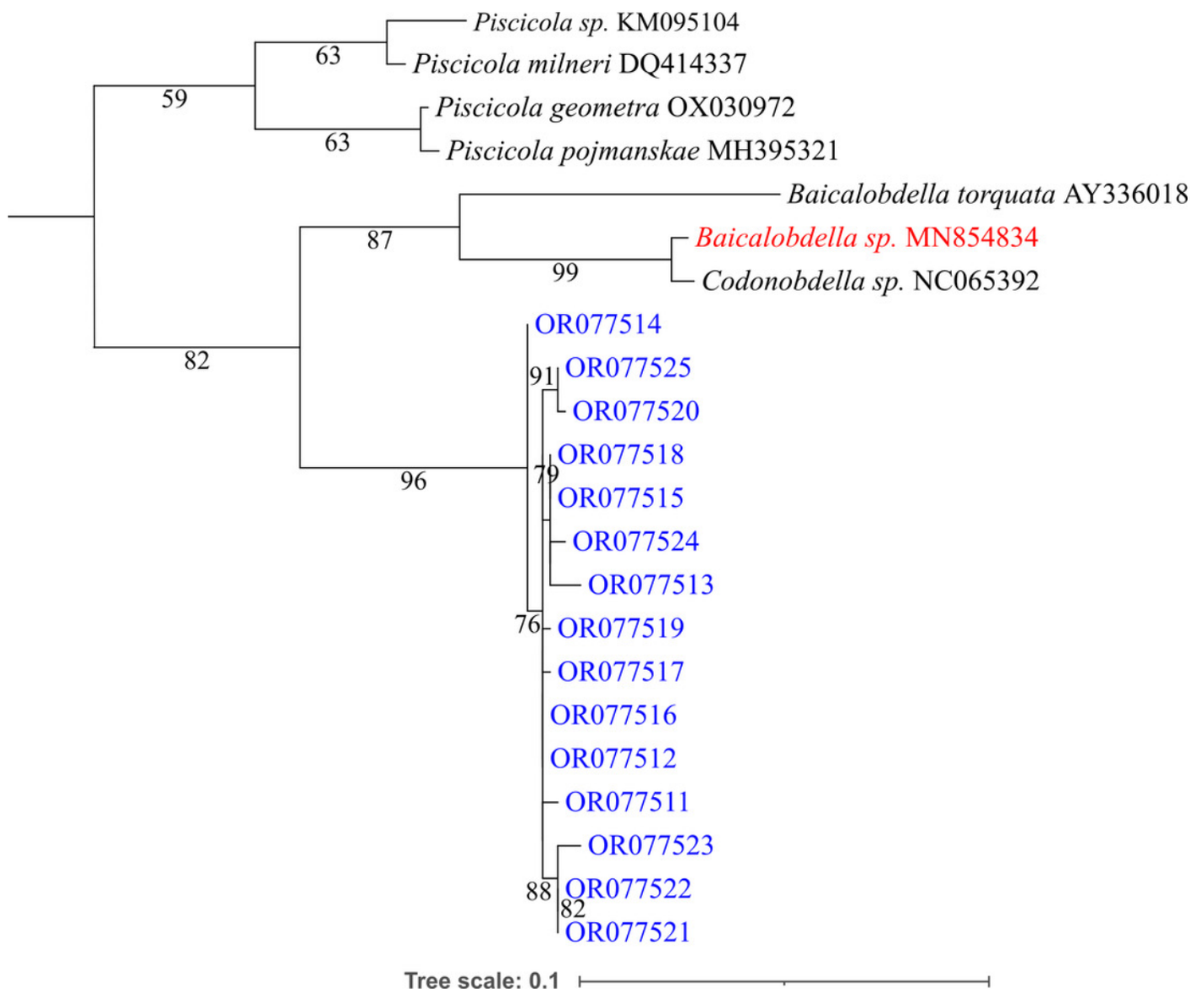


Figure 3

Distribution of latex microbeads 5 hours after injection into the central amphipod hemolymph vessel.

(a) Amphipod gill with latex microbeads and hemocytes. **(b)** Ciliate cells on the surface of gills with the microbeads inside them. **(c)** Content of leech body with latex microbeads. The pictures are merged photos obtained in brightfield and RFP channels with the same camera settings. Red arrows, latex microbeads; blue arrows, ciliates with microbeads inside. Scale bars: 100 μm (A, B), 50 μm (C).

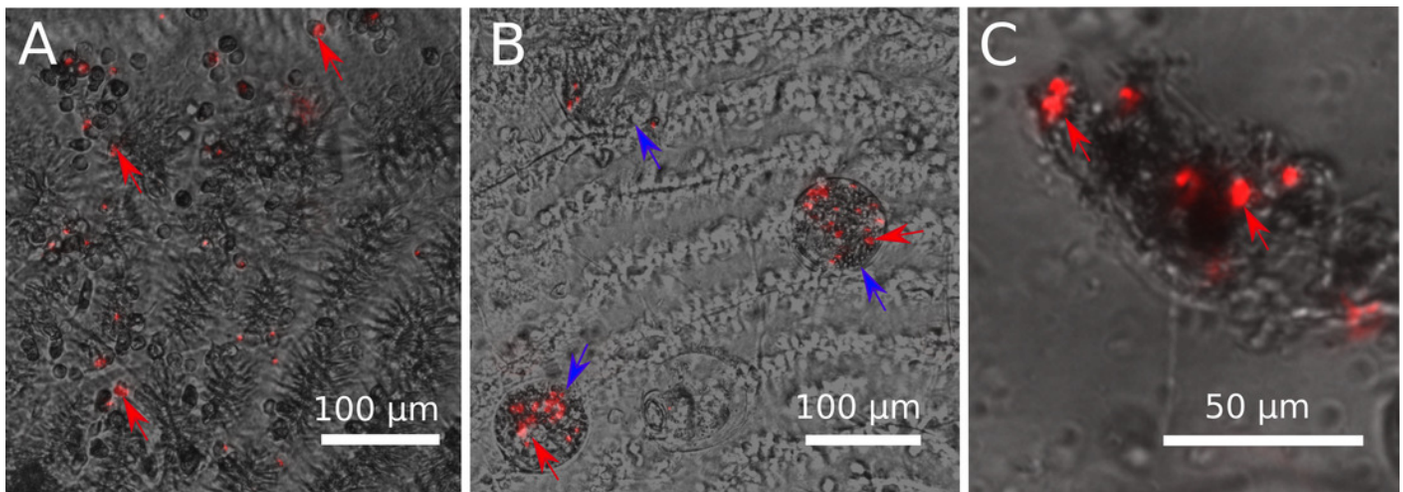


Figure 4

Characterization of *E. verrucosus* hemocytes using flow cytometry and microscopy.

Left panel shows the internal complexity (side scatter, SSC) against the cell size (forward scatter, FSC) of hemocyte populations (hyalinocytes and granulocytes), and other panels depict their respective phase contrast photos before (center) and after (right) attachment to a glass surface.

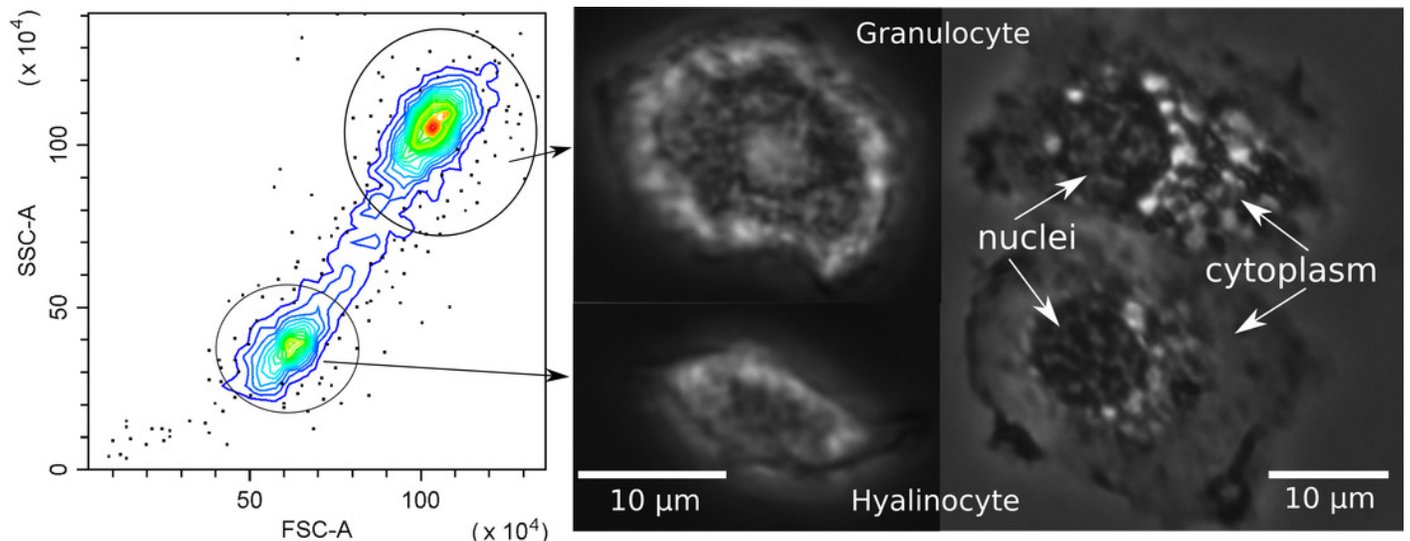


Figure 5

Different parameters of leech-infected non-infected amphipods *E. verrucosus* collected from natural environment and the leeches in different seasons.

(a) Total hemocyte count of infected and non-infected *E. verrucosus* collected in different seasons (n = 5-10). **(b)** Width of leeches in the middle of the body in different seasons. **(c)** Representative photos of leeches detached from *E. verrucosus* in different seasons. **(d)** Dependence of hemocyte count on summarized width of all leeches infecting the animal. The difference of the regression coefficient from zero is not statistically significant (p = 0.34). **(e)** Phenoloxidase activity in hemolymph of leech-free and leech-infected *E. verrucosus* collected in April 2023 (n = 10). Color legend is identical to panel (a). **(f)** Amount of glycogen in leech-free and leech-infected *E. verrucosus* collected in October 2022, February and April 2023 (n = 10-11). Color legend is identical to panel (a).

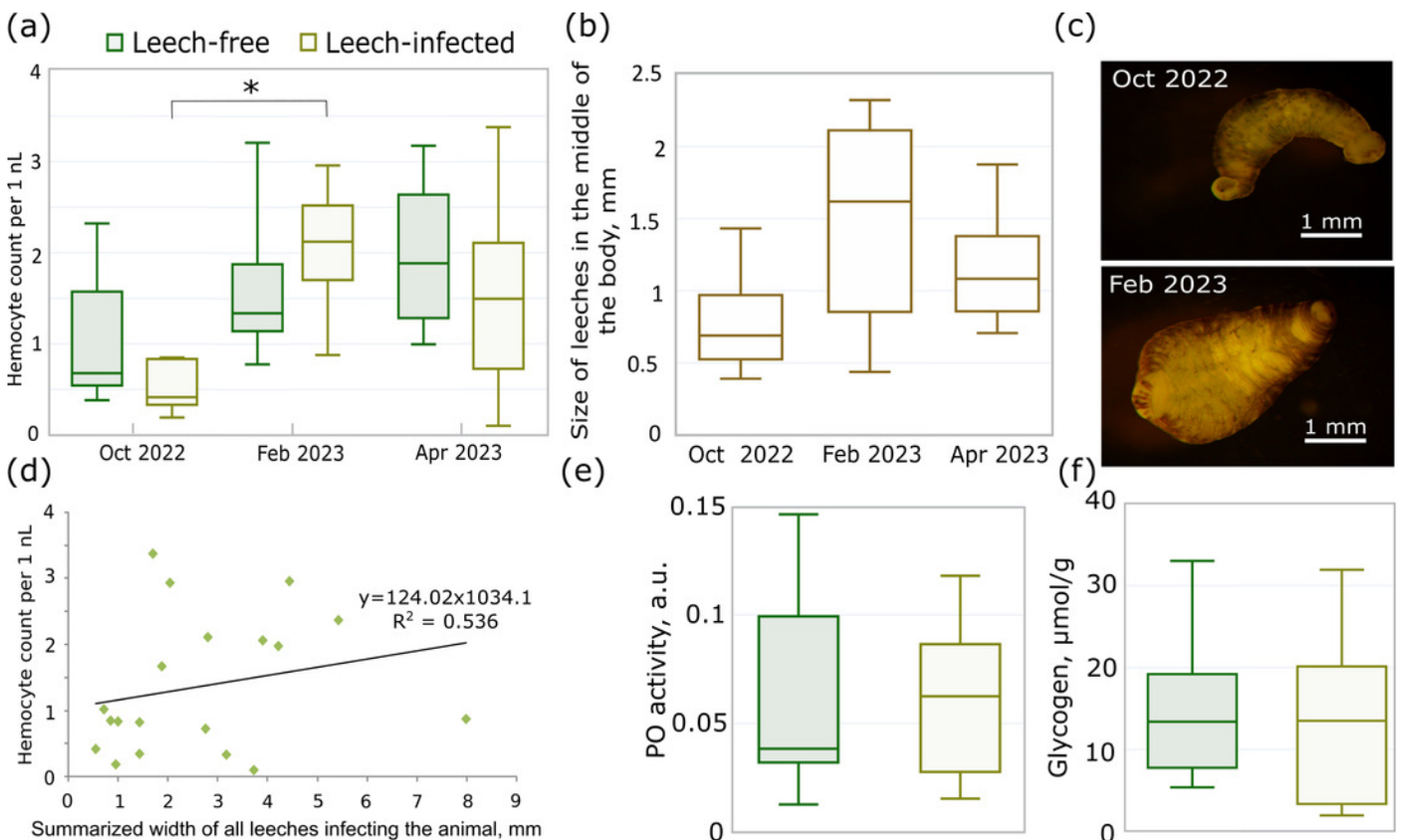


Figure 6

Intensity of the cellular immune response of hemocytes extracted in primary culture from leech-free and leech-infected amphipods *E. verrucosus*.

(a) Fractions of Sephadex microbeads partially and fully encapsulated by hemocytes after 24 hours of contact. (b) Example photos of microbeads' encapsulation in hemocyte primary culture, propidium iodide staining. Pink arrows — Sephadex microbeads, red arrows — aggregates of hemocytes. Photos in RFP channel were obtained at the same camera settings in different groups, but time of staining could be different.

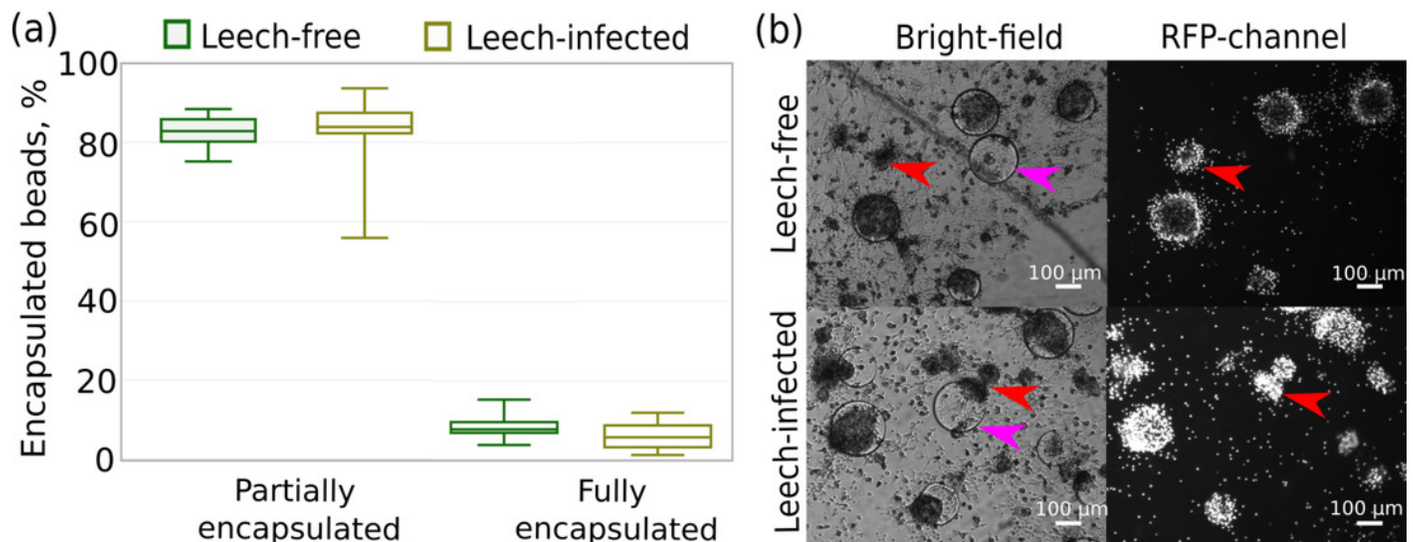


Figure 7

Immune cells in the hemolymph of leech-free and artificially leech-infected amphipods with and without bacterial injection.

(a) Total hemocyte count (n = 7-13). **(b)** Granulocyte fraction among all hemocytes (n = 7-13). The legend is identical for (a) and (b). Injection of bacteria to amphipod central hemolymph vessel was performed about 15 minutes before leech infection. Pair-wise comparisons of leech-free and leech-infected animals in each time point with Mann-Whitney U test with Holm's correction for multiple comparisons gave no statistically significant differences both for amphipods with and without bacterial infection.

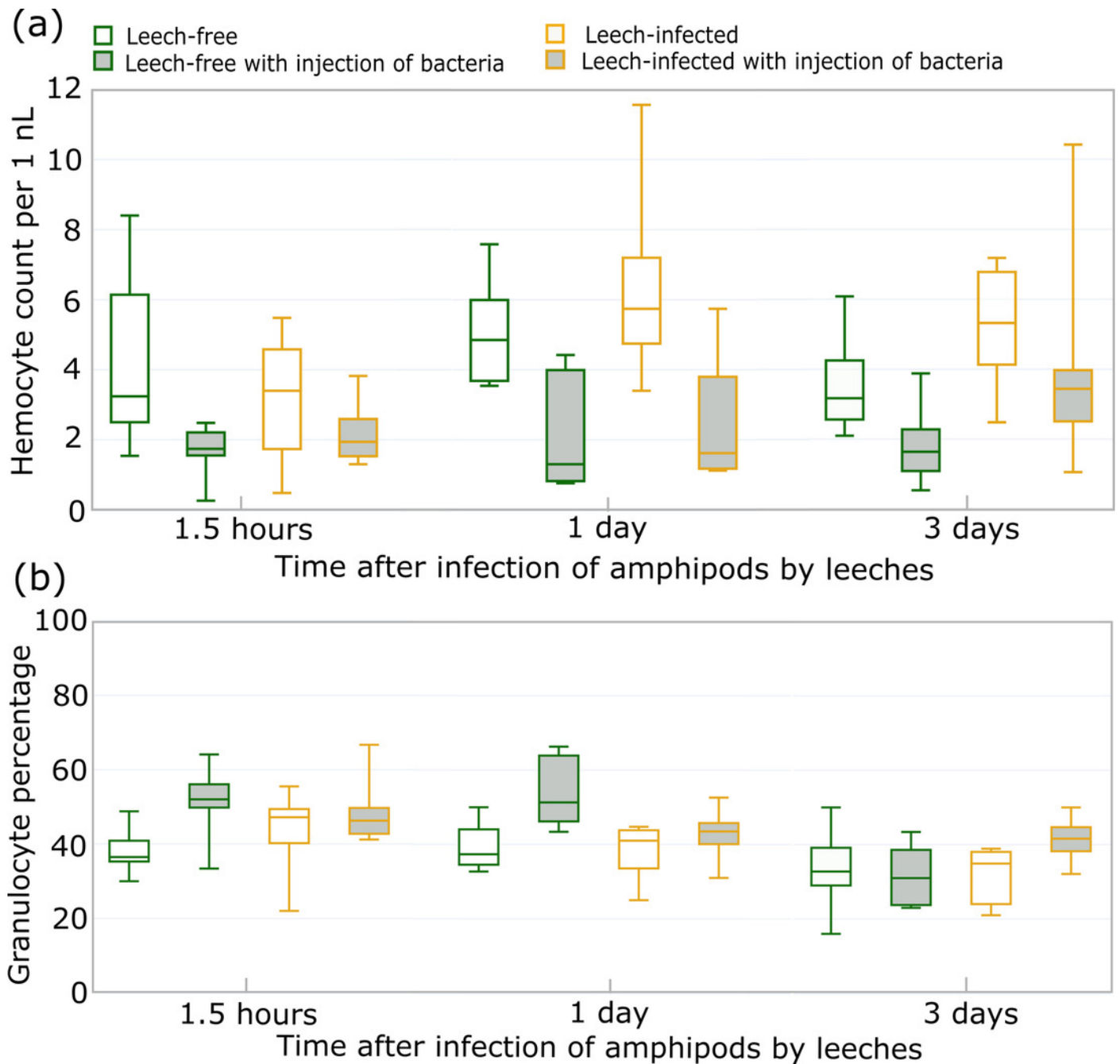


Table 1 (on next page)

Output of generalized linear model with the Gaussian distribution fitted to total hemocyte count and granulocyte percentage in leech-free and artificially leech-infected amphipods with and without bacterial injection (see Figure 7).

All interactions between factors were allowed, but the results only for three independent factors and their statistically significant interactions with substantial effect estimates are depicted here.

1

Variable	Total hemocyte count		Granulocyte percentage	
	Estimate, cells/ μ l	P-value	Estimate, %	P-value
Time	-15.1	0.248	-0.061	0.284
Leech	-514.1	0.537	4.149	0.256
Bacteria	-2846.1	< 0.001 ***	16.457	< 0.001 ***
Leech:Bacteria	603.3	0.601	-12.223	0.019 *

2