

# The crustacean Armadillidium vulgare, a new promising model for the study of cellular senescence

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## 1 The crustacean Armadillidium vulgare, a new

# 2 promising model for the study of cellular senescence

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#### 19 Abstract

20 Senescence, the decline of physiological parameters with increasing age, is a quasi-21 ubiquitous phenomenon in the living world. However, the observed patterns of senescence 22 can markedly differ between across species and populations, between sexes and even among 23 individuals. To identify the drivers of this variation in senescence, experimental approaches 24 are essential and involve the development of tools and new study models. In fact, current 25 knowledge of the senescence process is mostly based on studies on vertebrates and principal 26 information about senescence in invertebrates is mostly limited to model organisms such as 27 *Caenorhabditis elegans* or *Drosophila melanogaster*. In this context, we tested whether 28 biomarkers of vertebrate aging could be used to study senescence in a new invertebrate 29 model: the common woodlouse Armadillidium vulgare. More specifically, we looked for the 30 effect of age in woodlouse on three well established physiological biomarkers of aging in 31 vertebrates: immune cells (cell size, density and viability),  $\beta$ -galactosidase activity, and 32 Telomerase Reverse Transcriptase (TERT) (essential subunit of the telomerase protein) gene 33 expression. We found that the size of immune cells was higher in older individuals, whereas 34 their density and viability decreased, and that the  $\beta$ -galactosidase activity increased with age, 35 whereas the Telomerase Reverse Transcriptase (TERT) gene expression decreased. These 36 findings demonstrate that woodlouse display age-related changes in biomarkers of vertebrate 37 senescence, with different patterns depending on gender. Thus, the tools used in studies of 38 vertebrate senescence can be successfully used in studies of senescence of invertebrates such

39	as the woodlouse. The application of commonly used tools to new biological models offers a
40	promising approach to assess the diversity of senescence patterns across the tree of life.
41	
42	Keywords
43	Cellular senescence, immunosenescence, Telomerase Reverse Transcriptase (TERT), $\beta$ -
44	galactosidase activity.
45	
46	1. Introduction
47	Many theories have tried to explain why senescence is a quasi-ubiquitous phenomenon
48	in the living organisms. For instance, the disposable soma theory proposed the senescence
49	process as a result of damages accumulation over time. These damages are strongly
50	influenced by the environment, leading to trade-offs between the different functions (e.g.
51	between reproduction and somatic maintenance) and shaping a high diversity of senescence
52	patterns across species and populations, among individuals, and between sexes. One current
53	challenge is to understand the selective forces and mechanisms driving this diversity of
54	senescence patterns.
55	At the cellular level, senescence corresponds to the cellular deterioration leading to
56	stop the cellular cycle (Campisi & di Fagagna, 2007). As ageing is associated with cellular
57	senescence (Herbig et al., 2006; Wang et al., 2009; Lawless et al., 2010), many biomolecular
58	parameters potentially inform about senescence and can therefore be valuable tools for

59	studying this process (de Jesus & Blasco, 2012). For example, the evolution of the integrity
60	and efficiency of immune cells is particularly relevant to study cellular senescence because a
61	diminution of the number of effective immune cells with increasing age takes place in both
62	vertebrates (e.g. Cheynel et al., 2017) and invertebrates (e.g. Park et al., 2011). Another
63	marker used to study cellular senescence is the enzymatic activity of the $\beta$ -galactosidase. This
64	enzyme is a hydrolase that transforms polysaccharides in monosaccharides. The lysosomal
65	activity of this enzyme is increased when the cell enters in senescence (Dimri et al., 1995;
66	Itahana et al., 2007). This phenomenon occurs in senescent cells of many organisms ranging
67	from humans (Gary & Kindell, 2005) to honeybees (Hsieh & Hsu, 2011). Another protein
68	linked to the cellular senescence process is the telomerase, a ribonucleo protein complex
69	composed by two essential components, the telomerase reverse transcriptase (TERT) and the
70	telomerase RNA (TR) and other accessorial proteins (Podlevsky et al., 2007). Telomerase
71	lengthens the ends of telomeres (i.e. DNA sequences located at the end of chromosomes that
72	protect chromosome integrity and shorten after each cell division). Cell senescence arises
73	when the telomere length becomes critically short (Chiu & Harley, 1997; Shay & Wright,
74	2005). The telomerase activity depends on organism, age and also tissues (e.g. (Gomes et al.,
75	2010)). For instance, telomerase is active during the development before birth and after only
76	in stem and germ cells in humans (Liu et al., 2007; Morgan, 2013) while in the Daphnia
77	pulicaria, the telomerase activity in all tissues of the body decreases with increasing age
78	(Schumpert et al., 2015). The TERT is essential in the telomerase protein complex and has

been shown to be related to cell survival in humans (Cao *et al.*, 2002). The TERT has also
been detected in numerous species including vertebrates, fungi, ciliates and insects (Robertson
& Gordon, 2006; Podlevsky *et al.*, 2007).

82 As patterns of senescence are strongly diversified within the living world, it seems 83 essential to study organisms displaying markedly different life histories strategies to 84 understand the causes and mechanisms underlying this diversity. Thus, invertebrates are 85 increasingly used in experimental studies of senescence (Stanley, 2012; Ram & Costa, 2018). 86 In addition to share similarities with vertebrates in terms of senescence, they can be 87 manipulated experimentally and they are easier to be monitored throughout their entire 88 lifetime (Ram & Costa, 2018). These advantages make them models of choice for studying 89 senescence. Here, we propose the common woodlouse A. vulgare as a promising new model 90 for studying senescence. Woodlouse can live beyond three years and display sex-specific 91 senescence patterns in natural populations (Paris & Pitelka, 1962). In addition, one study has 92 already reported evidence of immuno senescence in this species (Sicard et al., 2010).

93

In this context, we tested the suitability of  $\beta$ -galactosidase activity, immune cell parameters and the TERT gene expression to cause age-specific responses in the common woodlouse *Armadillidium vulgare*. According to the literature, we expected an increase in  $\beta$ galactosidase activity, and a decrease of both TERT gene expression and immune cell

99 Pitelka, 1962), cellular senescence patterns are also expected to be sex-specific in <i>A. vulgare</i> .	98	viability and density in A. vulgare. As males have higher adult survival than females (Paris &
	99	Pitelka, 1962), cellular senescence patterns are also expected to be sex-specific in A. vulgare.

100

- 101 **2. Materials & Methods**
- 102

#### 2.1. Biological model

103 A. vulgare individuals used in the following experiments were derived from a wild 104 population collected in Denmark in 1982. These animals have been maintained on moistened 105 soil under the natural photoperiod of Poitiers (France 46.58°N, 0.34°E, 20°C) at 20°C fed ad 106 libitum with dried linden leaves and carrots. Crosses were monitored to control and promote 107 genetic diversity. For each clutch obtained, individuals were sexed, and brothers and sisters 108 were separated to ensure virginity. In common woodlouse, individuals molt throughout their 109 lives, with approximately one molt per month. During this process all the cells of the 110 concerned tissues are renewed at 20°C (Steel, 1980). However, the brain, the nerve cord and 111 gonads are not renewed during molting and are therefore relevant candidates for tissue-112 specific study of senescence in this species. Males and females were tested separately to 113 assess the impact of sex.

114

- 115 **2.2Measure of**  $\beta$ **-galactosidase activity**
- 116 Animals

117 To test the impact of age on the on  $\beta$ -galactosidase activity,180 individuals were used: 118 90 young (i.e. 6-months-old, 45 males and 45 females) and 90 old (2-years-old, 45 males and 119 45 females) individuals.

120 <b>Pro</b>	otocol
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121 Individuals were dissected separately in Ringer solution (Sodium Chloride 394 mM, 122 Potassium Chloride 2 mM, Calcium Chloride 2 mM, Sodium Bicarbonate 2 mM) and nerve 123 cord was removed. To obtain a sufficient amount of protein, we made pools of five nerve 124 cords (from five different individuals of the same age). The five nerve cords were filed in 500 125 µL of Lyse Buffer 1X (CHAPS 5 mM, Citric acid 40 mM, Sodium Phosphate 40 mM, 126 Benzamidine 0.5 mM, PMSF 0.25 mM, pH = 6) (Gary & Kindell, 2005), and then were 127 centrifuged at 15000g at 4°C for 30 minutes. The supernatant was taken and kept at -80°C 128 until its utilization. The protein concentration was determined by the BCA assay 129 (Thermofisher) and was homogenized at 0.1 mg/mL. The  $\beta$ -galactosidase activity was 130 measured as described by Gary and Kindell (2005). Briefly, 100 µL of extracted protein at the 131 concentration of 0.1 mg/mL were added to 100 µL of reactive 4-methylumbelliferyl-D-132 galactopyranoside (MUG) solution in a 96 well-microplate. The MUG reactive, in contact to 133  $\beta$ -galactosidase, leads by hydrolysis to the synthesis of 4-methylumbelliferone (4-MU), which 134 is detectable using fluorescent measurements. Measures were performed by the multimode 135 microplate reader Mithras (LB940 HTS III, Berthold; excitation filter: 120 nm, emission filter 136 460 nm) for 120 minutes. Two technical replicates were measured for each pool.

#### 137

141

#### 138 **2.3**Measure of immune cell parameters

#### 139 Animals

140 To test the impact of age on the immune cell parameters (i.e. density, viability, and

size) in A. vulgare, 60 mature individuals were used: 30 young (i.e. 1-year-old, 15 males and

- 142 15 females) and 30 old (3-years-old, 15 males and 15 females) individuals.
- 143 **Protocol**
- 144 To study the impact of age on the immune parameters, a hole was bored in the middle

145 of the 6<sup>th</sup> segment and 3  $\mu$ L of haemolymph were collected (per individual) with an

- 146 eyedropper and deposited promptly in 15 μL of anticoagulant solution(MAS-EDTA (EDTA 9
- 147 mM, Trisodium citrate 27 mM, NaCl 336 mM, Glucose 115 mM, pH 7, (Rodriguez et al.,
- 148 1995))). Then, 6 µL of Trypan blue at 0.4% (Invitrogen) were added to color the dead cells.
- 149 Thereafter, 10 µL of this solution were deposed in counting slide (Invitrogen Coutness®,
- 150 Thermofisher). The immune cell density, the immune cell viability and the immune cell size
- 151 were evaluated using an automated Cell Counter (Invitrogen Countess®).
- 152

#### 153 **2.4 Measure of TERT gene expression**

The identification of the Telomerase Reverse Transcriptase (TERT)gene was firstly performed from the *A. vulgare* genome (Chebbi *et al.*, 2019). In order to check whether this gene was present and preserved in crustaceans, phylogenetic analyses were carried out

157	upstream (see Supplementary materials 1, 2, 3 and 4). This gene has been found in crustacean
158	transcriptomes and the topology of the TERT gene tree follows the phylogenetic relationships
159	between the involved species (Supplementary material 3), suggesting a conserved role of the
160	TERT gene.
161	Gene expression
162	Animals
163	We tested the effect of age on the expression of TERT gene within 4 different age
164	groups: (1) 4-months-old, (2) 1-year-old, (3) 2-years-old and (4) 3-years-old. Females and
165	males were tested separately by pools of 5 individuals in 1-, 2-, 3-years-old groups and by
166	pools of 7 individuals in 4-months-old group. All conditions require 4 replicates for each sex.
167	176 individuals were used for this experiment. For each group we tested the expression level
168	of the TERT gene in two different tissues: the nerve cord (somatic line) and gonads (germinal
169	line).
170	Protocol
171	Animals were washed by immersion for 30s in a 30% sodium hypochlorite solution
172	followed by two 30s immersion in distilled water. Tissues were dissected in Ringer solution
173	(Sodium Chloride 394 mM, Potassium Chloride 2 mM, Calcium Chloride 2 mM, Sodium

174 Bicarbonate 2 mM) and deposited by specific tissues pools of 5 on TRIzol reagent 175 (Invitrogen) to extract RNA according to the manufacturer's protocol after a cell

176 disintegration using a Vibra Cell 75,185 sonicator (amplitude of 30%). Total RNA was

177	quantified by NanoDrop technology and was stored at -80°C until use. Reverse transcriptions
178	(RT) were made from 500ng of RNA previously extracted and using the kit SuperScript <sup>TM</sup> IV
179	Reverse Transcriptase (Thermo Fisher Scientific) according to the supplier's instructions.
180	Primers were designed using the identified gene: primer TERT_F: 5'-
181	AGGGAAAACGATGCACAACC-3' and primer TERT_R: 5'-
182	GTTCGCCAAATGTTCGCAAC- 3' (see Supplementary material 1). Quantitative RT-PCR
183	was performed using 0.6 $\mu l$ of each primer (10 $\mu M$ ), 2.4 $\mu l$ of nuclease-free water and 1.5 $\mu l$ of
184	cDNA template and the LightCycler LC480 system (Roche) with the following program:10
185	min at 95 °C, 45 cycles of 10 s at 95 °C, 10 s at 60 °C, and 20 s at 72 °C. Expression levels of
186	target genes were normalized based on the expression level of two reference genes previously
187	established: the Ribosomal Protein L8 (RbL8) and the Elongation Factor 2 (EF2) (Chevalier
188	<i>et al.</i> , 2011).

189

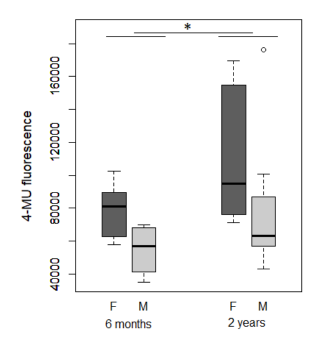
### 190 Statistics

191 All statistical analyses were performed using the R software (R. Core Team, 192 2016). The  $\beta$ -galactosidase activity was analyzed with linear mixed effect models using the 193 package lme4 (Bates *et al.*, 2014). As two technical replicates were measured for each pool, 194 the model including the pools fitted as a random effect, age and sex and their two-way 195 interaction as fixed factors.

196	Concerning the immune parameters, linear models with Gaussian distribution were
197	fitted to analyze variation in the cell size and viability. For the cell density, a linear model of
198	the cell number (log-transformed, (Ives & Freckleton Robert, 2015)) was fitted.
199	The level of TERT expression according to age in the two different tissues were
200	compared by a Kruskal-Wallis rank sum test in combination with Nemenyi's post hoc
201	multiple comparison test with the Tuckey correction using R package PMCMR.
202	
203	3. Results
204	β-galactosidase activity
205	The $\beta$ -galactosidase activity was higher in old (i.e. 2-years-old) than in young (i.e. 6-
206	months-old) individuals ( $\chi^2_1$ =6.15, p=0.013, Figure 1). We also detected a higher $\beta$ -
207	galactosidase activity in females than in males ( $\chi^2_1$ =7.26, p=0.007, Figure 1).
208	The thick line depicts the median, the box the interquartile range, and the whisker are bounded to the most
209	extreme data point within 1.5 the interquartile range. The outliers outside this range are displayed as open
210	circles. N= 24 pools of 5 individuals. * denotes $p < 0.05$

211

212





214 Figure 1: β-galactosidase activity according to age and sex in A. vulgare (F=females, M=males)

215

#### 216 **Immune cells parameters**

Cell size was larger in 3-years-old than in 1-year-old individuals ( $F_{1,58}$ =8.54, p=0.005, Figure 2A). Conversely, the cell density was higher in 1-year-old than in 3-years-old individuals ( $F_{1,58}$  =4.33, p=0.01, Figure 2B). Concerning the immune cell viability, a statistically significant interaction occurred between age and sex, with a relatively lower immune cell viability in 3-years-old females ( $F_{3,56}$ =6.85, p=0.01, Figure 2C). No sex effect was detected on cell size ( $F_{2,57}$ =0.76, p=0.38, Figure 2A) or cell density ( $F_{2,57}$ =0.32, p =0.57, Figure 2B).

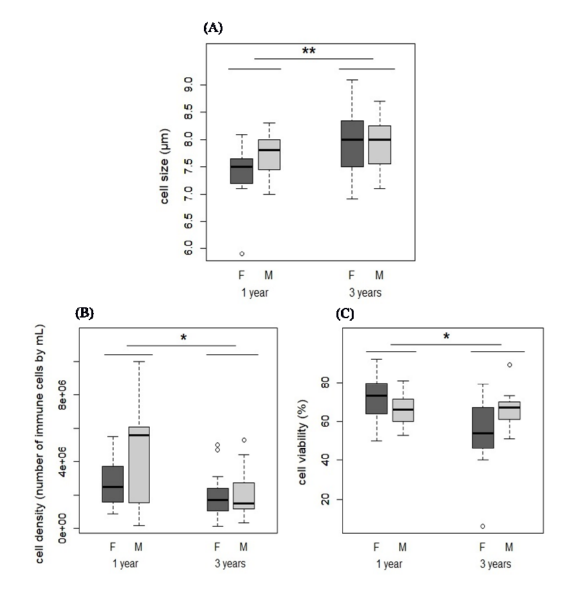




Figure 2: Immune cell size (A), density (B) and viability (C) according to age and sex in A. vulgare
(F=females, M=males)

Thethick line depicts the median, the box the interquartile range, and the whisker are bounded to the most extreme data point within 1.5 the interquartile range. The outliers outside this range are displayed as open circles.N= 60 individuals: 15 1-year-old females, 15 1-year-old males, 15 3-years-old females and 15 3-years-old males. \* denotes p<0.05 and \*\* denotes p<0.01

231

#### 232 **TERT gene expression**

The TERT gene expression decreased with increasing age in nerve cords ( $\chi^2_3$ =23.30, 233 234 p<0.001, Figure 3A). More precisely, the TERT gene expression was higher in 4-months-old individuals compared to 2-years-old and 3-years-old individuals (p=0.001 in both cases) and 235 236 in 1-year-old individuals compared to 3-years-old individuals (p=0.038), without any detectable sex effect ( $\chi^2_1$ =0.14, p=0.70, Figure 3A). In gonads, the TERT gene expression 237 was much higher in females ( $\chi^2_1$ =17.81, p<0.001, Figure 3B) and tended to decrease with 238 increasing age ( $\chi^2_3$ =7.5, p=0.057, Figure 3B) as the TERT gene expression tended to be 239 240 higher in 4-months-old females compared to 3-years-old females (p=0.054). In males, a general tendency was also observed ( $\chi^2_1$ =7.34, p=0.061, Figure 3B), the TERT gene 241 242 expression tending to be higher in 2-years-old individuals compared to 1-year-old and 3-

243 years-old individuals (p=0.14 and p=0.12, respectively, Figure 3B).

244

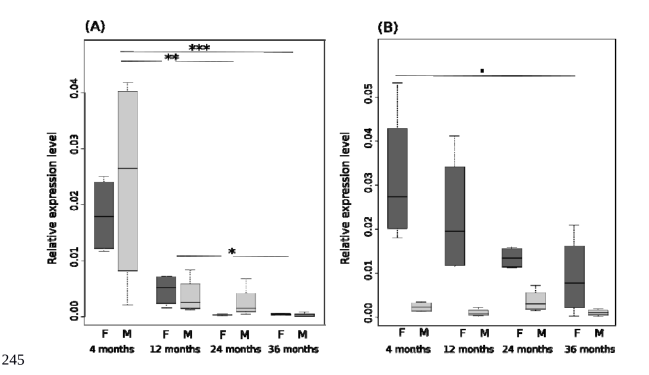


Figure 3: Relative expression level of TERT in (A) nerve cords and (B) in gonads in A. vulgare (F=females,
M=males.

Expression of each gene was normalized based on the expression of Ribosomal Protein L8 (RbL8) and Elongation Factor 2 (EF2) as reference genes. The thick line depicts the median, the box the interquartile range, and the whisker are bounded to the most extreme data point within 1.5 the interquartile range. N= 176 individuals: 284-months-old females, 28 4-months-old males, 20 1-year-old females, 20 1-year-old males, 20 2-years-old females, 20 2-years-old males, 20 3-years-old females, 20 3-years-old males. . denotes p<0.10, \*\* denotes p<0.01

253

**4. Discussion** 

In this study, we tested several effective physiological biomarkers of vertebrate senescence to assess whether they could also be used in invertebrates such as the common woodlouse. Immune cells showed an increase in their size and a decrease in their density and

258	viability with increasing age. In nerve cords, the activity of the $\beta$ -galactosidase enzyme
259	increased, whereas the TERT gene expression decreased with increasing age. These results
260	support the presence of increasing cellular senescence in A. vulgare with chronological age. In
261	contrast, in the gonads, the TERT gene expression was too low in males and was not
262	sufficiently variable between sexes to provide information on the cellular senescence status in
263	this tissue.

Our study is in line with previous studies that have already revealed the possibility of using vertebrate biomarkers in invertebrates (Hsieh & Hsu, 2011; Park *et al.*, 2011; Schumpert *et al.*, 2015). By testing a set of different physiological biomarkers of vertebrate senescence, often studied independently, our study supports both ideas that routinely used biomarkers in vertebrates can be adapted in invertebrates and that the senescence process is quasi-ubiquitous in the living world and can be expressed in a similar way in very different organisms.

Previous studies have shown that the probabilities to survive decrease with increasing age in *A. vulgare* (Paris & Pitelka, 1962). The cellular damages accumulated during the animal's life could be the cause of cell senescence and therefore the driving force behind actuarial senescence. (Harman, 1956; Barja, 2000; Barja & Herrero, 2000; Finkel & Holbrook, 2000). In *A. vulgare*, the 2- and 3-years-old individuals could have therefore accumulated more cellular damages during their lifetime, leading to the cellular senescence we report.

278	Our study also revealed a strong difference between sexes on the response of
279	biomarkers to age changes. At a given age, females display higher $\beta$ -galactosidase activity
280	and lower immune cell viability than males. Between-sex differences in lifespan have been
281	reported in A. vulgare with a longer lifespan in males than in females (Geiser, 1934; Paris &
282	Pitelka, 1962). Exact differences in actuarial senescence patterns (i.e. age-specific changes in
283	survival probabilities) remain to be quantified in A. vulgare but such differences are quite
284	common both in vertebrates and invertebrates (Tidière et al., 2015; Marais et al., 2018). One
285	of the main theory proposed to explain sex differences in longevity or senescence patterns
286	relies on different resource allocation strategies between sexes (Vinogradov, 1998;
287	Bonduriansky et al., 2008). The shorter lifespan in females A. vulgare, that allocate more
288	energy to reproduction than males (Paris & Pitelka, 1962) because they carry their offspring
289	in their marsupium during one month giving nutrients and protection, supports a role of
290	differential sex allocation.

Sex differences in resource allocation strategies could also be driven by environmental conditions (Shertzer & Ellner, 2002). Our physiological biomarkers of vertebrate senescence revealed sex differences, and as supported in Depeux et al., 2019, they could constitute useful tools to identify other factors involved in variations in senescence patterns, such as environmental stressors. Moreover, if these biomarkers seem to predict better the physiological age than chronological age notably in terms of survival and reproduction, they

297 could correspond to biomarkers of senescence in woodlouse (Baker & Sprott, 1988; Simm *et*298 *al.*, 2008; Sprott, 2010).

299	Our present study demonstrated that the physiological biomarkers of vertebrate
300	senescence respond to age changes in the common woodlouse, a new invertebrate model of
301	aging. These parameters that predict the chronological age of woodlouse individuals might
302	offer reliable biomarkers, especially if their measurements are related to both reproductive
303	and survival prospects more than to the chronological age of individuals. In this context, and
304	more broadly in the study of senescence and of the factors involved in its diversity, the
305	woodlouse model, which has physiological similarities with other invertebrates, could be a
306	model of choice to study sex-specific actuarial and cellular senescence.

307

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313

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321	
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