The dynamics of group formation among leeches

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INTRODUCTION

Animals from simple invertebrates to mammals and humans, not only interact with the environment but establish social interactions with their conspecifics (Anstey et al., 2009; Makris et al., 2009; Nagy et al., 2010; Sokolowski, 2010; Zhao et al., 2011). In lower invertebrates these interactions lead to a swarming behavior with the formation of groups of several tens and even thousands of conspecifics adopting the same behavior. Locusts can switch from a solitary to a social behavior, advantageous when competing for limited resources or during migration (Buhl et al., 2006; Bazazi et al., 2008; Anstey et al., 2009). Several species of fish form large shoals (Blaxter and Hunter, 1982; Hoare et al., 2000) often before migration (Makris et al., 2009). Birds form flocks able to fly collectively (Ballerini et al., 2008; Nagy et al., 2010). We have studied the emergence of group formation in leeches for two reasons: firstly, because it is possible to quantify their behavior easily and precisely (García-Pérez et al., 2005; Mazzoni et al., 2005; Bisson and Torre, 2011). Secondly, the nervous system of the leech Hirudo medicinalis has been extensively studied (Macagno, 1980; Muller et al., 1981) so that it is possible to relate its behavior to underlying neuronal networks (Kristan et al., 2005) and specific neuromodulators, such as serotonin (5-HT), dopamine (DA), and octopamine (Willard, 1981; Puhl and Mesce, 2008). In the present manuscript, we quantify the dynamics of formation of leech groups. When the density of conspecifics is low (less than five conspecifics in 700 cm²) leeches move independently, but when their density is doubled the formation of stable groups is observed. The critical density for group formation is lowered by elevating the level of intracellular 5-HT inside the leech bodies. Ablation experiments show that a neurobiological signal modulating group formation is initiated by sensory receptors located near the leech’s head and processed in the supraesophageal ganglion. We have detected an elevation of the level of endogenous 5-HT in leeches forming a large group and this observation raises the possibility that a somatic, i.e., an extrasynaptic release of 5-HT is involved in group formation. Indeed extrasynaptic release of 5-HT able to diffuse to remote receptor sites has been detected and characterized in several preparations where is associated to functional roles (Bunin et al., 1998; Bunin and Wightman, 1999; De-Miguel and Trueta, 2005; Kaushalya et al., 2008). At a formal level, the formation of stable groups can be explained by the establishment of a reinforcement dynamics (Cattuto et al., 2009; Stehlé et al., 2010) depending on the leech density and the intracellular level of 5-HT, which can be modeled in a mathematical way, very similar to that used to describe human aggregation (Stehlé et al., 2010; Zhao et al., 2011).

MATERIALS AND METHODS

ANIMALS AND PREPARATIONS

Adult leeches (Hirudo verbana) obtained from Ricarimpex (Eysines, France) were housed in groups of about 30 animals in 20-l tanks of artificial pond water [Instant Ocean salts (Aquarium Systems) diluted with deionized water to 1/1000 of ocean strength]. Leeches were kept in a circular tank (diameter: 30 cm, height: 6 cm), filled with artificial pond water kept at about 16°C, under daily illumination, and used within few months after shipping. Since Ricarimpex breeds leeches outdoors, we assume that they retain the seasonal rhythm established during development. Observation tanks were illuminated with a circular array of white light LEDs which provided dim, diffuse illumination with no abrupt spatial and/or temporal gradients. No additional sensory stimuli, such as chemical, mechanical, or visual inputs were intentionally applied. A transparent, plastic disk was used to keep leeches inside the tank.
Determination of the level of 5-HT by HPLC
Serotonin (5-hydroxytryptamine, 5-HT) was quantified in extracts of chains of leech ganglia by reverse-phase high performance liquid chromatography (HPLC). Leeches were positioned in the usual observation tank and after 3–5 h leeches which formed a group of at least eight conspecifics (gregarious leeches) and those who remained solitary were selected for the determination of their body level of 5-HT. Solitary and gregarious leeches were immediately submerged in liquid nitrogen for approximately 1 or 2 min and frozen leeches were placed on an ice cooled block. Chunks of ganglia from the tail to the head ganglion – usually composed by around 20 individual ganglia – were dissected out and placed in a 100-μl microhomogenizer containing 50 μl of ice-cold 0.15 M perchloric acid and homogenized for 3 min on ice. The homogenate was transferred to 1.5 ml Eppendorf tubes using a Hamilton syringe and centrifuged at 17500 g for 20 min at room temperature. The supernatant was transferred to a second Eppendorf tube and stored at −80°C until HPLC analysis. A sample of 100–200 μl of their blood was also extracted and treated as treated as chains of ganglia. 5-HT was quantified by reference to external standards.

Delivery of Neuromodulators
Some leeches were injected with specific amounts of 5-HT, ketanserin, or mianserin (Sigma-Aldrich). All neuromodulator stock solutions (5-HT: 1 mM, mianserin: 10 mM, ketanserin: 5 mM) were prepared using distilled water (except ketanserin, which was dissolved in 0.1 M HCl solution), stored at −20°C in 500 μl aliquots and defrosted in 15 min at room temperature. The final concentration was reached by dilution in our normal leech saline solution (in mM: 116 NaCl, 1.8 CaCl2, 4 KCl, 1.5 MgCl2, 10 glucose, 10 Tris maleate buffered to pH 7.4) immediately before use and delivery. We injected 300 μl of 200 μM 5-HT solution using 1 ml insulin syringes with 29 GA and 15 mm long needles. Injection of 150 μl containing 1 mM ketanserin or 2 mM mianserin solutions were followed by another injection of 150 μl containing 400 μM 5-HT solution. Each leech was gently held in one hand and the needle was inserted under the skin of the dorsal side, along its dorsal axis at a depth of about 10% of the body thickness. We calculated that the injected solution was diluted into the extracellular space roughly 25–30% of the leech volume, usually 3–4 ml. Therefore, the final drug concentration injected in the leeches is approximately one-fourth of the injected drug concentration. The body volume of an adult leech is 3–4 ml, while the overall blood volume is about 8–9% of the body mass, corresponding to 240–360 μl (Wenning and Meyer, 2007). Changes of body volume are well tolerated by leeches because they can increase their initial weight by 8–11 times during feeding (Dickinson and Lent, 1984), it is likely that an increase of the body volume (about 10% of its body mass) caused by the injection did not impair the animal’s movements.

Ablation Studies
Leeches were anesthetized with 8% ethanol in leech Ringer’s solution and iridectomy scissors were used to cut away the band of dorsal lip organs visible in the light microscope, with 1 or 2 mm of skin on either side including the rows of smaller lip organs detected with the scanning electron microscope (Elliott, 1986) but not visible in the light microscope. Sham operations were performed in the same way, except that a band of skin dorsally adjacent to the lip organs (along the dorsal surface of the head) was removed and the lip organs themselves were left intact. Leeches were allowed to recover for 1 day in water. The day after surgery, ablated leeches, injected with 300 μl of 200 μM 5-HT and sham-ablated leeches, were tested, along with normal controls, for social interactions.

Imaging
We used a color CCD camera (640 × 480 pixels of image size; model DFK 21BF04; The Imaging Source Europe) to image leeches from above and to monitor their movements (Figure 1A). The camera was connected via the FireWire-output to a frame grabber (PCI-1394; Texas Instruments) installed on a personal computer, able to process images in real-time. Leeches’ bodies were tracked at 7.5 Hz in RGB mode using a software program developed in Matlab language (MathWorks, Natick, MA, USA). The tracking algorithm is composed of two subroutines. The first subroutine is executed in real-time and extracts the borders of all the groups of leeches. Its purpose is to detect and store leech shapes. Each frame was converted into grayscale by means of the average method (i.e., by taking the average of the Red, Green, and Blue signals; Pratt, 1991). The resulting 8 bit grayscale image has values ranging between 0 (corresponding to white) and 255 (corresponding to black). A threshold equal to 50 is then applied to the grayscale image: all the pixels that have values major or equal to this threshold are set to value 255, 0 otherwise. We refer to this resulting image as the binary version of the frame (in which its pixels can assume only two possible values) and each leech – or group of leeches – appears as a black shape or blob. The extraction of the closed contours of each blob was achieved by applying a Radial Sweep algorithm (a contour tracing algorithm; Pavlidis, 1995) to the image, implemented in the Matlab function bwboundaries (Image Processing Toolbox, MathWorks, Natick, MA, USA). Each closed contour defines a blob representing a single leech or a group of leeches (Figures 1E,F, green lines). Blobs with an area below a threshold of 20 pixels2 (usually caused by leech shadows) were discarded, as the average area of a single, small leech, is major than 30 pixels2. The subroutine stores only the pixels belonging to the contour of the detected blobs so that it was possible to monitor leeches behavior for several hours, minimizing the storage size of the recording file.

The second subroutine is performed off-line and processes the previously extracted contours. Its purpose is to determine the number of leeches belonging to the blobs identified by their contour. In the generic frame k, each blob Bk can be labeled as the ith blob (with i that ranges between 1 and N, where N is the total number of blobs in the kth frame). If we overlap the frame k over the frame k − 1 we will see that blobs representing a non-moving leech or groups of leeches will match almost perfectly (the small differences being caused by recording noise), whereas blobs representing moving leeches or groups of leeches will share just a portion of their area (usually minor than 80%). Moreover, two blobs observed in frame k − 1, that merge (Figure 1E) between frames k − 1 and k will partially overlap with the resulting blob observed in frame k. Finally, a single blob observed in frame k − 1,
FIGURE 1 | The experimental set-up. A color CCD camera imaged from above the observation tank containing 10 leeches. (A) During the first 2h, leeches inspected the environment. (B) Following the searching phase a stable large group of leeches in mutual contact was observed. (C,D) Images shown in (A,B), after processing. Numbers in (C,D) indicate the number of conspecifics present in each group. A real-time algorithm extracted the borders of all leech groups (black blobs) and an off-line algorithm determined the number of leeches in each blob (black numbers). During the process of group formation, two kinds of events occurred: two groups could merge (E) forming a larger group, or a single group could split (F) forming two groups. During the searching phase, merge (G) and split (H) rates have similar values, because leeches are highly active and no formation of stable groups can be observed. Inset in panel h shows a zoom of Merge (black) and Split (red) on an enlarged scale. Merge and split plots are very similar because they are evaluated on a sliding time window of size 5 min. Subtle differences between these two signals determine the group formation dynamics.

that splits (Figure 1F) between frames \(k - 1\) and \(k\) will partially overlap with the resulting blobs observed in frame \(k\). The area of intersection between two blobs \(j\) and \(i\) (the first belonging to frame \(k - 1\), \(B_{i}^{k-1}\), and the second to frame \(k\), \(B_{j}^{k}\)) can be calculated by performing a multiplication between the binary image containing only \(B_{i}^{k-1}\) and the binary image containing only \(B_{j}^{k}\). These two images can be obtained by filling the contours of the two blobs, using a polygon fill algorithm (Hearn and Baker, 1997) implemented in the Matlab function poly2mask. The multiplication is equivalent to performing a logical AND operation: the number of resulting pixels is the area of intersection between the two blobs.

We can now define the overlap matrix \(\Omega_{ij}^k\) as:

\[
\Omega_{ij}^k = \frac{A\left(B_{i}^{k-1} \cap B_{j}^{k}\right)}{A\left(B_{i}^{k-1}\right)}.
\]

where \(A\) stands for “area” and the symbol \(\cap\) indicates intersection between blobs: the element \(\omega_{ij}\) of the overlap matrix represents the area of intersection between the blob \(i\) in frame \(k\) and the blob \(j\) in frame \(k - 1\) divided by the area of the latter blob. The overlap matrix has size \(N \times M\), where \(N\) is the number of blobs in frame \(k\) and \(M\) the number of blobs in frame \(k - 1\). If, for example, row number 3 has more than one non-zero entry, then blob number 3 in frame \(k\) is formed by a merge of two or more blobs in frame \(k - 1\) (Figure 1E). If, for example, column number 5 has more than one non-zero entry, then blob number 5 in frame \(k - 1\) splits in two or more blobs in frame \(k\) (Figure 1F). The central assumption of the algorithm is that the number of leeches belonging to blob \(B_{i}^{k-1}\) in frame \(k - 1\), multiplied by the non-zero entries \(\omega_{ij}\) of the overlap matrix \(\Omega_{ij}^k\) give a non-integer estimate of the number of leeches belonging to blobs \(B_{j}^{k}\) that have non-zero area of intersection with \(B_{i}^{k-1}\). This assumption is supported by the experimental evidence that a leech, in a time interval of 130 ms, cannot vary its shape area in a substantial way and it can be viewed as a rigid body. Let \(g_{0}\) be the \(M \times 1\) vector containing the number of leeches belonging to each of the \(M\) blobs found in frame 0. The non-integer estimate of \(g_{1}\) (i.e., the \(N \times 1\) vector containing the number of leeches belonging to each of the \(N\) blobs found in frame 1) is given by:

\[
g_{1} = \Omega_{1} \cdot g_{0}.
\]

This estimate is also affected by additive noise \(\mu\):

\[
g_{1} = \Omega_{1} \cdot g_{0} + \mu.
\]

Minimizing the effect of the additive noise \(\mu\) on the estimate \(g_{1}\) corresponds to minimizing the quantity:
\[(g_1 - \Omega^1 \cdot g_0)^2\]  

(4)

This, in turn, corresponds to solving a quadratic programming problem

\[g^* = \arg\min \left\{ (g_1 - \Omega^1 \cdot g_0)^T I (g_1 - \Omega^1 \cdot g_0) \right\} + 0.5\]  

(5)

Where \(g^*\) is the integer estimate of \(g_1\) (obtained using the floor operator), i.e., the estimate of the number of leeches belonging to the blobs observed in frame 1. The design matrix \(I\) (identity matrix) is also positive-definite: this ensures that the problem has a global minimizer. However, the problem needs to be constrained: first, the size of two blobs in frame 1 that are the result of a split of a blob in frame 0 must be equal to the size of the single blob in frame 0. Moreover, the sum of all blob sizes in every frame must be equal to the number of leeches in the tank. A constraint on the merge was not necessary, since it is included in the latter constraint. These two constraints can be included in a single equation, where \(\Lambda_{eq}\) is a matrix whose entries are either 0 or 1, opportune\n
derived from \(\Omega^1\) (its last column having all entries equal to 1, in order to describe the second constraint) and the first column \(b_{eq}\) is \(g_0\) while all the entries of the second column are equal to the number of leeches in the tank. Finally, we also constrained the number of leeches belonging to a blob, to stay between two reasonable bounds that are dependent on the area of the blob itself: for example, a blob with area 60 pixels\(^2\) is likely to contain one leech or at the most three leeches well in contact. We estimated the normal distributions of the areas of a group, given its size (for size ranging from 1 to 20). Then, each blob area was assigned to the closest area distribution, using the Mahalanobis metric.

\[d = \frac{(\alpha - \mu)^2}{\sigma}\]  

(6)

Where \(\mu\) and \(\sigma\) are the mean and the standard deviations of the area distribution under consideration: a distance is calculated for each of the 20 distributions, and the minimum one was retained. The closest area distribution (in terms of the Mahalanobis distance) determined the “most likely” size of the considered blob. Thus, the lowest bound \(l\) for its size was the size associated to the most likely area distribution to which it belongs, minus 2, while the upper bound \(u\) is the size associated to the most likely area distribution to which it belongs, plus 2.

\[\begin{align*}
\begin{cases}
\begin{aligned}
\text{argmin} & \left\{ (g_1 - \Omega^1 \cdot g_0)^T I (g_1 - \Omega^1 \cdot g_0) \right\} + 0.5 \\
\Lambda_{eq} \cdot g^* &= b_{eq} \\
l \leq g^* &\leq u
\end{aligned}
\end{cases}
\end{align*}\]  

(7)

The quadratic programming problem is solved by using the Matlab function quadprog, starting from an initial condition: the sizes associated to each blob in frame 0, which are manually set by the operator, i.e., \(g_0\). The algorithm estimates the group sizes of the blobs in frame 1, then in frame 2 and so on. We applied this algorithm to 89 experiments lasting for a time that varied from 1 h to more than 8 h with a number of leeches varying from 5 to 20. The structure of the problem and the constraints imposed seem to guarantee a good performance of the algorithm, with an error of ±1 leech per blob of size major than 1, at the most.

**THE MODEL OF GROUP FORMATION**

The model of group formation describes the dynamics of splits and merges of leech groups. The model is based on the reinforcement mechanism already used to model face-to-face human interactions (Stehlé et al., 2010; Zhao et al., 2011). The reinforcement mechanism implies that the longer is the lifetime of a group the less likely it is that its composition changes. This mechanism is responsible for the broad distribution of lifetime of social groups observed in the present experiments with leeches as well as in human face-to-face interactions (Cattuto et al., 2010). In addition to this mechanism we include a second reinforcement mechanism mediated by the role of 5-HT level in the blood, in agreement with the experimental observations reported in the present manuscript. We assume in particular that the level of 5-HT in the blood increases the leech sociability by increasing the probability of merges. Moreover we assume that, as the number of social interactions increases, the level of 5-HT in the blood of the animals increases.

We start from initial conditions in which leeches are connected in groups of size 2. Each group \(i\) is characterized by its size \(n_i\) and its duration \(\Delta t = t - t_i\) where \(t_i\) is the last time in which group \(i\) was modified. At each time step first we update the following parameters determining the dynamics of group formations (splits and merges of social groups):

- The variable \(s\), indicating the 5-HT level in the blood of the animals increases proportionally to the number of contacts \(n^2\) (with multiplicative constant \(c_1\)) and is at the same time degraded with probability \(c_2\). Therefore \(s\) is updated at each time according to

\[s \rightarrow (1 - c_2) s + c_1 \langle n^2 \rangle;\]  

(8)

- The parameter \(\lambda\) modulates the probability that there is a merge in the dynamics of group formation. We assume that \(\lambda\) is activated by \(s\), the 5-HT level in the blood of the animals, and therefore we take

\[\lambda = \frac{s}{s + K_{met}};\]  

(9)

The constant \(K_{met}\) sets the typical scale for the 5-HT level in the blood to activate the “social behavior” and enhances the probability of merges in the group formation dynamics.

At each time we run the dynamics of group formation:

- We choose one random group \(i\);

One conspecific of group \(i\) splits from the group with probability

\[D_n (\Delta t) = \frac{a_n}{\Delta t + b_n}.\]  

(10)

This probability encodes for a reinforcement dynamics responsible for the broad distribution of group lifetimes.
longer leeches interact with each other the less likely they are to split from their own group. We adopt a minimal hypothesis, summarized by Eq. 10 previously used to model face-to-face social interactions in humans during scientific conferences (Steinhil et al., 2010; Zhao et al., 2011) The constants $a_n$ and $b_n$ are responsible for the shape of the lifetime distribution of a group of size $n$.

- If the split occurs, two options are allowed:
  - The animal merges with another group $j$ of size $n_j$ and duration $\Delta t_j$ with probability proportional to $\lambda$.
  $\prod_j = \lambda \sum_j \frac{D_{n(j)}(\Delta t_j)}{\sum \Delta t_r}$. \hspace{1cm} (11)
  - The animal remains isolated with probability $(1 - \lambda)$.

Finally we update the time of this system according to the expression

$$t \rightarrow t + \frac{1}{\text{Number of groups}}$$ \hspace{1cm} (12)

RESULTS

We positioned up to 20 leeches H. medicinalis in an observation tank filled with artificial pond water. The tank had a diameter of 30 cm with a height of 6 cm, where leeches could swim, crawl, and search around. A CCD camera viewing the observation tank from above allowed a precise quantification of leech motion and behavior (Garcia-Perez et al., 2005; Mazzoni et al., 2005; Bisson and Torre, 2011). When 10 leeches were in the tank, they inspected the environment (Figure 1A). During this phase, leeches swam (red arrows) or explored the new environment with their head, keeping the tail sucker attached to the bottom of the tank (blue arrow). After 2–3 h leeches appeared to the viewing CCD camera aggregated in a compact silhouette (Figure 1B) formed by almost all the conspecifics. When the number of leeches was less or equal to 5, after an initial transient exploration, leeches stopped moving and rested in isolation. Therefore leeches, as several other species (Buhl et al., 2006; Ballerini et al., 2008; Bazazi et al., 2008; Anstey et al., 2009; Makris et al., 2009; Nagy et al., 2010) exhibit a transition from solitary to social behavior when their density is above a critical threshold.

In order to quantify the dynamics of group formation, we developed a suitable software: Images (640 × 480 pixels) were acquired at 7.5 Hz and binarized (Figures 1C,D) and an algorithm determined in each image $I_i$ ($i = 1,\ldots, K$) the number of distinct groups $N_n(i)$ formed by $n$ conspecifics (see numbers near black shapes in Figures 1C,D). The algorithm was also able to determine merges, i.e., when two groups of leeches unified in a single group (Figure 1E) and split, i.e., when a group of leeches broke in two subgroups (Figure 1F). From these data we determined:

- $M(i\Delta t)$ and $S(i\Delta t)$ the number of merges($M$) and splits($S$) in the time widow $[i\Delta t, (i + 1)\Delta t]$ (Figures 1G,H).
- $N^r(i\Delta t)$ the mean number of groups in the time widow $[i\Delta t, (i + 1)\Delta t]$ (Figure 2B).
- $<N_n>$ $i\Delta t)$ the mean size of groups in the time widow $[i\Delta t, (i + 1)\Delta t]$ (Figure 2C).

The algorithm was also able to determine a suitable software: Images (640 × 480 pixels) were acquired for additional 3–6 h, i.e., the usual duration of an experiment. We investigated in detail the dynamics when 5 (7 experiments), 6 (3 experiments), and 10 (10 experiments) leeches were in the observation tank. The group size distribution under the assumption of independent behavior (i.e., constant merge and split rates, regardless group size) is expected to be proportional to $\exp(-\lambda n)$, where $\lambda$ is a constant and $n$ is the group size(Gueron and Levin, 1995). When less than five leeches are in the observation tank, the frequency $f(n)$ of finding a group of $n$ leeches is proportional to $\exp(-\lambda n)$ blue bars in Figure 2M, theoretical fit in red line) in agreement with the notion of an independent behavior when the leech density is below a given threshold (Bisson and Torre, 2011). When the number of conspecifics in the tank is increased to more than five, $f(n)$ is not proportional anymore to $\exp(-\lambda n);$
FIGURE 2 | Statistics of group formation. (A) Time evolution of merges (black line) and of splits (red line) in a time window of 5 min. (B) Time evolution of number of groups in a time window of 1 min (gray line) and in 5 min (black line). (C) Time evolution of average group size in a time window of 1 min (gray line) and in 5 min (black line). (A–C) Refer to experiments with five leeches. (D–F) Time evolution of density of groups over 4 h for three different experiments. Color coded scale from dark blue (0 groups) to deep red (5 groups). (G–L) As in (A–F) but with 10 leeches. Inset in panel g shows a zoom of Merge (black) and Split (red) on an enlarged scale. Merge and split plots are very similar but not identical and small differences determine group formation. (M–O) Probability density of group size in the presence of 5, 6, and 10 conspecifics respectively. Plots in (A,G) are averages from 7 and 10 experiments, respectively. Plots in (B–F) and (H–L) are based on a single experiment. Plots in (M–O) are averages from 7, 3, and 10 experiments, respectively.
Figures 2N,O, theoretical fit in red line), indicating the onset of a cooperative behavior in which individual leeches do not remain solitary for long. In order to identify the neurobiological basis of this behavior, we asked whether this transition was mediated by the accumulation in the tank of molecules/pheromones secreted by the leeches. We positioned 20 leeches in the tank and after the formation of a large group of more than 13 conspecifics, leeches were removed from the tank and – leaving the same water – replaced by five naïve leeches. These naïve leeches did not form a group, even though they were sensing the same local chemical medium where previously a large group of conspecifics was formed. Therefore, we posit that group formation is not likely triggered by chemicals secreted into the bath.

STATISTICS OF TRANSITIONS BETWEEN DIFFERENT GROUPS

We have also analyzed the frequency of transitions $p_{ij}$ between groups of different sizes. The most frequent transitions were those from groups of $n$ conspecifics to groups of $n+1$ or $n-1$ conspecifics, i.e., merges and splits usually involved one animal leaving or joining a group. This pattern was clearly observed in the presence of 10 (Figure 3B, estimates over 10 experiments) and – to a lesser extent – of five conspecifics (Figure 3A, estimates over 7 experiments). The lifetimes distribution of groups of size $n$ when 5 (Figure 3C) and 10 leeches (Figure 3D) were in the observation tank could be fitted by the equation

$$\alpha_n / (\beta_n + \Delta t) \gamma_n$$

(Equation 14)
With $\alpha_n$, $\beta_n$, and $\gamma_n$ as free parameters. The value of $\beta_n$ controls the average lifetime of a group of $n$ conspecifics (Figure 3F): it has a high value for long lasting groups and varies from 2 to 16 s. The exponent $\gamma_n$ in Eq. 14 determines how stable a group of $n$ conspecifics (Figure 3G) is: when 10 leeches were in the tank, the value of $\gamma_n$ was close to 1.8 for groups composed by three to five conspecifics, otherwise it was close to 1. Groups of three to four conspecifics had a shorter mean lifetime, indicating that these groups are unstable and either split in smaller groups or merged with other leeches. Groups of more than five leeches are more stable because there is a larger possibility that individuals of the group decide to leave it or split it. In leeches, however, in order to explain the experimental data we need to suppose that the split probability depends on the group size.

In order to understand the mechanisms leading to the formation of a large group of conspecifics we analyzed in detail the events preceding it. The average number of moving leeches (m.l.) $<N_{\text{moving}}>$ decreased from an initial value around 0.7 and 1.4 m.l./s for groups of 5 and 10 leeches respectively (Figures 4A,D, average values over 15 and 17 experiments, respectively) to less than 0.2 m.l./s after 3 h. However, after 2–3 h, five leeches in the tank moved more often than 10 leeches (compare Figures 4A,D) and continued to merge and split (Figures 4B,E).

During the initial exploratory phase, encounters among 10 leeches are more frequent: indeed, the average number of contacts $<C(i\Delta t)>$ (Figures 4C,F) was consistently higher with 10 leeches than with 5, because encounters among conspecifics are easier in a more crowded environment. In the presence of 10 leeches, the number of merges–splits oscillated during the first hour, but had large positive peaks in a time window of 10–30 min (asterisks in Figure 4E) preceding the formation of a stable group (G letters in Figure 4E). The results of Figure 4 indicate two features of group formation: (1) – leeches explore for 2–3 h the new environment before coming to rest and (2) – if during the exploratory phase leeches encounter enough conspecifics a reinforcement dynamics is triggered and leeches stop and rest not in isolation but in groups.
THE ROLE OF 5-HT IN GROUP FORMATION
In order to understand the biological basis of group formation, we asked which neurotransmitters could be involved. An elevation of the body concentration of 5-HT shifts locusts from a solitary to a social behavior (Anstey et al., 2009). We isolated from solitary leeches their entire spinal cord from the tail to the head ganglion and we took a sample of 100–200 μl of their blood and we repeated the same dissection from gregarious leeches, which had formed a group of at least eight conspecifics. We measured the concentration of 5-HT with High Pressure Liquid Chromatography: the concentration of 5-HT in ganglia from the spinal cord of gregarious leeches increased by 37% (with a significance of 0.0493; one tail t-test. N = 5 solitary leeches, six gregarious leeches) over that measured in solitary leeches. Also the level of 5-HT in the blood of gregarious leeches was higher but at a lower extent: indeed the level of 5-HT in the blood of gregarious leeches was 25% higher than in solitary leeches (with a significance of 0.04; one tail t-test. N = 4 solitary leeches, 7 gregarious leeches). These experiments were performed from May to September when leeches are more gregarious (see Figure 8). We repeated the same experiments in winter when leeches are less gregarious and do not form easily groups (see Figure 8) and in this period of the year changes of level of 5-HT between gregarious and solitary leeches could not be determined with an adequate statistical significance.

Therefore we studied group formation when five leeches were injected with 300 μl of 200 μM 5-HT solution. In this case (Figure 5A, gray bars: 15 experiments involving five not injected leeches, black bars: 12 experiments involving five injected leeches), groups of three or four conspecifics occur more frequently than what is expected from an independent behavior (red bars). The difference between merges and splits, (Figure 5B and to be compared with Figure 4) approached values close to 0, with positive peaks (asterisks) preceding the formation of a group (G). The average number of contacts <Ci(Δt)> for five leeches injected with 5-HT was higher than for the case of five uninjected leeches (Figure 5C). When an isolated leech was injected with 300 μl of 200 μM 5-HT and was moved to a tank where a large group of conspecifics had formed, the injected leech quickly joined the large group (see red trajectory in Figure 5D). The results of these experiments did not have the seasonal variation of spontaneous group formation described in Figure 8.

There are many types of 5-HT receptors and several of them have been identified in the leech nervous system: in leeches, 5-HT can activate cationic channels (Sanchez-Armass et al., 1991; Catarsi and Drapeau, 1997; Burrell et al., 2001), anionic channels (Sanchez-Armass et al., 1991; Ali et al., 1998; Burrell et al., 2001), and affect its own reuptake (Bruns et al., 1993; Calviño et al., 2005; De-Miguel and Trueta, 2005). The role of these distinct

![Figure 5](https://www.frontiersin.org)
actions of 5-HT can be dissected by using antagonists such as ketanserin blocking cationic 5-HT gated ionic channels 5-HT_2A and 5-HT_2C (Sanchez-Armass et al., 1991; Mar and Drapeau, 1996) or mianserin blocking 5-HT gated ionic channels (Gaudry and Kristan, 2009).

In contrast to what observed when a leech is injected with 5-HT (Figure 5D), leeches injected with 150 μl of 1 mM ketanserin (Figure 6A) or of 150 μl of 2 mM mianserin (Figure 6B) and subsequently moved to a tank where a large group of conspecifics had formed, remained solitary and rarely joined the large group. This behavior was also observed when the injection of the antagonist was followed by a subsequent injection of 5-HT (Figure 6C, eight experiments under saline conditions, seven under ketanserin, six under mianserin, eight under 5-HT). Leeches injected with a 5-HT antagonist were picked from the water tank where either they were solitary or formed large groups of conspecifics. Their behavior following the injection of 5-HT antagonist was not dependent whether they were solitary or gregarious in the water tank. Leeches injected with a saline solution (cyan line in Figure 6D) remained solitary, as well as leeches injected with both 5-HT and mianserin or ketanserin (red and green lines, respectively), while those injected with 5-HT joined large groups (blue trajectory). Not injected leeches remained solitary, presumably because they could not encounter their conspecifics during the exploratory phase. The simultaneous injection of ketanserin and mianserin into leeches is expected to have a larger effect of the injection of a single 5-HT antagonist.

**NEUROBIOLOGICAL MECHANISMS UNDERLYING GROUP FORMATION**

Leeches are often observed exploring other conspecifics’ bodies by flaring the mouth over their skin, chemoreception could be at basis of group formation. Chemoreception in leeches is mediated by sensilla, ciliated, button-like structures lining the edge of the dorsal lip (Dickinson and Lent, 1984; Elliott, 1984, 1986; Lent and Dickinson, 1984; Gaudry et al., 2010). There are two classes of sensilla: structures with a size of 25–30 μm readily detectable with a light microscope as unpigmented spots in the skin, and smaller structures with a size of 8–10 μm flanking the larger organs not easily detectable with a light microscope (Figure 7A, green dots). The chemical signals detected by sensilla are conveyed to the supraesophageal and subesophageal ganglion. These two ganglia form the head brain of the leech and are connected to the chain of 21 ganglia constituting the central nervous system of the leech (Figure 7A, blue structure).

Therefore we investigated the effect of the surgical removal of the dorsal lip sensilla visible in the light microscope (Figure 7B, red dashed line enclosing green dots). Sham ablations were done in the same way and a band of the dorsal skin adjacent to the lip organs was removed while leaving sensilla intact (Figure 7B, red dashed line not enclosing green dots). After ablation, leeches

![FIGURE 6](image)

**FIGURE 6** | The effect of 5-HT antagonists. (A, B) Black shapes indicate large groups of nine and seven leeches, respectively and the leeches profile (black blob with green circle) injected with 150 μl of 1 mM ketanserin (A) or 2 mM mianserin (B) solutions. Trajectories of injected leeches in red. In both panels, the initial and final positions of the injected leech are marked by the green and the blue dot respectively. Injection of 5-HT antagonists was followed by another injection of 150 μM 5-HT solution. (C) Percentage of time spent by injected leeches in contact with the large group of conspecifics, when leeches were injected with 300 μl of saline solution, with 150 μl of 1 mM ketanserin or 2 mM mianserin solutions, followed by another injection of 150 μl of 400 μM 5-HT solution, or 300 μl of 200 μM 5-HT solution. (D) Time evolution of the mean size of the group joined by the injected leech under the same treatments described in (C) cyan, blue, red, and green lines refer to leeches injected with a saline solution, 5-HT, mianserin, and ketanserin respectively. Student’s t-test, *p < 0.05.
were allowed to recover for 1 day in water and exhibited a normal behavior with frequent episodes of swimming and crawling. After recovery from surgery, ablated, and sham-ablated leeches, injected with 300 μl of 200 μM 5-HT were tested. Ablated leeches, either 5-HT injected (N = 5) or not (N = 7), remained solitary (Figures 7C,D, see legend, Student’s t-test, *p < 0.05.), while not ablated (N = 8) and sham-ablated 5-HT injected leeches (N = 6) joined the other leeches (Figures 7C,D, see legend Student’s t-test, p < 0.05.), indicating that group formation depended on the presence of intact sensilla.

It is possible to disconnect the supraesophageal ganglion from the leech central nervous system by cutting the neuronal projections from the supraesophageal to the subesophageal ganglion (Figure 7B, red vertical line). Leeches with a disconnected supraesophageal ganglion recovered from surgery and 1 day later they swam and crawled as intact leeches do. When these leeches were injected with 5-HT, they remained solitary and did not join and mix with other leeches and behaved as intact leeches not injected with 5-HT (Student’s t-test, p < 0.05, N = 3).

SEASONAL VARIABILITY

Deviations from an independent behavior and the formation of large groups of conspecifics were observed both in well-fed and hungry leeches. However, a careful examination of data collected over more than 2 years of experiments showed a seasonal variability; leeches received from the supplier (Ricarimpex, see Materials and Methods) in late spring or summer were more active and prone to social aggregation, while those received in winter did not move very much and appeared to be less “sociable” and more solitary. Leeches supplied by Ricarimpex were kept outside and therefore were exposed to seasonal changes. We measured several properties of group formation mediating data over a time window of 3 months. In all our experiments the water temperature did not change more than 2˚C, from winter to summer, and its average value was about 16˚C.

The tendency to group formation was quantified by computing the average group size (Figure 8A), the average number of contacts established within 4 h (Figure 8B), the average velocity during the initial exploratory phase (Figure 8C) and the average number of moving leeches per minute (Figure 8D) when 10 leeches were in the observation tank (N = 62 experiments uniformly performed over 2 years). By using all four criteria, we observed a recurrent dependence between group formation and the season. Leeches were more prone to form large groups between May and September, corresponding to warm weather in continental Europe. During European winter, leeches were less active and more solitary. These results show that the transition from solitary to social behavior is a complex process, likely to depend on the overall state.
of the animals, also influenced by seasonal variations and possibly by the length of daylight.

DISCUSSION

Group formation is an important aspect of social interactions (Sokolowski, 2010), occurring when conspecifics make collective decisions and adopt a similar behavior. Group formation can happen for different ethological reasons, but its dynamics – as we show in this study – can be similar in distant species such as leeches and humans (Stehlé et al., 2010; Zhao et al., 2011), when the population of leeches exceeds a critical value, they aggregate in large groups. Similar conspecifics aggregation has been observed in simple model systems, such as the worm C. elegans the fruit fly Drosophila (Sokolowski, 2010), in fish shoals (Makris et al., 2009), in locust swarms (Anstey et al., 2009) but also in higher primates and humans (Goodson and Thompson, 2010). Group formation in several species is not primarily caused by a mutual attraction among conspecifics but is produced by aggregation around abiotic factors, – in worms, for instance – as a response to oxygen levels (De Bono and Bargmann, 1998). Leeches are negative phototactic (Elliott, 1973; Sawyer, 1986b) and they tend to move away from a light source, therefore the observed group formation could have been simply their way to escape from the light and find a shelter among other leeches. When five or less leeches were placed in the observation tank of a given area they hardly noticed each other and their motion and behavior followed the pattern indicating independence with a high statistical significance (Bisson and Torre, 2011). When a higher number of leeches were placed in the same tank, a different dynamics was observed, difficult to reconcile with negative phototaxis. Similarly, group formation observed in a smaller number of leeches injected with 5-HT strongly indicated the emergence of social interactions, i.e., reciprocal attraction among conspecifics.

Similarly to what observed in locusts (Anstey et al., 2009), the hormone 5-HT participates in the onset and control of group formation. 5-HT plays a role similar to oxytocin, arginin, and vasopressin. These non-peptides mediate social interactions and behavior in several species from invertebrates (Donaldson and Young, 2008) to rodents (Insel, 2010) and possibly also in humans (Goodson and Thompson, 2010). In humans, 5-HT is involved in mood regulation and its disruption is at the basis of several kinds of mood disorders (Chiao, 2010). The segmental ganglia of the leech contain a network of 5-HT releasing neurons (Lent and Frazer, 1977; De-Miguel and Trueta, 2005) modulating the swimming motor program (Ort et al., 1974; Kristan and Nusbaum, 1982; Friesen, 1989). These neurons are one pair of large Retzius cells, one pair of dorso-lateral (DL) and ventro-lateral (VL) interneurons, and two pairs of medial interneurons (cells E and M; Lent et al., 1991). 5-HT is known to influence social behavior and in particular social ranking in lobsters (Kravitz, 1988) and in crayfish (Teh et al., 1996).

NEUROBIOLOGICAL MECHANISMS FOR GROUP FORMATION

Sensilla play a strategic role in chemoreception and several associated behaviors such as feeding (Dickinson and Lent, 1984; Elliott, 1984, 1986; Lent and Dickinson, 1984; Gaudry et al., 2010). The presence of conspecifics is detected by chemoreceptors located in the sensilla present in the dorsal lip and the associated neurobiological signal is transmitted through the supraesophageal ganglion...
FIGURE 9 | Modeling group formation in leeches and comparison with experimental data. (A) Time evolution of number of groups from a simulation of Eqs 3–5 with five leeches and using the parameters $K_{\text{met}} = 30; c_1 = 1; c_2 = 0.3; a_1 = 1, a_2 = 1, a_3 = 0.8, a_4 = 0.8, a_5 = 0.4, a_n = 0.3$, for $n > 5; b_n = 10a_n$, for $n > 0$. (B) Time evolution of number of groups in three different experiments with five leeches. (C) Time evolution of the average group size from a simulation of Eqs 3–5 with five leeches. (D) Time evolution of the average group size in three different experiments with five leeches. (E–H) As in (A–D), but with 10 leeches. (I,J) Comparison of the lifetime distributions that result from experimental data (blue line) and simulated data (magenta line) for groups of size 1 (I) and 5 (J) in experiments with five leeches. (K,L) As in I–J but for groups of size 1 (K) and 10 (L) in experiments involving 10 leeches.
to the central nervous system of the leech, where it promotes the increase of the intracellular level of 5-HT. Ablation experiments indicate a specific neurobiological pathway mediating the establishment of social interactions. Impairment of the dorsal lip or of the supraesophageal ganglion compromises conspecifics detection, as in fake group experiments (Bisson and Torre, 2011) and abolishes social attitude. There are many types of 5-HT receptors and several of them have been identified in the leech nervous system, where 5-HT can activate cationic channels, anionic channels, and affect its own reuptake (Calviño et al., 2005). Ketanserin blocking cationic 5-HT gated ionic channels 5-HT_{2A} and 5-HT_{2C} and mianserin blocking 5-HT gated ionic channels make leeches more solitary (Figure 6) suggesting 5-HT could promote group formation by activating the whole population of 5-HT receptors. As the body concentration of 5-HT in ganglia of gregarious leeches is by 37% larger than in solitary leeches we propose that social interactions occur because of an elevation of the body level of 5-HT.

Supraesophageal leech ganglia, similarly to all leech ganglia, have two large Retzius cells able to secrete 5-HT both from the soma and synaptic terminals (Sawyer, 1986a). Retzius neurons are the major serotoninergic neurons in the leech central nervous system and indeed contain approximately half of the total concentration of 5-HT (Coggeshall et al., 1972). We speculate, therefore, that 5-HT released extrasynaptically from the soma of Retzius cells in the supraesophageal ganglion diffuse into the leech body and is the source of the measured elevation of 5-HT. 5-HT diffusing out of the release site will have a variety of actions, such as autoinhibition of 5-HT releasing Retzius cells (Cercós et al., 2009) and may also produce slow electrical responses, as happens in dopaminergic neurons (Sombers, 2009) and ultimately triggers group formation. The time course of group formation occurs in 1 h or so and therefore acts on a time scale more compatible with an extrasynaptic release of 5-HT (Bunin et al., 1998; Bunin and Wightman, 1999; De-Miguel and Trueta, 2005; Kaushalya et al., 2008).

**MODELING GROUP FORMATION**

During the analysis of leech aggregation, we observed that the lifetimes distribution of leech groups (Eq. 14 and Figures 3C,D) were very similar to those seen among humans (Stehlé et al., 2010; Zhao et al., 2011). Therefore, we decided to model group formation among leeches, on the basis of the reinforcement dynamics used to describe human social interactions (Stehlé et al., 2010; Zhao et al., 2011). We assumed that leech aggregation (Figure 3B) is determined primarily by transitions of the type

\[
\begin{align*}
\text{n} & \to \text{n} + 1 \quad \text{(a merge)} \\
\text{n} & \to \text{n} + 1 \quad \text{(a split)}
\end{align*}
\]

in which a single conspecific leaves or joins another group. As suggested by Figures 3C,D, the probability \(D_n(\Delta t)\) that an animal makes a transition and leaves a group of \(n\) conspecifics which has persisted for a time \(\Delta t\) is

\[
D_n(\Delta t) = \frac{a_n}{\Delta t + b_n}.
\]

This is the simplest and minimal hypothesis able to explain the distribution of lifetimes of Eq. 14. Eq. 14 indicates the existence of a reinforcement dynamics so that the longer a group lasts the less likely it is to split. The constants \(a_n\) and \(b_n\) set the temporal scale of this reinforcement dynamics and the limit of this probability for \(\Delta t \to 0\).

This transition will be a merge with another random group with probability \(D_n(\Delta t)\) or a split with probability \(D_n(\Delta t)(1 - \lambda)\). In agreement with the presumed role of 5-HT (Figures 5 and 6) merges are promoted by the intra body concentration of 5-HT \(s\), as

\[
\lambda = \frac{s}{s + K_{\text{met}}}. \tag{16}
\]

This expression assumes that the probability to merge with another group is an activated process modulated by the parameter \(\lambda\) reminiscent of Michaelis–Menten kinetics. \(K_{\text{met}}\) sets the typical scale for \(s\) at which the activation process takes place. We assumed also that \(s\) increases when leeches encounter their conspecifics and that \(s\) is subsequently degraded. These assumptions are modeled as:

\[
\frac{ds}{dt} = c_1 N_{\text{contacts}}(t) - c_2 s.
\]

\[
N_{\text{contacts}} = \sum_{i=1}^{G} n_i^2
\]

Where \(c_1\) is the rate at which 5-HT concentration increases for each group contact and \(c_2\) is the degradation constant.

From Eqs 3–5 and the parameter values reported in the legend we simulated the time evolution of the number of groups in the presence of 5 (Figure 9A) and 10 leeches (Figure 9C): the formation of a large group of conspecifics is observed for 10 leeches and not for 5 as experimentally observed (Figures 9B,D). Also the simulated time course of group size (Figures 9E,G) formation is in agreement with what experimentally observed (Figures 9F,H). The simple model reproduces qualitatively also the lifetime distribution of groups of conspecifics (Figures 9I–L). The essence of the reinforcement dynamics, leading to the formation of a large group with 10 but not with 5 leeches is contained in Eqs 4–5: the asymptotic rate of merges \(\lambda\) increases with the number of conspecifics, because the number of contacts is higher with a larger number of conspecifics (Figure 4).

These results indicate that group formation is caused by a reinforcement dynamics that can be modeled by similar equations in different species, such as leeches and humans. The neuromodulator 5-HT promotes group formation in leeches and its involvement in interpersonal relationships among humans is well-established (Goodson and Thompson, 2010). Therefore group formation, important aspect of social interactions in leeches and in humans, share common dynamical properties and specific neuromodulators.

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