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# Optic Flow Based Visual Guidance: From Flying Insects to Miniature Aerial Vehicles 

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## 1. Introduction

Insects and birds have been in the sensory-motor control business for more than 100 million years. The manned aircraft developed over the last 100 years rely on a similar lift to that generated by birds' wings. Aircraft designers have paid little attention, however, to the pilot's visual sensor that finely controls these wings, although it is definitely the most sophisticated avionic sensors ever known to exist. For this reason, the thinking that prevails in the field of aeronautics does not help us much grasp the visuo-motor contol laws that animals and humans bring into play to control their flight. To control an aircraft, it has been deemed essential to measure state variables such as barometric altitude, groundheight, groundspeed, descent speed, etc. Yet the sensors developed for this purpose - usually emissive sensors such as Doppler radars, radar-altimeters or forward-looking infrared sensors, in particular - are far too cumbersome for insects or even birds to carry and to power. Natural flyers must therefore have developed other systems for controlling their flight.
Flying insects are agile creatures that navigate swiftly through most unpredictable environments. Equipped with "only" about one million neurons and only 3000 pixels in each eye, the housefly, for example, achieves 3D navigation at an impressive 700 bodylengths per second. The lightness of the processing system at work onboard a fly makes us turn pale when we realize that this creature actually achieves just what is being sought for in the field of aerial robotics: dynamic stabilization, 3D autonomous navigation, ground avoidance, collision avoidance with stationary and nonstationary obstacles, tracking, docking, autonomous takeoff and landing, etc. Houseflies add insult to injury by being able to land gracefully on ceilings.
The last seven decades have provided evidence that flying insects guide themselves through their environments by processing the optic flow (OF) that is generated on their eyes as a consequence of their locomotion. In the animal's reference frame, the translational OF is the angular speed $\omega$ at which contrasting objects in the environment move past the animal (Kennedy, 1939; Gibson, 1950; Lee, 1980; Koenderink, 1986).
In the present chapter, it is proposed to summarize our attempts to model the visuomotor control system that provides flying insects with a means of autonomous guidance at close range. The aim of these studies was not (yet) to produce a detailed neural circuit but rather to obtain a more functional overall picture, that is, a picture that abstracts some basic control
principles (Marr, 1982). We attempted to determine the variables the insect really measures (see also: Taylor \& Krapp, 2008), the actions it takes to control its flight and the causal and dynamic relationships between the sensory and motor variables involved.
Our progress on these lines was achieved by performing simulation experiments and testing our control schemes onboard miniature aircraft. Like our early terrestrial "robot-mouche" (robot Fly) (Pichon et al., 1989; Franceschini et al., 1992), these aerial robots are based on the use of an electronic OF sensor (Blanes, 1986) inspired by the housefly Elementary Motion Detectors (EMD), which we previously studied in our laboratory (Rev.: Franceschini, 1985; Franceschini et al., 1989).


Figure 1. Head of the blowfly Calliphora erythrocephala with its panoramic compound eyes (left). The right part shows about $2 \%$ of the housefly retinal mosaic. A cluster of micrometersized photoreceptors is located in the focal plane of each facet lens : 6 outer receptors R1-6 surround a central cell R7 (prolonged by an R8, not seen here). These are the natural autofluorescence colors of the receptors observed in vivo under blue excitation (Franceschini et al., 1981a,b), after « optical neutralization of the cornea » (From Franceschini, 2007)
Section 2 recalls some aspects of the fly visual system. Section 3 focuses on the realisation of fly-inspired OF sensors. Section 4 examines the problem of ground avoidance. Section 5 introduces the OF regulator. In Section 6 the miniature helicopter we built is described, equipped with an electronic $O F$ sensor and an $O F$ regulator. The extent to which the $O F$ regulator accounts for the actual behavioral patterns observed in insects is discussed in Section 7. Section 8 introduces the dual OF regulator that is able to control both speed and distance to the walls in a corridor. In Section 9, another visuomotor control system is presented which enables an aerial robot to stabilize in yaw by visually fixating a target with hyperacuity, on the basis of a principle inspired by the fly. Section 10 ends up by discussing the potential applications of these insect-derived principles to the navigation of aerial vehicles - in particular Micro Aerial Vehicles (MAVs) and Micro Space Vehicles (MSVs).

## 2. The fly visual system and its motion sensitive neurons

Each compound eye consists of an array of ommatidia. The frontend of each ommatidium is a facet lens that focusses light on a small group of photoreceptor cells (Fig. 1, right). The fly retina is among the most complex and best organized retinal mosaic in the animal kingdom. It has been described in great details, with its different spectral types of photoreceptor cells, polarization sensitive cells and sexually dimorphic cells. There exists a typical division of labour within the retina (Rev.: Franceschini, 1984; Hardie, 1985):

- The two central photoreceptor cells, R7-8, display various spectral sensitivities that are randomly scattered across the retinal mosaic, as attested by the various R7 autofluorescence colors (Fig. 1, right). R7 and R8 are thought to participate in color vision.
- The outer 6 photoreceptor cells (R1-R6) all have the same, wide-band spectral sensitivity. They participate, in particular, in motion detection (Rev.: Buchner, 1984, Heisenberg \& Wolf, 1984; Riehle \& Franceschini, 1984 ). In this visual pathway, signal-to-noise ratio is improved by the presence of an ultraviolet sensitizing pigment that enhances the quantum catch (Kirschfeld \& al., 1977), and by an exquisite opto-neural projection called "neural superposition" (Braitenberg, 1967; Kirschfeld, 1967; Kirschfeld and Franceschini, 1968). The R1-R6 photoreceptors therefore make for a high sensitivity ("scotopic") system (Kirschfeld and Franceschini, 1968).
To estimate the OF, insects use motion sensitive neurons. In flies, part of the 3rd optic ganglion called the Lobula Plate (LP) appears as a genuine "visual motion processing center". It comprises approximately 60 uniquely identifiable neurons, the LP tangential cells (LPTC) that analyze the OF field resulting from the animal's walking or flying. Some of these neurons transmit their electrical signals via the neck to thoracic interneurons that will drive the wing-, leg-, and head-muscles. Other neurons (in particular H1, see Fig. 2b) send their signals to the contralateral eye. The LPTCs are actually large-field collator neurons that pool the electrical signals from many retinotopic input elements called 'Elementary Motion Detectors' (EMDs) (Rev.: Hausen \& Egelhaaf, 1989; Egelhaaf \& Borst, 1993; Hausen, 1993; Krapp et al., 1998; Borst \& Haag, 2002, Taylor \& Krapp, 2008). The cellular details underlying an EMD has not been fully identified in any insects - nor in any vertebrates.
Taking advantage of the micro-optical techniques we had developed earlier (Rev.: Franceschini, 1975), we were able, however, to activate a single EMD in the eye of the living housefly by stimulating single identified photoreceptor cells within a single ommatidium while recording the response of an identified motion sensitive neuron (H1) in the LP (Riehle \& Franceschini, 1984, Franceschini, 1985, 1992 ; Franceschini et al., 1989). We applied pinpoint stimulation to two neighboring photoreceptors (diameter $\cong 1 \mu \mathrm{~m}$ ) within the selected ommatidium (Fig.2a) by means of an optical stimulation instrument (Fig. 2d) whose main objective was quite simply the facet lens itself (diameter $\cong 25 \mu \mathrm{~m}$, focal length $\cong 50$ $\mu \mathrm{m}$ ). This exact instrument (a rotating triple beam incident light 'microscope-telescope') served to : (i) select a facet lens, (ii) select 2 out of the 7 receptors (R1 and R6), and (iii) stimulate them successively with $1 \mu \mathrm{~m}$-light spots. Sequential stimulation produced an 'apparent motion' that would simulate a real motion within the small visual field of the selected ommatidium. The H1-neuron responded by a conspicuous increase in spike rate, as long as the phase relationship between the two stimuli mimicked a movement occurring in the preferred direction (see Fig. 2c, top trace). The null response observed for the reverse sequence (Fig 2c, bottom trace) attests to the remarkable sequence discriminating ability of an EMD, which is sensitive to directional motion (Franceschini et al., 1989, Franceschini, 1992).


Figure 2. (left) : (a-c) Principle of the experiment aimed at deciphering the principle of motion vision in flies, using optical stimulation of single photoreceptors. (d) Triple-beam incident light "microscope-telescope" that delivers a $1 \mu \mathrm{~m}$ light spot to two neighboring photoreceptor cells, R1 and R6, successively (see (a)). A microelectrode (c) records the electrical response (nerve impulses) of the motion sensitive neuron H1 to this "apparent motion" (From Franceschini et al., 1989)
From many experiments of this kind, in which various sequences of light steps and/or pulses were applied to selected receptor pairs, we established an EMD block diagram and characterized each block's dynamics and nonlinearity (Franceschini, 1985, 1992; Franceschini et al., 1989). While not unveiling the cellular details of the EMD circuit, our analysis allowed the EMD principle to be understood functionally, paving the way for its transcription into another, man-made technology.

## 3. From biological to electronic optic flow sensors

In the mid 1980's, we designed a neuromorphic optic flow sensor (Blanes, 1986, 1991, Franceschini et al., 1986), the signal processing scheme of which was inspired by what we had learned from the fly EMD. The OF is an angular speed $\omega$ that corresponds to the inverse of the time $\Delta t$ taken by a contrasting feature to travel between the visual axes of two adjacent photoreceptors, separated by an angle $\Delta \varphi$. Our OF sensor processes this delay $\Delta \mathrm{t}$ so as to generate a response $\omega_{\text {meas }}$ that grows monotonically with the inverse of $\Delta t$, and hence with the optic flow $\omega=\Delta \varphi / \Delta \mathrm{t}$ (Fig. 3a). Short delays $\Delta \mathrm{t}$ give higher voltage outputs and vice versa.


Figure 3. (a) Principle of the optic flow sensor derived from our electrophysiological analyses of the housefly's EMD (Blanes, 1986, Franceschini et al., 1986). (b) purely analogue version (weight 5 grams) built in 1989 for the Robot-Fly, whose compound eye housed a ring of 114 EMDs of this type (Pichon et al., 1989 ; Blanes, 1991 ; Franceschini \& al., 1992) (c) hybrid (analogue + digital) version (size : $7 \mathrm{~mm} \times 7 \mathrm{~mm}$, mass 0.2 grams) based on a microcontroller and built using Low Temperature Co-fired Ceramics technology (LTCC) (Pudas et al., 2007)
Our scheme is not a "correlator scheme" (cf Hassenstein \& Reichardt, 1956, Reichardt, 1969) and corresponds to the class of "feature-matching schemes"(Ullman, 1981), where a given feature (here a change in intensity that may represent a moving edge) is extracted and tracked in time. The photodiode signal of each channel is first bandpass filtered (Fig. 3a) - resembling the analog signal measured in the large monopolar neurons of the fly lamina (Laughlin, 1984). The next step consists of hysteresis thresholding and generation of a unit pulse. In the EMD version built in 1989 for the Robot-Fly (Fig. 3b), the unit pulse from one channel was sampling a long-lived decaying exponential function generated by the other channel (Blanes, 1991), via a nonlinear circuit called a minimum detector, to give an output $\omega_{\text {meas }}$ that is virtually equal to the OF, $\omega=\Delta \varphi / \Delta \mathrm{t}$ (Fig. 3a). The thresholding operation makes the voltage output virtually invariant to texture and contrast and the circuit responds as well to natural scenes (Portelli et al., 2008). A very similar EMD principle has been conceived, independently, a decade later by C. Koch's group at CALTECH, where it became known as the "facilitate and sample" velocity sensor
(Kramer et al., 1995). Yet another variant of our original "time of travel" principle was proposed another decade later (Moeckel \&Liu, 2006).
Since our original analog design (Blanes, 1986, 1991; Franceschini et al, 1986), we have built various versions of OF sensors based on this very principle. In the EMD currently used onboard our aerial robots, the signals are processed using a mixed (analog + digital) approach (Ruffier et al., 2003). Such OF sensors can be small and lightweight (the smallest one weighs only 0.2 grams: Fig. 3c) to mount onto MAVs. Several OF sensors of this type were also recently integrated on a miniature FPGA (Aubépart et al. 2004, 2008; Aubépart \& Franceschini, 2007). A different kind of OF sensor was designed recently (Barrows et al., 2001) and mounted on a model plane (Barrows et al, 2003; Green \& Barrows, 2004).

## 4. The problem of ground avoidance



Figure 4. Definition (a) and measurement of the ventral optic flow $\omega$ experienced by an insect (or a robot) flying in translation in the vertical plane. (b) an EMD of the type shown in Fig. 3, like the EMDs driving some insects' neurons (Kien, 1975; Olberg, 1981; Ibbotson, 2001), is able to measure the ventral OF, i.e., the angular speed $\omega$ at which a contrasting feature moves under the flying agent (From Franceschini et al., 2007)
The ventral OF experienced in the vertical plane by flying creatures - including aircraft pilots - is the apparent angular velocity $\omega$ generated by a point directly below on the flight track (Gibson et al., 1955; Whiteside \& Samuel, 1970). As shown in figure 4a, the ventral OF depends on both the groundspeed $V_{x}$ and the groundheight $h$ and is equal to the ratio between these two variables:

$$
\begin{equation*}
\omega=\mathbf{V}_{\mathrm{x}} / \mathrm{h}\left[\mathrm{rad} . \mathrm{s}^{-1}\right] \tag{1}
\end{equation*}
$$

We know that flies and bees are able to react to the translational $O F$ independently of the spatial texture and contrast (David, 1982; Kirchner \& Srinivasan, 1989, Srinivasan et al., 1991, 1993; Baird et al., 2005). We also know that some insects' visual neurons may be involved in this reaction because they respond monotonically to $\omega$ with little dependence on texture and contrast (Kien, 1975; Olberg, 1981; Ibbotson, 2001; Shoemaker et al., 2005). Neurons facing downwards can therefore act as ventral OF sensors, and thus assess the $V_{x} / h$ ratio (figure 4). Based on laboratory experiments on mosquitoes and field experiments on locusts, Kennedy put forward an "optomotor theory" of insect flight, according to which flying insects
maintain a "preferred retinal velocity" with respect to the ground below (Kennedy, 1939, 1951). In response to wind, for example, insects may adjust their groundspeed or groundheight to restore the apparent velocity of the ground features. Kennedy's hypothesis has been repeatedly confirmed during the last 30 years: both flies and bees were found to maintain a constant OF with respect to the ground while cruising or landing (David, 1978; Preiss, 1992; Srinivasan et al., 1996, 2000, Baird et al., 2006).
The problem is how insects may achieve this feat, since maintaining a given OF is a kind of chicken-and-egg problem, as illustrated by Eq.1: an insect may hold its ventral OF, $\omega$, constant by adjusting either its groundspeed (if it knows its groundheight) or its groundheight (if it knows its groundspeed). Besides, the insect could maintain an OF of $1 \mathrm{rad} / \mathrm{s}$ (i.e., $57^{\circ} / \mathrm{s}$ ), for instance, by flying at a speed of $1 \mathrm{~m} / \mathrm{s}$ at a height of 1 meter or by flying at a speed of $2 \mathrm{~m} / \mathrm{s}$ at a height of 2 m : there is an infinitely large number of possible combinations of groundspeed and groundheight that will give rise to the same "preferred $\mathrm{OF}^{\prime}$.
Drawing on the experience we had with OF-based visual navigation of a terrestrial robot (Pichon et al., 1989; Franceschini et al., 1992), we attempted early to develop an explicit flight control scheme for aerial navigation in the vertical plane. Our first tentative step on these lines was not particularly successful, because we were cornered by the general notion that prevailed in those days that insect navigation relies on gauging range (Kirchner \& Srinivasan, 1989; Srinivasan et al., 1991; Franceschini et al., 1992; Srinivasan, 1993). In the experimental simulations we performed in 1994, for example (Mura \& Franceschini, 1994), we assumed that the insect (or the robot) would know its groundspeed $V_{x}$ (by whatever means), so that by measuring $\omega$ it would be able to gauge the distance $h$ from the ground (Eq.1) and react accordingly to avoid it. Although this procedure may be justified in robotics (see, e.g., Barber et al, 2005; Srinivasan et al., 2006; Garratt \& Chahl, 2008), where groundspeed can be determined via GPS, this makes the way insects operate all the more elusive.
In 1999, we established (in simulation) how a rotorcraft (or an insect) might be able to follow a terrain (Fig. 5a) and land (Fig. 5b) on the sole basis of OF cues, without measuring its groundspeed and groundheight (Netter \& Franceschini, 1999).


Figure 5. (a) Simulation of nap-of-the-earth flight of an helicopter equipped with an eye whose 20 pixels cover a frontal FOV of $75^{\circ}$ in the vertical plane, centered at $40^{\circ}$ below the horizon (initial conditions: height: 5 m , speed $2 \mathrm{~m} / \mathrm{s}$; iteration step 1 s ). (b) Landing was initiated by linearly decreasing the aircraft horizontal speed at every iteration while retaining the same request to maintain a reference OF value (From Netter \& Franceschini, 1999)
The landing manoeuver was achieved under permanent visual feedback from a 20 -pixel forward looking eye (with 19 EMDs). The driving force causing the progressive loss in altitude was the decrease in the horizontal flight speed, which occurred when the rotorcraft (or the insect) was about to land - either volontarily or because of a head wind. The landing trajectory obtained in this simulation (Fig. 5b) already resembles the final approach of bees landing on a
flat surface (Srinivasan et al, 1996). The principle was first validated onboard FANIA, a miniature tethered helicopter having a single (variable pitch) rotor, an accelerometer and a forward looking eye with 20 pixels (and, therefore, 19 EMDs) arranged in the frontal meridian (Netter \& Franceschini, 2002). This $0.8-\mathrm{kg}$ rotorcraft had three degrees of freedom (surge, heave and pitch). Mounted at the tip of a flight mill, the robot lifted itself by increasing its rotor collective pitch. Upon remotely inclining the servo-vane located in the propeller wake, the operator made the helicopter pitch forward by a few degrees so that it gained speed and, therefore, climbed to maintain a reference OF with respect to the terrain below. FANIA jumped over contrasting obstacles by increasing its collective pitch as a function of the fused signals from its 19 EMDs (see Fig. 8 in Netter \& Franceschini, 2002).

## 5. The "optic flow regulator"

In spite of this early success to explain how an insect could navigate on an OF basis, we considered that Kennedy's insightful "optomotor theory" was calling for a clear formalization that would bring to light:

- the flight variables really involved
- the sensors really required
- the dynamics of the various system components
- the causal and dynamic links existing between the sensory output(s) and the variable(s) to be controlled
- the points of application of the various disturbances that insects may experience
- the variables insects have to control to compensate for these disturbances.

We came up with an autopilot called OCTAVE (OCTAVE stands for Optical altitude Control sysTem for Autonomous VEhicles) that is little demanding in terms of neural (or electronic) implementation and could be just as appropriate for insects as it would be for aircraft (Ruffier \& Franceschini, 2003; Franceschini et al., 2003). A ventral OF sensor was integrated into a feedback loop that would drive the robot's lift, and thus the groundheight, so as to compensate for any deviations of the OF sensor's output from a given set point (Ruffier \& Franceschini, 2003, 2004a,b, 2005; Franceschini et al., 2007). This simple autopilot (Fig. 6a) enabled a miniature helicopter to perform challenging tasks such as take-off, terrain following, reacting suitably to wind, and landing (see Section 6).


Figure 6. (a) The OF regulator OCTAVE (bottom red feedback loop) controls the lift $L$, and consequently the groundheight, at all times so as to maintain the ventral optic flow $\omega$ constant and equal to the set point $\omega_{\text {set }}$. (b) Flies and bees, like helicopters, pitch forward to increase their forward thrust, and hence their airspeed. As long as they pitch forward by $\Theta<10^{\circ}$, the lift component L does not incur any major loss (From Franceschini et al., 2007)

The OCTAVE autopilot can be said to be an OF regulator. The word 'regulator' is used here in the strictest acceptation of the word in the context of control theory, where it denotes a feedback control system designed to maintain an output signal constantly equal to a given set point. The Watt flyball governor from the $18^{\text {th }}$ century, for instance, was not only one of the first servomechanisms ever built: it was also the very first angular speed regulator. It served to maintain the rotational speed of a steam engine shaft at a given set point, whatever interferences occurred as the result of unpredictable load disturbances. The Watt regulator was based on a rotational speed sensor (meshed to the output shaft), whereas our OF regulator is based on a noncontact rotational speed sensor - the OF sensor - that measures the ventral OF - again in rad/s.
Specifically, the OF signal $\omega_{\text {meas }}$ delivered by the OF sensor (see Fig. 6a, bottom) is compared with the OF set point, $\omega_{s e t}$. The comparator produces an error signal: $\boldsymbol{\varepsilon}=\omega_{m e a s}-\omega_{s e t}$ which drives a controller adjusting the lift $L$, and thus the groundheight $h$, so as to minimize $\boldsymbol{\varepsilon}$. All the operator does is to set the pitch angle $\Theta$ and therefore the airspeed (see figure 6a): the $O F$ regulator does the rest, that is, it adjusts the groundheight $h$ proportionally to the current groundspeed $V_{x}$, keeping the OF, i.e., the $V_{\sqrt{ }} / h$ ratio constant in the steady state:

$$
\begin{equation*}
\mathrm{V}_{\mathrm{x}} / \mathrm{h}=\omega \cong \omega_{\mathrm{meas}} \cong \omega_{\text {set }}=\mathrm{constant} \tag{2}
\end{equation*}
$$

Upon looking at figure 6 b , one might be tempted to object that controlling $F$ (via the rotor speed) will affect not only $L$ but also $T$, and hence $V_{x}$. This coupling is negligible, however, because the ensuing change in $T$ is much smaller than the change in $L$ (by a factor of at least $5.67=\operatorname{cotan} 10^{\circ}$, because the maximum speed of $3 \mathrm{~m} / \mathrm{s}$ is already attained for $\Theta=10^{\circ}$ ).

## 6. A miniature helicopter equipped with an "optic flow regulator"



Figure 7. (a) Miniature helicopter (MH) equipped with an OF sensor (Fig. 3) and an $O F$ regulator (Fig. 6). The operator sets the forward pitch, and thus the airspeed. The gaze remains automatically oriented downwards. (b) The 100-gram robot lifts itself and circles around CCW at speeds up to $3 \mathrm{~m} / \mathrm{s}$ and heights up to 3 m over the arena, giving rise to the flight patterns given in Fig. 8-10 (From Ruffier \& Franceschini, 2003)
We tested the idea that insects may be equipped with a similar OF regulator by comparing the behavior of insects with that of a 'seeing helicopter' placed in similar situations. The robot we built (figure 7a) is a miniature helicopter (MH) equipped with a simple, 2-pixel
ventral eye driving an EMD acting as an OF sensor (Fig. 3 and 4b). The 100-gram robot is tethered to an instrumented flight mill consisting of a light pantographic arm driven in terms of its elevation and azimuth by the MH's lift and forward thrust, respectively (Fig. $7 b)$.


Figure 8. Flight parameters monitored during a 70-meter flight (consisting of about 6 laps on the test arena: figure 7 b ) performed by the miniature helicopter (MH) (Fig. 7a), equipped with an OF regulator (figure 6a). The complete journey over the randomly textured pattern (shown at the bottom of (a)) includes take-off, level flight and landing
(a) Vertical trajectory in the longitudinal plane. (b) Groundspeed $V_{x}$ monitored throughout the journey. (c) Output $\omega_{\text {meas }}$ of the OF sensor showing the relatively small deviation from the OF set point $\omega_{s e t}$ (red curve). (d) Actual OF, $\omega$ (calculated as $V_{\sqrt{ }} / h$ ) resulting from the behavioral reaction: $\omega$ is seen to have been held relatively - but not perfectly - constant throughout the journey, even during the takeoff and landing maneuvers where the groundspeed varies considerably as shown in (b) (From Franceschini et al., 2007)

Any increase in the rotor speed causes the MH to rise, and the slightest (operator mediated) forward ("nose-down") tilting by a few degrees produces a forward thrust component that causes the MH to gain forward speed. The flight mill is equipped with ground-truth azimuthal and elevation sensors that allow the position and speed of the MH to be monitored at high accuracy and in real time. Since the MH purpose was to demonstrate a basic principle, it was equipped with an elementary ventral eye composed of only two photoreceptors driving a single Elementary Motion Detector (EMD) built according to the "travel time" principle shown in Fig. 3 (Ruffier \& Franceschini, 2003, Ruffier et al., 2003).

## 7. Comparison of robots' and insects' behavioral patterns

The OCTAVE control scheme (Fig. 6) produced the helicopter behavioral patterns shown in figures 8-10. The robot's behavior was found to account for a series of puzzling, seemingly unconnected flying abilities observed by many authors during the last 70 years in various species (fruitflies, honeybees, moths, mosquitoes, dung-beetles, migrating locusts and butterflies, and birds) (Franceschini et al., 2007). Most of the data published in these studies are qualitative, but recent quantitative findings on honeybees' landing performances can also be explained on the basis of this simple control scheme, including the constant descent angle observed in the bee's final approach (Srinivasan et al., 1996, 2000).
Let us now examine these various behavioral patterns, assuming the insect to be equipped with the OF regulator shown in Fig. 6 (Franceschini et al., 2007).

### 7.1 Take-off

An insect that increases its forward thrust by pitching forward (Fig. 6b) like a fly or a helicopter (David, 1978), is bound to rise into the air because the OF regulator (Fig. 6a) will constantly increase the groundheight $h$ proportionally to the current groundspeed $V_{x}$, to maintain the OF constant (Eq.2).
The performances of the MH illustrate this point quite clearly (figure 8a, left part). Starting at position 1 (with the rotor axis oriented vertically and the lift just balancing the weight), the MH is remotely commanded to pitch 'nose-down' ( $\Theta$ from $0^{\circ}$ to $+10^{\circ}$ rampwise, Fig. 6b). The ensuing increase in groundspeed (figure $8 b$ ) automatically causes the micro-flyer to rise (figure 8a) because the feedback loop slaves $h$ to $V_{x}$, effectively maintaining the actual ratio $\omega=V_{\sqrt{ }} h$ relatively constant throughout take-off (Fig. 8d). Upon reaching a steady groundspeed of $3 \mathrm{~m} \cdot \mathrm{~s}^{-1}$ (Figure 8 b ), the MH can be seen to have cruised at a steady groundheight (of around one meter: figure 8a), which depends on the OF set point (here, $\omega_{\text {set }}=3$ rad.s ${ }^{-1}$ : figure 8 c ).

### 7.2 Terrain following

An increase in relief absolute altitude is regarded as a "disturbance" that causes a reduction in the local groundheight $h$. This disturbance will be compensated for by an increase in flight altitude (see figure 6a). Figure 9a shows that the MH is able to clear a gentle slope. Figure $9 b$ confirms the prediction made at the end of Section 5 that the groundspeed hardly changes in spite of the changes in the mean flight force F needed to change the altitude (see figure 6b).
Insects are able to follow a terrain or a canopy and the above scheme may explains how they achieve this feat. Migrating butterflies crossing narrow canyons fly down into the gully and
across the bottom. When traversing dense forests, they clear the canopy at a distance roughly equal to their previous height above the ground, and once they have crossed this large obstacle, they re-descend (Srygley \& Oliveira, 2001).


Figure 9. Terrain following by the miniature helicopter (Fig. 7a) recorded at 4 different groundspeeds ( $1 \mathrm{~m} / \mathrm{s} ; 1.5 \mathrm{~m} / \mathrm{s} ; 1.8 \mathrm{~m} / \mathrm{s} ; 3 \mathrm{~m} / \mathrm{s}$ ). The altitude $\mathbf{z}-1$ plotted in (a) is that of the landing gear which is at a distance $\mathrm{l}=0.3 \mathrm{~m}$ under the eye. The higher the speed, the higher the clearance from the terrain (From Ruffier \& Franceschini, 2003)
Figure 9 shows that when the MH is set to travel at a higher speed, it automatically maintains a greater height above the terrain, giving rise to a "safe" flight. This is precisely the flight pattern that was observed in dung-beetles: they fly higher at higher groundspeeds and lower at lower speeds (Steiner, 1953). Likewise, bees cruise at a height roughly proportional to their horizontal flight speed (see Fig. 7c in Srinivasan et al., 2000). In other words, there is a similarly tight link between groundspeed and groundheight in insects and our helicopter.

### 7.3 Flying against wind

Locusts and other migrating insects have long been known to descend under headwind and ascend during lull (Kennedy, 1951, Srygley \& Olivera, 2001). Under headwind conditions, Braüninger (1964) observed that bees were flying at such a low speed and so close to the ground that he was able to accompany them on his bicycle all the way to the nectar source.
Windspeed disturbs the OF autopilot but the point of application of this disturbance differs from that of the relief disturbance (see Fig. 6a). A light headwind (produced by a fan) was observed to reduce the MH groundspeed (Fig. 10b, dotted curve). The reduction in speed forced the groundheight $h$ to decrease in similar proportion (figure 10a, dotted curve). Since the speed increased again when the robot left the wind-swept region of the arena, the MH rose again. This MH flight pattern therefore accounts particularly well for the field data cited above.
A strong head wind reduced the groundspeed so much that it forced the MH to land (Fig. 10a, continuous curve). Landing was smooth, however, because $h$ was decreased under
visually closed loop, in proportion to $V_{x}$. This finding is highly reminiscent of what occurs in many insects and birds, which descend by headwind and settle when the wind grows stronger (Kennedy, 1939, 1951, Williams, 1965; Alestam, 1990). Given the natural decrease in wind speed that occurs close to the earth in the boundary layer, these reactions can be said to be "ecological" because they allow the insects to travel farther for the same amount of energy (this point is discussed extensively in Ruffier \& Franceschini, 2005). Whether for terrain following or for the reaction to head wind, the control scheme (Fig. 6) underlines the direction of causality: it is the change in groundspeed that induces the change in groundheight, and not the other way round (this point is discussed further in Section 8).


Figure 10. Reaction of the miniature helicopter (MH) to head wind (dashed curves: low head wind, $\boldsymbol{V}_{w}=0.5 \mathrm{~m} / \mathrm{s}$; continuous curves: strong head wind, $\boldsymbol{V}_{w}=1.5 \mathrm{~m} / \mathrm{s}$ ). The OCTAVE autopilot (Fig. 6a) makes the MH react to the reduction in speed (b) by reducing its altitude (a). The strong wind forced the MH to land at negligible forward speed (continuous curves). In these experiments, a $30 \times 20 \mathrm{~cm}$ planar airfoil was mounted perpendicular to the travel axis to catch the wind better, emphasizing the effect (From Ruffier \& Franceschini, 2004)

### 7.4 Flying over mirror-smooth water

Bees crossing mirror-smooth water during foraging trips were reported to fly lower and lower until crashing head-first into the water. They were rescued from drowning only when the water surface was rippled or when a slatted bridge was placed on the water surface to provide sufficient contrast (Heran \& Lindauer, 1963). Recent studies on bees trained to fly across a lake did not quite confirm these early results (Tautz et al., 2004), but the (large) lake may not have been as ripple-free as the (small) flooded quarry used in the original experiments.
A featureless expanse of water no longer provides the animal's eye with any contrasting features, and the OF sensor will no longer respond. If we make the feedback signal $\omega_{\text {meas }}=0$ in figure 6 a, the error signal $\boldsymbol{\varepsilon}$ becomes large and negative (equal to $\omega_{s t t}$ ), which leads to a fatal decrease in groundheight $\boldsymbol{h}$. The OF regulator therefore may account for the puzzling finding that bees plunge straight into calm water during their foraging trip. The closed feedback loop irrevocably pulls the insect down whenever the OF sensor fails to respond. A similarly disastrous tendency was observed in the MH when it was flying over a ground that was made deliberately blank over a distance of 1.5 meters. A simple way to rescue the MH flight from this dangerous situation consisted in holding the last OF measurement for some time (e.g., 0.5 seconds, see figure 9 in Ruffier \& Franceschini, 2005).

### 7.5 Landing

Video recording of landing honeybees showed that their grazing landings on a flat surface occur with a constant slope (Srinivasan et al., 2000). Bees were assumed to meet two requirements to be able to land smoothly: "(i) adjusting the speed of forward flight to hold constant the angular velocity of the image of the surface as seen by the eye, (ii) making the speed of descent proportional to the forward speed"(Srinivasan et al., 2000). Our OF regulator scheme (Fig. 6a) controls neither the forward flight speed nor the descent speed, yet predicts that the bee should land with a constant slope, as actually observed in the MH flight pattern (Fig. 8a, right part).
Both MH's and bees' landing behavior can be explained in the same way because the surge dynamics can be described in both cases by similar (first order) linear differential equations (see Supplements in Franceschini \& al., 2007).
Landing of the MH is initiated by commanding it to gradually pitch backwards to the vertical (at arrowhead 3 in figure 8a), which induces a gradual loss of speed (figure 8b). The groundheight $h$ is then bound to decrease proportionally to $V_{x}$ since the feedback loop strives to make $V_{x} / h=$ constant (Eq. 2).
The final approach actually starts when the MH has regained its completely upright position (arrowhead 4 in Fig. 8a). From this moment onwards, $V_{x}$ decreases exponentially with the "surge time constant" $\tau_{M H}=2.15$ s (see figure 4 in Franceschini \& al., 2007) and the feedback loop forces $h$ (and therefore its derivative $d h / d t$, i.e., the descent speed $V_{z}$ ) to decrease with the same time constant. The descent speed:groundspeed ratio $V_{z}(\mathrm{t}): V_{x}(\mathrm{t})$, that is, the descent slope, will therefore remain constant throughout the final approach, as actually observed in both landing bees (Srinivasan et al., 2000) and the landing MH (figure 8a).
In (Franceschini \& al., 2007), we established that the ensuing descent angle $\alpha$ [rad] can be calculated as follows :

$$
\alpha=-\arctan \left(\frac{1}{\omega_{s e l} \tau}\right)
$$

where $\omega_{s e t}=$ OF set point [rad.s ${ }^{-1}$ ], and $\tau=$ surge time constant [s] of the $\mathrm{MH}\left(\tau_{M H}=2.15 \mathrm{~s}\right)$.
This means that the slope of the final approach depends on only two parameters: the surge time constant and the OF set point.
If the constant-slope landing glide observed in bees reflects the presence of an OF regulator onboard these insects, then by substituting the landing data obtained on bees (Srinivasan et al., 2000): $\alpha_{B E E}=-28^{\circ}$; $\omega_{\text {SetBEE }}=500$ deg.s ${ }^{-1}$, into Eq. 3, we can determine the bee's surge time constant:

$$
\tau_{\mathrm{BEE}}=0.22 \mathrm{~s}
$$

It is striking that this value predicted by our OF regulator model matches closely the time constant values $(0.22 \mathrm{~s}, 0.27 \mathrm{~s}, 0.29 \mathrm{~s}$ and 0.57 s$)$ that we derived from the data of Srinivasan and coworkers (2000) on the exponential decrease in bees' groundheight with time. This makes the OF regulator scheme (Fig. 6a) an appealing hypothesis for the control system that is implemented onboard the bee.

## 8. A dual optic flow regulator for joint speed control and lateral obstacle avoidance

The model presented above differs from another one where the OF would control the groundspeed $V_{x}$ rather than the groundheight $h$ (Preiss \& Kramer, 1984; Srinivasan et al., 1996, 2000; Baird et al., 2005). If the ventral OF controlled $V_{x}$ instead of $h$ in figure 6a, this would lead to strikingly different flight patterns from those observed in insects by previous authors, as follows:
(i) instead of following a slanting terrain, as migrating butterflies (Williams, 1965; Srygley \& Oliveira, 2001) and our MH do, the insect would gradually decelerate until touching the rising ground at a negligible speed, thus inopportunely interrupting its journey.
(ii) instead of descending in a headwind and rising in a tailwind, as occurs in honeybees (Braüninger, 1964), locusts (Kennedy, 1951), dung-beetles (Steiner, 1953), mosquitoes (Kennedy, 1939), and our MH (figure 10), the insect would compensate for the unfavourable headwind by increasing its airspeed without changing its groundheight.
These two models can be reconciled, however, if we add the hypothesis that another OF regulator, based on OF sensors from the lateral parts of the eyes, may be in charge of controlling the groundspeed $V_{x}$.
In line with the OCTAVE autopilot, we designed the LORA III autopilot (LORA III stands for Lateral Optic flow Regulator Autopilot, mark III), which is able to control both the forward speed $V_{x}$ of an aerial vehicle and its lateral distances $D_{r}$ and $D_{l}$ to the two walls of a corridor (Fig. 11).


Figure 11. The LORA III autopilot is a dual OF regulator that enables a fully actuated hovercraft to navigate in a (straight or tapered) corridor by controlling its forward speed $V_{x}$ and its distance $\boldsymbol{D}_{r}, \boldsymbol{D}_{l}$ to the walls jointly, without requiring any speed or distance measurements (From Serres et al., 2008a)
We showed the feasibility of this scheme in simulation experiments where a miniature hovercraft navigates in a straight or tapered corridor (Serres et al., 2008a). Our hovercraft is equipped with two additional lateral thrusters that make it fully actuated. This means that it is capable of independent side-slip and forward slip, like hymenopteran insects (Zeil et al., 2008). The groundspeed is constrained by the environment, as observed on bees navigating in a
straight or tapered corridor (Srinivasan et al., 1996). The clearance to the walls is constrained by the environment as well, but the robot does not show a systematic "centering behavior", in contrast with former observations on bees (Kirschner \& Srinivasan, 1989). However, the robot's "wall-following behavior" observed is fully consistent with the results of our own behavioural experiments on bees navigating in a large corridor (Serres et al. 2008b).
LORA III is based on only two OF sensors (one looking to the right, one to the left). The forward speed $V_{x}$ is controlled by the error signal $\varepsilon_{\text {Fwd }}$ between the sum of the two OFs (right and left) and the forward OF set point $\omega_{\text {setFwd }}$. The lateral clearance from the walls is controlled by the error signal $\boldsymbol{\varepsilon}_{\text {side }}$ between the larger of the two OFs and the sideways OF set point $\omega_{\text {setside }}$. The LORA III dual OF regulator accounts particularly well for both types of behaviours. The typical "wall-following behavior" that we observed on both the bees and the robot can be shown to degenerate into a "centering behavior" for particular relative values between the two OF set points $\omega_{\text {setFwd }}$ and $\omega_{\text {setSide }}$ (Serres et al., 2008a, Serres, 2008). The advantage of this control scheme is that it determines both the forward speed and the clearance from the walls on the sole basis of two parameters - two OF set points - without any needs for measuring forward speed, lateral distances and corridor width, that is, without using any velocimeters or range finders. The simulation experiments performed in straight or tapered corridors were shown to mimic the honeybee's behaviour remarkably well (Serres et al., 2008a).

## 9. OSCAR as a yaw control autopilot

Electrophysiological recordings on freely flying flies have revealed that the muscle attached to the base of the retina (Burtt \& Patterson, 1970; Hengstenberg, 1971) causes retinal microscanning : the whole retinal mosaic (Fig.1a) translates repeatedly at about 5 Hz by a few micrometers underneath the facet mosaic, causing the visual axes to rotate by a few degrees (Franceschini \& Chagneux, 1997). Such a purely rotational OF is exploited on-board another aerial robot we built: robot OSCAR (Fig. 12a).



Figure 12. (a) OSCAR is a 100-gram twin-engine robotic demonstrator that sees. It is equipped with an elementary eye (visible on top), composed of a lens and only two photoreceptors driving an EMD of the type shown in Fig. 3 (Photo by H. Raguet) (b) Smooth pursuit (tracking) of a dark edge (contrast $\mathrm{m}=0.4$ ) displaced sinusoidally at 0.15 Hz in a frontal plane situated 1.3 meters from the eye (From Viollet \& Franceschini, 1999b)

This project was based on the assumption that microscanning in flies operates in connection with motion detection. On the basis of this fly's retinal microscanning process, we developped an optronic sensor (Viollet \& Franceschini, 1999a), and a novel aerial robot, equipped with this sensor (Viollet \& Franceschini, 1999b, 2001). OSCAR is attached to a thin, 2-meter long nylon wire secured to the ceiling of the laboratory. It is free to adjust its yaw by driving its two propellers differentially, and it does so in spite of the many (aerodynamic or pendular) disturbances it encounters.
Thanks to its microscanning eye, this miniature (100-gram) robot is not only able to fixate a nearby "target" (a dark edge or a bar) steadily but also to track it at angular speeds of up to $30^{\circ} / \mathrm{s}$ - a value similar to the maximum tracking speed of the human eye. It cannot, however, estimate the distance from the target since it deals only with the rotational OF.
The OSCAR sensor is able to detect a bar subtending an angle much smaller than the interreceptor angle $\Delta \varphi=4^{\circ}$, making it applicable for nonemissive power line detection onboard fullscale helicopters (Viollet et al., 2006). The OSCAR sensor is also able to locate an edge with great positioning accuracy ( 0.1 degrees, which is 40 times smaller than $\Delta \varphi$ ): OSCAR has acquired hyperacuity (Viollet \& Franceschini, 2001). This property makes the principle appealing for accurate stabilization of various platforms (Franceschini et al., 2005). Inspired by the fly's thoracic halteres that have long been known to act as gyroscopes, we equipped the OSCAR robot with an additional inertial feedback based on a MEMS rate gyro. The interplay of the two (visual and inertial) control loops enhances both the dynamic stability and performances of the tracking system (Viollet \& Franceschini, 1999b, 2001). More recently, we introduced a mechanical decoupling between eye and body, and implemented a genuine vestibulo-ocular reflex that was key to maintaining the robot's gaze perfectly fixed on the target (Viollet \& Franceschini, 2005, Kerhuel et al., 2007). Robot OSCAR II is a novel 100gram aerial robot whose gaze, decoupled from the heading, remains robustly locked onto a target in spite of various perturbations that we deliberately applied to the body. It is the gaze orientation of OSCAR II that ultimately controls its heading (Viollet et al, this volume).

## 10. Summary and conclusion

In this chapter, we reported about our attempts to understand how flying insects could behave on the basis of OF cues. We presented several bio-inspired, OF based visual guidance principles and showed that they may be helpful in the context of autonomous guidance of MAVs over a terrain or in a corridor. The principles we came up with differ markedly from a current OF based navigation strategy. The latter requires that OF sensors be associated with a groundspeed sensor (based on GPS, for example) in order to determine the groundheight (see Eq 1) to achieve terrain following or landing (Barrows et al., 2003; Barber et al., 2005; Srinivasan et al., 2006; Garrat \& Chahl, 2008).
OCTAVE and LORA III autopilots harness the power of the translational OF in a quite different manner. They do not need to estimate any speed or range, and they do not aim to achieve any "speed holding" or "range holding" abilities. They aim instead at producing safe behavior by a process that can be called "OF hold". These autopilots consist of feedback control loops, called «optic flow regulators». Their block diagrams (Figures 6 \& 11) pinpoint which variables are to be either measured or controlled or regulated, and give the causal and dynamical relationships between these variables. Three (ventral, forward and sideways) OF regulators strive to maintain a given OF $\omega$ constant by acting upon the lift, the forward thrust
or the side thrust, respectively. OF sensors are shown to be sufficient for these tasks and there is no need to measure (or estimate) groundspeed, groundheight and clearance to the walls in a corridor. Simulation experiments and robotic demonstrators showed that risky maneuvers such as automatic takeoff, ground avoidance, level flight, terrain following, landing, centering, wall following, and groundspeed control can all be successfully performed on the basis of these three OF regulators. Moreover, these control schemes were found to account for a series of puzzling, seemingly unconnected flying abilities observed by many authors over the last 70 years in numerous insect species, suggesting that $O F$ regulators may well be implemented onboard insects (see Sections $7 \& 8$ ).
Three parameters - the three OF set points $\omega_{s e t}$ of Fig. 6 and 11 - would suffice to determine the flight behavior, at least in the simplified, random but richly contrasted environments considered here. Onboard the insect, these set points may depend on either innate, internal or external factors.
Electronic implementation of an OF regulator is not very demanding - nor is its neural implementation - since it relies upon a few linear operations (such as summation, difference and various filters) and nonlinear operations (such as minimum or maximum detection). The worthiest component of an OF regulator is the OF sensor itself. Like the honeybee's Velocity Tuned (VT) neurons (Ibbotson, 2001), the neuromorphic OF sensors we have been building since 1985 deliver an output that grows monotonically with the OF $\omega$, with little dependence on spatial frequency and contrast (see Section 3). They respond over a 10 -fold range in OF (from $40^{\circ}$ s to $400^{\circ} / \mathrm{s}$ : Ruffier et al., 2003; Aubépart \& Franceschini, 2007). This range is largely sufficient because each OF sensor needs only to detect a deviation $\boldsymbol{\varepsilon}$ from the OF set point. The miniature helicopter flight pattern illustrates this point: the ventral $O F$ regulator produces a behavior such that the OF output $\omega_{\text {meas }}$ varies little throughout the journey, from takeoff to landing (Fig. 8c). A similar observation holds for the forward and sideways OF regulators (see Figs 7c,e \& 12c,e in Serres et al., 2008a).
Each OF sensor at work on our proof of concept robots is characterized by a low resolution. The inter-receptor angle in the eyes of OCTAVE, LORA III and OSCAR robots is around $4^{\circ}$, a value close to the interommatidial angle ( $5^{\circ}$ ) of the fruitfly's eye (Franceschini, 1975). The field of view (FOV) of the eyes and the provocatively small number of pixels ( 2 pixels per ventral or lateral eye) will obviously need to be increased. Covering the frontal part of the FOV, in particular, will be essential to sense and avoid obstacles directly in the flight path, and here the retinal microscanning principle (Section 9) could be a great help (Mura \& Franceschini, 1996; Viollet \& Franceschini, 2001). A recent study showed that a more frontally oriented EMD may provide OCTAVE autopilot with a feedforward anticipatory signal that helps the robot climb steeper rises (Ruffier \& Franceschini, 2008).
The control schemes we proposed are simple but thus far limited to the use of the maximal OF sensed either ventrally or laterally, perpendicular to the heading direction. Once developed beyond the state of the current minimalistic robotic demonstrators, OCTAVE and LORA III principles could potentially be harnessed to guide a MAV indoors or through complex terrains such as mountainous canyons or urban environments. Since the control systems are parsimonious and do not rely on GPS or any emissive (power-hungry) sensors such as RADARs or LADARs, they promise to match the draconian constraints imposed upon micro aerial vehicles, in terms of size, mass and consumption.
For the same reasons, these autopilots could potentially be developped for robotic missions on other planets or moons. A Martian lander could make use of the OCTAVE principle,
since it ensures safe automatic landing (see Section 7.5). A rover flying in Martian canyons and gullies could make use of the LORA III principle, since both the craft's groundspeed and its clearance to the walls would automatically depend on the canyon's width (see Section 8, and Serres et al, 2008a). Whether on Earth or on another celestial body, the craft would need to receive from the Earth based (or orbital) station only three low bandwidth signals: the values of the OF set points.

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## 12. References

Alestam, T. (1990). Bird Migration. Cambridge University Press
Aubépart, F. \& N. Franceschini., N. (2007). Bio-inspired optic flow sensors based on FPGA: application to micro-air vehicles. Journal Microprocessors and Microsystems 31, 408419
Aubépart, F., El Farji, M. \& Franceschini, N. (2004). FPGA implementation of elementary motion detectors for the visual guidance of micro-air vehicles. Proceedings of IEEE International Symposium on Industrial Electronics (ISIE'2004), Ajaccio, France, pp. 7176
Aubépart, F., Serres, J., Dilly, A., Ruffier, F. \& Franceschini., N. (2008). Field programmable gate arrays (FPGA) for bio-inspired visuo-motor control systems applied to microair vehicles, In: Aerial Vehicles, published by In-Tech
Baird, E., Srinivasan, M.V., Zhang, S.W. \& Cowling., A. (2005). Visual control of flight speed in honeybees. Journal of Experimental Biology 208, 3895-3905
Baird, E., Srinivasan, M.V., Zhang, S.W., Lamont R. \& Cowling, A. (2006).Visual control of flight speed and height in the honeybee. In: From Animals to Animats 9, S. Nolfi et al. (Eds.), 4095, 40-51
Barber, D.B.; Griffiths, S..R.; McLain, T.W.;\& Beard, R.W. (2005) Autonomous landing of miniature aerial vehicles. In: Proceedings of American Institute of Aeronautics and Astronautics Conference
Barrows, GL, Neely, C. \& Miller, K.T. (2001). Optic flow sensors for MAV navigation. In : Fixed and flapping wing aerodynamics for micro-air vehicle applications. Progress in Astronautics and Aeronautics, Vol. 195, 557-574
Barrows, GL; Chahl, J.S.; Srinivasan MV. (2003) Biologically inspired visual sensing and flight control. Aeronautic Journal 107, 159-168
Blanès, C. (1986) Appareil visuel élémentaire pour la navigation à vue d'un robot mobile autonome. MS thesis in Neuroscience, University of Aix-Marseille II, Marseille, France

Blanès, C. (1991). Guidage visuel d'un robot mobile autonome d'inspitation bionique. PhD thesis, Polytechnical Institute, Grenoble, France (thesis work at the Neurocybernetics lab, CNRS, Marseille, France)
Borst, A. \& Haag, J. (2002). Neural networks in the cockpit of the fly. Journal of Comparative Physiology A188, 419-437
Braitenberg, V. (1967). Patterns of projection in the visual system of the fly, I/Retina-Lamina projections. Experimental Brain Research 3, 271-298
Bräuninger, H.D. (1964). Uber den Einfluss meteorologischer Faktoren auf die Entfernungsweisung im Tanz der Bienen. Zeitschrift für Vergleichende Physiologie 48, 1-130
Buchner E. (1984). Behavioral analysis of spatial vision in insects In: Photoreception and Vision in Invertebrates, Ali, M. (Ed.), New-York : Plenum, 561-621
Burtt, J. \& Patterson, J. (1970). Internal muscle in the eye of an insect. Nature 228, 183-184
David, C. (1978). The relationship between body angle and flight speed in free-flying Drosophila. Physiological Entomology 3, 191-195
David, C. (1982). Compensation for height in the control of groundspeed by Drosophila in a new 'barber's pole' wind tunnel. Journal of Comparative Physiology A 147, 1432-1351
Egelhaaf, M. \& Borst, A. (1993). Movement detection in Arthropods. In : Visual motion an its role in the stabilization of gaze, F. Miles \& J. Wallman, Eds.,Amsterdam: Elsevier, 5377
Franceschini N. (1975). Sampling of the visual environment by the compound eye of the fly: fundamentals and applications, in: Photoreceptor Optics, A. Snyder, R. Menzel (Eds), Berlin : Springer, Chapt. 17, 98-125
Franceschini, N. (1984). Chromatic organisation and sexual dimorphism of the fly retinal mosaic. In: Photoreceptors. Borsellino, A. and Cervetto, L. (Eds.), pp. 319-350
Franceschini, N. (1985). Early processing of colour and motion in a mosaic visual system. Neuroscience Research, Supplement 2, 517-549
Franceschini, N. (1992). Sequence-discriminating neural network in the eye of the fly. In: Analysis and Modeling of Neural Systems, F.H.K. Eeckman (Ed.), Norwell, USA, Kluwer Acad. Pub., 142-150.
Franceschini, N. (2007). Sa majesté des mouches. In: Voir l'Invisible, JP.Gex (Ed.), Paris: Omnisciences (2007), pp 28-29
Franceschini, N. (2008). Towards automatic visual guidance of aerospace vehicles: from insects to robots. Acta Futura : Proceedings of the ESA International Symposium on Advanced Concepts : «A Bridge to Space» Noordwick, (in press)
Franceschini, N. \& Chagneux, R. (1997). Repetitive scanning in the fly compound eye. Proceedings of the 25th Göttingen Neurobiology Conference (N. Elsner, H. Wässle, Eds.) Stuttgart, : G. Thieme, 279
Franceschini, N., Blanes, C. \& L. Oufar, L. (1986). Appareil de mesure, passif et sans contact, de la vitesse d'un objet quelconque. Technical Report ANVAR/DVAR, №51549, Paris.
Franceschini, N., Pichon, J.M. \& Blanès, C. (1992). From insect vision to robot vision. Philosophical Transactions of the Royal Society London B 337, 283-294
Franceschini, N., Kirschfeld, K., Minke, B. (1981a). Fluorescence of photoreceptor cells observed in vivo. Science 213, 1264-1267
Franceschini, N., Hardie, Ribi, W., Kirschfeld, K. (1981b). Sexual dimorphism in a photoreceptor. Nature 291, 241-244

Franceschini, N., Riehle, A. \& Le Nestour A. (1989). Directionally selective motion detection by insect neurons. In: Facets of Vision, D.G. Stavenga and R.C. Hardie (Eds.), Berlin, Springer, 360-390.
Franceschini, N., Ruffier, F. \& Serres, J. (2007). A bio-inspired flying robot sheds light on insect piloting abilities. Current Biology 17, 329-335
Franceschini, N.; Viollet, S. \& Boyron, M. (2005). Method and device for hyperacute detection of an essentially rectilinear contrast edge, and system for fine following and fixing of said contrast edge. International Patent PCT/FR2005/11536
Franceschini, N., Ruffier, F., Viollet, S. \& Boyron, M. (2003). Steering aid system for altitude and horizontal speed, perpendicular to the vertical, of an aircraft and aircraft equipped therewith. International Patent PCT/FR2003/002611
Garratt M.A. \& Chahl, J.S. (2008). Vision-based terrain following for an unmanned rotorcraft. Journal of Field Robotics 25, 284-301
Gibson, J.J. (1950). The perception of the visual world, Boston: Houghton Mifflin
Gibson J.J., Olum, P. \& Rosenblatt, F. (1955). Parallax and perspective during aircraft landings. American Journal of Psychology 68, 372-395
Green, WF, Oh, Y. \& Barrows, G. (2004).Flying insect inspired vision for autonomous axial robot maneuvers in near earth environments. Proceedings of the IEEE International Conference on Robotics and Automation (ICRA04) 2343-2352
Hardie R.C. (1985). Functional organization of the fly retina. Progress in Sensory Physiology 5, Ottosson D. (Ed.), Springer, Berlin
Hassenstein, B. \& Reichardt, W. (1956) Systemtheoretische Analyse der Zeitreihenfolgen undVorzeichenauswertung bei der Bewegungsperzeption des Rüsszelkäfers Chlorophanus. Zeitschrift für Naturforschung 11b, 513-524
Hausen , K. (1993). Decoding of retinal image flow in insects. In : Visual motion an its role in the stabilization of gaze, F.A. Miles and J. Wallman (Eds.) Amsterdam : Elsevier, 203235
Hausen, K. \& Egelhaaf, M. (1989) Neural mechanisms of course control in insects. In: Facets of vision; D.G. Stavenga \& R.. Hardie, Berlin, Springer pp. 391-424
Hengstenberg, R. (1971). Das Augenmuskel System der Stubenfliege Musca domestica. Kybernetik 2 56-77
Heran, H. (1955). Versuche über die Windkompensation der Bienen. Naturwissenschaften 5, 132-133
Heran, P. \& Lindauer, M. (1963). Windkompensation und Seitenwindkorrektur der Bienen Flug über Wasser. Zeitschrift für vergleichende Physiologie 47, 39-55
Heisenberg, M, Wolf ,R. (1984).Vision in Drosophila. Berlin, Springer.
Ibbotson, M.R. (2001). Evidence for velocity-tuned motion-sensitive descending neurons in the honeybee. Proceedings of the Royal Society, London B 268, 2195-2201
Kennedy, J.S. (1939). Visual responses of flying mosquitoes. Proceedings of the Zoogical Society of London 109, 221-242
Kennedy, J.S. (1951). The migration of the desert locust (Schistocerca gregaria Forsk.) I. The behaviour of swarms. Phil. Transactions of the Royal Society, London, B 235, 163-290
Kerhuel, L., Viollet, S. \& Franceschini, N. (2007). A sighted aerial robot with fast gaze and heading stabilization. Proceedings of the IEEE International Conference on Intelligent Robots and Systems (IROS), San Diego, USA, pp. 2634-2641

Kien, J. (1975) Neuronal mechanism subserving directional selectivity in the locust optomotor system. J. Comp. Physiol. 113: 161-179
Kirchner, W.H. \& Srinivasan, M.V. (1989). Freely moving honeybees use image motion to estimate distance. Naturwissenchaften 76, 281-282
Kirschfeld, K. (1967). Die Projektion der optischen Umwelt auf das Raster der Rhabdomere im Komplexauge von Musca. Experimental Brain Research 3, 248-270
Kirschfeld, K.; Franceschini, N. (1968). Optische Eigenschaften der Ommatidien im Komplexauge von Musca. Kybernetik 547-52
Kirschfeld, K.; Franceschini, N., Minke, B. (1977). Evidence for a sensitizing pigment in fly photoreceptors » Nature 269, 386-390
Koenderink, J.J. (1986). Optic flow. Vision Research 26, pp. 161-79
Kramer, J., Sarpeshkar, R. \& Koch C. (1995). An analog VLSI velocity sensor. In Proceedings of IEEE International ymposium on Circuits and Systems, Seattle, USA, pp 413-416
Krapp, H, Hengstenberg, B, Hengstenberg, R.(1998). Dendritic structure and receptive-field organisation of optic flow processing interneurons in the fly, Journal of Neurophysiology 79, 1902-1917.
Laughlin, S. (1984). The role of parallel channels in early visual processing by the arthropod compound eye. In: Photoreception and vision in invertebrates. M.A. Ali Ed., Plenum, 457-481
Lee, D. N. (1980). The optic flow field: the foundation of vision. Philosophical Transactions of the Royal Society London B 290, 169-179
Marr, D. (1982). Vision, San Francisco: Freeman
Moeckel, R. \& Liu S.C. (2007). Motion detection circuits for a time-to-travel algorithm. Proceedings IEEE Int. Symposium Circuits and Systems (ISCAS07), pp. 3079-3082
Mura, F. \& Franceschini., N. (1994). Visual control of altitude and speed in a flying agent. In : From Animals to Animats III, D. Cliff et al. (Eds), Cambridge: MIT Press, 91-99
Mura, F.; Franceschini, N. (1996). Obstacle-avoidance in a terrestrial mobile robot provided with a scanning retina. Proceedings IEEE International Symposium on Intelligent Vehicles. N. Aoki and I. Masaki, eds., Seikei University, Tokyo, Japan, pp. 47-52.
Netter, T. \& Franceschini, N. (1999). Neuromorphic optical flow sensing for nap-of-the-Earth flight. In : Mobile Robots XIV, D. Gage and H. Choset (Eds), SPIE, Vol. 3838, 208-216
Netter, T. \& Franceschini, N. (2002). A robotic aircraft that follows terrain using a neuromorphic eye. In : Proceedings of the IEEE International Conference on Intellignent Robots and Systems (IROS) Lausanne, Switzerland, pp. 129-134
Olberg R.M. (1981) Object- and Self- Movement detectors in the ventral nerve cord of the dragonfly. J. Comp. Physiol. 141, 327-334
Pichon, J-M., Blanès, C. \& Franceschini, N. (1989). Visual guidance of a mobile robot equipped with a network of self-motion sensors. In : Mobile Robots IV, W.J. Wolfe , W.H. Chun (Eds.) Bellingham, U.S.A : SPIE , Vol. 1195, 44-53

Portelli, G., Serres, J., Ruffier F. \& Franceschini, N. (2008). An Insect-Inspired Visual Autopilot for Corridor-Following. Proceedings of the $2^{\text {nd }}$ Biennial IEEE Int. Conference on Biomedical Robotics and Biomechatronics, BioRob 08, Scottsdale, USA, pp. 19-26
Preiss, R. (1992). Set point of retinal velocity of ground images in the control of swarming flight of desert locusts. Journal Comparative Physiology A 171, 251-256

Preiss, R. \& Kramer, E. (1984). Control of flight speed by minimization of the apparent ground pattern movement. In: Localization and Orientation in Biology and Engineering, D. Varju and H. Schnitzler, eds. (Berlin, Springer), 140-142
Pudas, M. ; Viollet, S. ; Ruffier, F. ; Kruusing, A. ; Amic, S. ; Leppävuori, S. \& Franceschini, N. (2007). A miniature bio-inspired optic flow sensor based on low temperature cofired ceramics (LTCC) technology, Sensors and Actuators A 133, 88-95
Reichardt, W. (1969): Movement perception in insects. In: Processing of Optical Data by Organisms and by Machines, W. Reichardt (Ed.), New York: Academic Press , 465-493
Riehle, A. \& Franceschini, N. (1984). Motion detection in flies: parametric control over ONOFF pathways. Experimental Brain Research 54, 390-394
Ruffier, F. \& Franceschini, N. (2003) OCTAVE, a bioinspired visuo-motor control system for the guidance of Micro-Air Vehicles, In: Bioengineered and Bioinspired Systems, A. Rodriguez-Vazquez et al., Eds., Bellingham, U.S.A , SPIE, Vol. 5119, 1-12
Ruffier, F. \& Franceschini, N. (2004a). Visually guided micro-aerial vehicle: automatic takeoff, terrain following, landing and wind reaction. Proceedings IEEE International Conference on Robotics and Automation (ICRA04), New Orleans, USA, pp. 2339-2346
Ruffier, F. \& Franceschini, N. (2004b). Optic flow based AFCS for rotorcraft automatic maneuvring (terrain following, takeoff and landing). Proceedings of the 30th European Rotorcraft Forum, AAF/CEAS, Marseille, 71.1-71.9.
Ruffier, F. \& Franceschini, N. (2005). Optic flow regulation: the key to aircraft automatic guidance. Robotics and Automomous Systems 50, 177-194
Ruffier, F. \& Franceschini, N. (2008). Aerial robot piloted in steep relief by optic flow sensors. Proceedings of the IEEE/RSJ International Conference on Intelligent Robots and Systems (IROS08), Nice, France, pp. 1266-1271.
Ruffier, F.; Viollet, S.; Amic, S. \& Franceschini, N. (2003). Bio-inspired optical flow circuits for the visual guidance of micro-air vehicles. Proceedings the IEEE International Symposium on Circuits and Systems (ISCAS), Bangkok, Thaïland, III, pp. 846-849
Serres, J. (2008). De l'abeille au robot: la «régulation du flux optique», PhD Thesis, University of Montpellier, France (Thesis work from the Biorobotics lab, CNRS, Marseille, France).
Serres, J.; Dray, D.; Ruffier, F. \& Franceschini, N. (2008a). A vision-based autopilot for a miniature air vehicle: joint speed control and lateral obstacle avoidance. Autonomous Robots 25, 103-122
Serres, J.; Masson, G.; Ruffier, F. \& Franceschini, N. (2008b). A bee in the corridor : centring and wall following. Naturwissenschaften, 95 (12) 1181-1187
Shoemaker, P.A. ; O'Caroll, D.C. \& Straw, A.D. (2005). Velocity constancy and models for wide-field visual motion detection in insects. Biological Cybernetics, 93, 275-287
Srinivasan, M.V. (1993). How insects infer range from visual motion. In: Visual motion and its role in the stabilization of gaze. F.A. Miles and J. Wallman, Eds. Amsterdam : Elsevier, 139-156
Srinivasan, M.; Thurrowgood, S. \& Soccol, D. (2006). An optical system for guidance of terrain following in UAVs. Proceedings of the IEEE International Conference on video and signal based surveillance AVSS06
Srinivasan, M.V.; Lehrer, M.; Kirchner, W.H. \& Zhang, S.W. (1991). Range perception through apparent image speed in freely flying honeybees. Visual Neuroscience 6, 519-535

Srinivasan, M.V., Zhang, S.W. and Chandrashekara K. (1993). Evidence for two distinct movement-detecting mechanisms in insect vision. Naturwissenchaften 80, 38-41.
Srinivasan, M.V., Zhang, S.W., Lehrer, M. \& Collett., T. (1996). Honeybee navigation en route to the goal: visual flight control and odometry. Journal of Experimental Biology, 199, 237-244
Srinivasan, M.V., Zhang, S.W., Chahl, J.S., Barth, E. \& Venkatesh, S. (2000). How honeybees make grazing landings on flat surface. Biological Cybernetics 83, 171-183
Srygley, R.B. \& Oliveira, E.G (2001). Orientation mechanisms and migration strategies within the flight boundary layer. Insect movements: mechanisms and consequences. T.P. Woiwod, D.R. Reynolds and C.D. Thomas (Eds.), CAB international, 183-206
Steiner, G. (1953). Zur Duftorientierung fliegender Insekten. Naturwissenchaften 19, 514-515
Tautz, J., Zhang, S., Spaethe J., Brockman A., Si, A. \& Srinivasan M.V. (2004). Honeybee odometry: Performance in varying natural terrain. PLOS Biology, 2, 915-923
Taylor G.K. \& Krapp, H.G. (2008). Sensory systems and flight stability : what do insects measure and why? In : Advances in Insect Physiol. 34 : Insects mechanisms and control, J. Casas and S.J. Simpson (Eds.) Amsterdam : Elsevier, 231-316

Ullman, S. (1981). Analysis of visual motion by biological and computer systems, Computer 14, 57-69
Viollet, S. \& Franceschini, N. (1999). Biologically-Inspired Visual Scanning Sensor for Stabilization and Tracking. Proceedings of the IEEE International Conference on Intelligent Robots and Systems (IROS 99) Kyon-gyu, Corea, pp. 204-209
Viollet, S. \& Franceschini, N. (1999). Visual servo system based on a biologically-inspired scanning sensor. Proceedings of the Conference on sensor fusion and decentralized control in robotics II, Bellingham, U.S.A : SPIE Vol. 3839, 144-155
Viollet, S. \& Franceschini, N. (2001). Super-Accurate Visual Control of an Aerial Minirobot
In : Autonomous minirobots for Research and Edutainment, U. Rückert, J. Sitte,
U. Witkowski (Eds.), Heinz Nixdorf Institut, Padderborn, Germany, 215-224

Viollet, S. \& Michelis, J. \& Franceschini, N. (2006). Toward a bio-inspired nonemissive powerline detection system. Proceedings of the 32th European Rotorcraft Forum, ERF06, Maastricht, Netherlands, Paper N ${ }^{\circ}$ AD09
Viollet, S. \& Franceschini, N. (2005). A high speed gaze control system based on the Vestibulo-Ocular Reflex. Robotics and Autonomous Systems 50 147-161
Viollet, S., Kerhuel, L. \& Franceschini, N. (2008). A novel biomimùetic steering control system for sighted vehicles. In : Aerial Vehicles, published by In-Tech
Whiteside, T.C. \& Samuel, G.D. (1970). Blur zone. Nature 225, 94-95
Williams, C.P. (1965). Insect migration. London: Collins, second edition
Zeil, J., Boeddeker, N. \& Hemmi, J.M. (2008).Vision and the organization of behavior, Current Biology 18, 320-323


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