Nitric oxide synthase-dependent NADPH-diaphorase activity in the optic lobes of male and female Ceratitis capitata mutants

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The gas nitric oxide (NO) is recognized as an important signalling molecule in the nervous system (Moncada et al., 1991; Bredt & Snyder, 1990), not only in mammals (Bredt & Snyder, 1990; Garthwaite, 1991; Garthwaite et al., 1988; Snyder, 1992; Vincent & Kimura, 1992), but also in all the vertebrate groups (Williams et al., 1994; Brüning et al., 1994; Holqvist et al., 1994; Pisu et al., 2002) and in several invertebrates (Martinez, 1995; Müller & Bicker, 1994; Kurzin et al., 1996; Gibbs & Truman, 1996; Elphick et al., 1996; Pisu et al., 1999).

NO is synthesized from L-arginine and molecular oxygen by the enzyme NO-synthase (NOS) (Bredt & Snyder, 1990), that, using as cofactor NADPH, displays NADPH diaphorase (NADPHd) activity (Garthwaite, 1991; Vincent, 1994; Johansson & Carlberg, 1995); the formation of NO is a Ca$^{2+}$/calmodulin-dependent process (Garthwaite et al., 1988).

NOS containing neurones are present throughout the mammalian central nervous system (CNS) and recently a gene encoding a protein with 43% aminoacid identity to rat neural NOS was reported in D. melanogaster (Regulski & Tully, 1995) providing further evidence that NO is a signalling molecule in the insect CNS (Elphick et al, 1996). In parallel with mammalian studies, the research in invertebrate focused on role for NO in olfaction (Gelperin et al., 1994; Muller & Bicker, 1994) and learning and memory (Robertson et al., 1994). All these studies suggest the high conservation of this signalling system throughout the animal kingdom (Martinez, 1995; Elofsson et al., 1993; Ribeiro et al., 1997; Salieo et al., 1996; Elphick et al., 1993a).

NO is also found at all levels of the vertebrate visual system (Vincent & Kimura, 1992; Kalamarov et al., 1993) and it is thought to be implicated in the visual processes and in the visually guided behaviour of some insects (Elphick et al., 1996;
Bacigalupo et al., 1995; Bicker & Schmachtenberg, 1997).

Since NO is extremely labile and NOS displays NADPHd activity, frequently the method used to demonstrate nitrinergic elements in the brain consisted in the histochemical reaction for NADPH-diaphorase (NADPHd; Thomas & Pearse, 1964). The most important and attractive reason for the interest of neuroanatomist and neurobiologist in this technique arose when NADPHd was identified as a marker for neuronal nitric oxide synthase (Hope et al., 1991). Thus, the relative simple NADPHd histochemical technique was widely used to identify NO producing elements in the brain of representatives of all vertebrate classes (Luebke et al., 1992; Brüning, 1993; Panzica et al., 1994; Alonso et al., 1995; Smeets et al., 1997; Arévalo et al., 1995; Muñoz et al., 1996; Alonso et al., 2000).

The NADPHd staining method in invertebrate has been validated by purification of the locust NOS and demonstration of co-localization for both NOS and NADPHd activities (Elphick et al., 1994).

In a previous work (Conforti et al., 1999), we have found that NADPHd activity is present in the optic lobes of two different strains of the medfly Ceratitis capitata (Diptera, Tephritidae), a wild type eye colour and a white eye mutant line. Here we give histochemical evidence of the differences in NADPHd activity in the optic lobes of four eye colour mutant strains of C. capitata with the aim to further demonstrate a functional relationship between NOS dependent NADPHd activity in the optic lobes and the visual activity of the medfly C. capitata. The orange eye (or) and orange eye, white pupa (or,wp) mutants have orange coloured eyes; the white eye M360 (w,M360) strain presents white-yellow eye and the white eye Heraklion (w,Heraklion) mutant phenotype lacks eye colour pigmentation.

In particular, we have compared the pattern of NADPHd staining, paying attention to the possible role of a sex-dependent NOS activity in the optic lobes of C. capitata.

In fact, it has been previously observed that in many insects some visually guided behaviour patterns differ between the sexes (Franceschini et al., 1981). In particular, in the blowfly Calliphora erythrocephala dimorphism is expressed by differences in the shapes of analogous neurons in males and females, as well as by the presence of some cells in only one sex (Strausfeld, 1980). Sexual dimorphisms, structural differences between the sexes, have been described in the brains of many vertebrate species.

**Materials and Methods**

**Insect strains**

Four adult strains of C. capitata were used: or; or, wp (both orange eyed); w,M360; w,Heraklion (both white eyed). White eye Heraklion is homozygous for the white eye (w) allele of the white locus (w) (Torti et al., 1994). Our experiments were carried out both on males (5 specimens per strain) and females (5 specimens per strain). The four strains were reared in our laboratory at the same conditions (25°C and 65% RH under 12 hours light-12 hours dark conditions).

Visual activities were tested by observation during the routine maintenance and on mating behaviour.

**Tissue preparation**

The exoskeleton was dissected from the anterior part of the puparium and from the anterior face of the head to expose the brain and the optic lobes. Heads were fixed with 4% paraformaldehyde in 0.1 M phosphate buffer, pH 7.35, for 2 h. After washing in the same buffer for 30 min, they were stored in cryoprotective 25% sucrose-phosphate buffer overnight at 4°C and then frozen in liquid nitrogen. Serial 14 μm sections were cut using a cryostat in the frontal plane.

**NADPHd histochemistry**

The sections, from different mutants and sexes, were contemporaneously incubated for 1 hour at 37°C in the dark in the following medium [modified from Van Noorden and Frederiks, 1992 (Fortini & Bonini, 2000)]: 0.1 M phosphate buffer, pH 7.35, containing 15% (w/v) polyvinyl alcohol, 0.5 mM β-NADPH (Sigma, M0, USA), 0.2% Triton-100 and 5 mM nitro blue tetrazolium (NBT, Sigma). After incubation, the sections were rinsed in 0.1 M phosphate buffer, pH 7.35, then mounted on glycerine gelly and viewed/photographed with a Zeiss Axioskop microscope.

Control preparations were stained with the NADPH-diaphorase procedure after: (i) omission of NADPH, (ii) omission of nitro blue tetrazolium, (iii) replacement of NADPH with NAD. In addition, some sections were pre-incubated with 1 mM
L-NNA (N\text{-nitro-L-arginine}) dissolved in glycine-NaOH buffer, pH 8.5. L-NNA is a specific inhibitor of NOS activity and, consequently, of NO production. Control preparations were not stained, except for \textit{w},M360, that showed persistent labelling in the retina.

**Evaluation of staining intensity**

In order to make objective evaluations, the staining intensity of each area of the optic lobes was determined by scanning colour photographs of sections with a colour scanner (AGFA ARCUS II) and then using Adobe Photoshop 5.0 for computerized image analysis. The staining intensity was expressed as percentage black in the grey scale. All the intensity values obtained from the control reaction omitting NADPH were subtracted from the correspondent intensity values obtained after specific NADPHd reaction. Data are mean ± S.D. of five individuals for each stage (five sections for each individual and fifteen measurements for each section). No significant differences in the staining intensity were seen among individuals. The differences between the neuropiles of the phenotypes were analyzed by the Tukey's HSD Multiple Comparisons test.

**Results**

**Anatomical description of an insect brain**

The head of an insect is formed from a set of fused segmental units, whose precise number, perhaps seven, has proved difficult to establish. Likewise the head contains a set of segmental ganglia, fused to form a brain, called cerebrum. This gangliar complex can be divided into three main areas: proto-, deuto-, and tritocerebro, each one containing nerve centres.

About half of the entire complement of neurones in a Dypteran insect form the optic lobes that serve the compound eyes. In fact, large compound eyes with a facet lens for each \textit{pixel} in the optical image are a distinctive feature of insects; the basic unit of these compound eye is called ommatidium. Beneath these eyes much of the brain is devoted to vision.

The first stages of vision involve point-by-point transformation and coding of image on the retina. This is done by neurones in the optic lobes, whose columnar architecture of almost crystalline regularity beautifully reflects their function. Signals from the eye pass successively through three ganglia: the lamina, the medulla and the lobula. In the outer two ganglia – the lamina and the medulla – the neurones are arranged to form columnar modules, with a set of cells repeated beneath each ommatidium.

The first and smallest ganglion is the lamina; it receives direct inputs from the retina, as well as feedback from the medulla. Signals passed from the lamina to the medulla, finally enter in the lobula.

**Distribution of NOS labelling intensity in the visual system of \textit{or}; \textit{or},wp; \textit{w},M360; \textit{w},Heraklion mutant of \textit{C. capitata}**

The intensity of NADPHd labelling differs in the optic lobes of the four phenotypes and may be sex dependent.

\textit{w},M360 mutant. All the areas of the \textit{w},M360 optic lobes were labelled (Figure 1 a, b, c, d). In particular, a strong intensity of NADPHd staining was found in the retina and in the outer lamina, with respect to the medulla and lobula. However, the immunostaining was stronger in the males. In the male (Figure 1 a, b), the labelling is associated with distinct layers of fibers and varicosities along the whole medulla and in the lamina. In the female (Figure 1 c, d) all the areas of the optic lobe displayed a generally lower staining intensity. In particular, the labelling was higher in the retina and in the lamina whereas the medulla and the lobula were very weakly labelled. Several fibers and varicosities were clearly detectable in the medulla and in the lobula. Both the monopolar cell layer of the outer lamina, and the cell bodies adjacent to the medulla were weakly stained in both sexes.

\textit{or} mutant. The optic lobes of males and females of the \textit{or} strain displayed different NADPHd staining intensities (Figure 2 b, d). All the areas of the optic lobes were strongly labelled in the \textit{or} male (Figure 2 b). The staining was most intense in the retina and in the lamina, whereas the inner neuropiles displayed less NADPHd staining. The photoreceptors showed intense immunostaining and in the lamina and in the outer medulla fibres and varicosities were also clearly labelled. The monopolar cell layer of the outer lamina was weakly stained. On the contrary, in the \textit{or} females (Figure 2 d) only the retina displayed intense labelling; the staining was weak in the lamina and it decreased in the medulla and in the lobula. Low immunostaining was present in the monopolar layer and only sporadically a few fibers and varicosities were labelled in
the outer medulla.

\textit{w,Heraklion mutant}. The \textit{w,Heraklion} optic lobes were stained by NADPHd reaction (Figure 2 a, c) and, like the or and \textit{w,M360} mutants, they displayed different labelling intensities in males (Figure 2 a) and females (Figure 2 c). In particular, lower intensity of staining was found in the medulla and in the lobula, with respect to the retina and the lamina in both sexes. However, the retina and the lamina were strongly labelled in the male. Cell bodies of the monopolar layer were not stained, while in the lamina and outer medulla several fibers and varicosities were labelled.

\textit{or,wp mutant}. The \textit{or,wp} optic lobes did not display different staining intensities in males and females (not shown in figures). The labelling was generally strong, particularly in the retina and in the lamina. The lamina cartridge showed strong labelling in both sexes. In the outer medulla, fibers and varicosities were stained while the monopolar cell layer of the outer lamina was weakly labelled.

\textbf{Intensity values of NADPHd activity of \textit{or}; \textit{or,wp}; \textit{w,M360} and \textit{w,Heraklion mutant optic lobes}}

The variation of NADPHd staining intensity in the optic lobes of males and females of the four strains was considered (Figure 3). The pattern of staining distribution was similar in the optic lobes of the four phenotypes: the retina and lamina generally displayed a higher degree of staining than the medulla and lobula.

The comparison between males and females of
each strain and among all the flies of the same sex (Figure 4) showed that all the areas of the optic lobes were generally strongly labelled in the male of w,M360 strain and, much more, in the or mutants, whereas significant differences in intensities were not evident in the or,wp strain. In the w,Heraklion strain, the female showed a slightly lower staining with respect to the male, but this difference was not highly significant. Moreover, it appeared that, while in the male the intensity of NADPHd staining increased when the eye pigmentation intensified, in the female this relationship was not present.

**Discussion**

**Technical considerations**
Since the enzymatic activity of NOS requires a cofactor NADPH, histochemistry has been used extensively to localize NOS through the reduction of tetrazolium salts to an insoluble formazan reaction product. The large use of NADPHd staining procedure indicates that it is sensitive and specific for several tissues, after careful fixation with formaldehyde (Eldred, 2000) and therefore it is more reliable in the detection of NOS in the animal kingdom. In fact we have recently demonstrated NOS immunoreactivity in *C. capitata* strains (Conforti et al., 2002); however, results required
hard criticism for their evaluations, that was possible since we have already found clear differences of NADPHd reactivity between $w$ and $w^+$ strains (Conforti et al., 2001).

**NOS-dependent NADPHd pattern and role in the male optic lobes**

NOS has been characterized in brain of various insects, such as the honeybee *Apis mellifera*, the fruitfly *Drosophila melanogaster*, the locust *Schistocerca gregaria* and the cricket *Acheta domestica* (Elphick et al., 1996; Müller, 1994) but the centres of visual information processing exhibit very different levels of staining among the species. While in *Drosophila* NADPH diaphorase staining appears to be almost absent in visual neuropiles, *Apis* exhibits an intermediate and *Acheta* and *Schistocerca* a very strong labelling in the visual neuropiles (Müller, 1997). The extreme differences in NADPH diaphorase labelling in the visual system of the various insects suggest a role of the NO system which is more likely characteristic for certain species than a conserved function. Here we have used the NADPHd histochemical technique on a single species, the medfly *C. capitata*, comparing the staining pattern of the optic lobes among four eye colour mutant strains, i.e., $or, or, wp, w, M360$ and $w, Heraklion$. Previously we demonstrated the piv-

Figure 5. Staining intensity values (mean ± S.D.) of NADPHd activity of $or, or, wp, w, M360$ and $w, Heraklion$ mutants optic lobes. Comparison between all the flies of the same sex: while in the males the intensity of NADPHd staining increased when the eye pigmentation intensified, in the females this relationship was not present.
otal role of nitric oxide (NO) in the modulation of the chemosensory information, that seems to be implicated also in visual processes and visually guided behaviour of many insects, using NADPHd staining on two different strains of the medfly *C. capitata* (Conforti et al., 1999).

Our study revealed a peculiar pattern of NADPHd staining in the optic lobes of the species with respect to those previously observed in other insects. These results showed that the staining pattern was similar in the optic lobes of the four strains: the retina and lamina generally displayed a higher degree of staining than the medulla and lobula.

The most remarkable feature was the difference in the labelling intensity of the visual tissue among the four eye colour phenotypes. The optic lobes of the four strains displayed high staining along the whole retina and lamina, whereas the medulla and the lobula showed lower NADPHd labelling. However, the optic lobes of the phenotypes with less pigmented eye (*w*, Heraklion & *w*, M360) displayed a weaker NADPHd staining, at least in the male (see below), than those of the phenotypes with a more intense eye pigmentation.

Independent studies of artificial populations of *D. melanogaster* agree that where wild type and laboratory eye colour mutants are in competition, the mutants are at a distinct disadvantage (Torti et al., 1997). There are observations which point to mating behaviour as a source of disadvantage to the competing mutant flies: it was shown that wild type males are more effective than mutant males when competing for the same females (Geer & Green, 1962). Also in response to light *D. melanogaster* eye colour mutants vary in their ability to orient themselves, compared to wild type eye colour flies (Fingerman, 1952; Spieth & Hsu, 1950). These results emphasized the conclusion that a difference in visual recognition or discrimination accounts for the advantage of one type of male in the light. Moreover, visual activities were tested during the routine maintenance and on mating behaviour of wild type (*w*+) and white eye Heraklion (*w*, Heraklion) mutant of *C. capitata*. These tests showed the higher competitiveness of the wild type males in their mating choice, compared to the white eye mutant males (Conforti et al., 1998; Conforti et al., 1999). Based on these observations, the marked NADPHd staining difference among *or*, *or*, *wp*, *w*, M360 and *w*, Heraklion visual neuropiles, which are specifically devoted to the analysis and elaboration of the visual signals, suggested a role for NO in the regulation of the visual information processes of *C. capitata*. Moreover, the observation of the correlation between the NADPHd staining and eye pigmentation suggested a possible relationship between the intensity of eye pigmentation and the protection of the visual system.

In particular, a neurodegenerative disease, analogous to the human condition retinitis pigmentosa, was analyzed in *Drosophila* (Fortini & Bonini, 2000). Mutant variants of several *Drosophila* photoreceptor cell-specific proteins, including rhodopsin, structural proteins and other factors needed for rhabdomere integrity all cause gradual, light-independent degeneration. Therefore the intensity of eye pigmentation could favour the protection of the visual system.

**Relationship between NOS-dependent NADPHd and sex**

The intensity of NADPHd staining differed in the optic lobes of the four phenotypes and moreover it appeared to differ at least in some strains, between the two sexes. All the areas of the optic lobes were generally strongly labelled in the male, except for the *or*, *wp* mutant, which did not display significant labelling differences between sexes. Considering the pattern of labelling intensity distribution in the four strains, we observed that in the male the intensity of NADPHd staining increased when the eye pigmentation intensified. As previously stated, these results suggested a possible relationship among NOS, eye pigmentation and visual information processes in the *C. capitata* male. Otherwise, the female displayed a very heterogeneous NADPHd labelling and no correlation between eye pigmentation and NOS was shown. In comparison with the males, the females of the *or*; *w*, M360 and *w*, Heraklion strains displayed weak staining, whereas the *or*, *wp* females appeared strongly labelled. In particular, the weak staining of the optic lobes of the *or* mutant, characterized by orange eye coloration, did not allow a correlation between intensity of eye pigmentation and NOS in the female. These results could suggest that nitric oxide production may be dependent upon behavioural differences linked to vision function in the two sexes of *C. capitata*.

Several works (Franceschini et al., 1981; Strausfeld, 1980) analyzed sexual dimorphism in insect vision: some visually guided behaviour pat-
terns differ between the sexes and presumably information processing by nerve cells also differs in males and females. Recent anatomical studies of houseflies Musca domestica have demonstrated that certain visual neurons are present only in males. Also in the blowfly C. erythrocephala another kind of sexual difference in the neural architecture of the visual system was reported. In this fly dimorphism was expressed by differences in the shapes of analogous neurons in males and females, as well as by the presence of some cells in only one sex (Franceschini et al., 1981).

The widely accepted view that NO also plays an essential role also in mating behaviour and that there is an evident relationship between NOS and sexual dimorphism, were confirmed by Hadeishi & Wood’s work (Hadeishi & Woods, 1996) that reported that NO is present in the mating behaviour circuitry of the brain of the male Syrian hamster. Obviously chemosensory and hormonal stimuli are essential for mating in the male Syrian hamster; NO is implicated in the regulation of male sexual behaviour, and NOS is present in the limbic system. Moreover, NO has a pivotal role in male courtship behaviour of the urodele crested newt (Triturus cristatus) but not that of the females (Zerani & Gobbetti, 1996).

A number of experiments also displayed sexual differences in the morphology and the function of neurons in the brains of vertebrates and in particular mammals (Goto & Goto, 2000; Grachev & Apkarian, 2000; Cooke et al., 1998; Raisman & Field, 1973; Nottebohm & Arnold, 1976).

In the human brain sexual dimorphism was linked not only to anatomical but also to chemical features. The orbital frontal cortex and sensorimotor cortex showed gender dependence, with women demonstrating increased metabolite concentrations compared to man (Nopoulos et al., 2000). Data indicate that the morphology and chemistry of brain may be sex-dependent.

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**References**


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