Molecular and behavioural analysis of the 
no-on-transientA (nonA) gene 
of Drosophila virilis.

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by

Susanna Campesan
Department of Genetics
University of Leicester

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"A fly got into the transmitter pod with me that first time... My teleporter turned into a gene splicer, a very good one... You look so pretty... Help me, please, help me."

-BRUNDEFLY

(Seth Brundle to Veronica in The Fly, D.Cronenberg, dir. -Brooksfilms Toronto, 1986)
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ABBREVIATIONS USED

ANOVA  Analysis of variance
A_x  Absorbance at a wavelength of x nm
bp  Base pair
BSA  Bovine serum albumin
°C  Degrees centigrade
Ci  Curie
dATP  Deoxyadenosine triphosphate
dCTP  Deoxycytidine triphosphate
DEPC  Diethyl pyrocarbonate
dGTP  Deoxyguanosine triphosphate
DNA  Deoxyribonucleic acid
DNase  Deoxyribonuclease
dNTP  Deoxynucleoside triphosphate
DTT  Dithiothreitol
dTTP  Deoxythymidine triphosphate
EDTA  Ethilenediaminetetraacetic acid
gr, mg, µg  Grammes, milligrammes, microgrammes
IPTG  Isopropyl-β-D-thiogalactopyranoside
Kb  Kilobase pairs
KV  Kilo Volts
L, ml, µl  Litres, millilitres, microlitres
M, mM, µM  Molar, millimolar, micromolar
min  minutes
mRNA  messenger ribonucleic acid
Myr  Million years
nm  nanometres
OD  Optical density
pmol  picomole
r.p.m.  revolutions per minute
RNA  Ribonucleic acid
RNase  Ribonuclease
s, ms  Seconds, milliseconds
SDS  Sodium dodecyl sulfate
TEMED  N,N,N',N'-tetramethyl-ethylenediamine
tRNA  Transfer ribonucleic acid
U  Unit (of enzymatic activity)
X-gal  5-bromo-4chloro-3indoly1-β-D-galactoside
ABSTRACT

The nonA gene of Drosophila melanogaster influences vision, courtship song and viability, and encodes a protein inferred to bind single-stranded nucleic acids.

This study describes the molecular and functional characterisation of the nonA gene of Drosophila virilis. The main goal of the project was to establish whether the D.virilis nonA gene rescued the viability and behavioural defects of nonA(♂) mutants, and whether it carried species-specific information concerning the lovesong. The D.virilis nonA gene was cloned and sequenced. The overall structure of the D.virilis nonA gene was similar to that of D.melanogaster. Nucleotide and amino acid sequence comparisons revealed a highly diverged and repetitive N-terminus, followed by a conserved C-terminal region. Study of the promoter region highlighted islands of homology containing putative transcription factor binding sites. In addition, the D.virilis gene was found to display an unusual codon usage bias in comparison to other D.virilis or D.melanogaster genes. Finally, protein secondary structure predictions revealed differences between the D.virilis and D.melanogaster polypeptides.

P element-mediated transformation was used to assess the ability of the D.virilis nonA clone to rescue viability, visual and song defects of D.melanogaster flies lacking nonA (nonA(♂)) and an adjacent and partially overlapping lethal gene l(1)l19e. Complete rescue of the viability indicated that the transgene contained an entire functional D.virilis l(1)l19e gene. Visual studies assessing the transformant flies' optomotor response suggested that the D.virilis transgene completely rescued this phenotype. Lovesong analysis initially presented difficulties in distinguishing between incomplete rescue or transfer of species-specific song characteristics. However, extensive statistical analyses suggested that some D.virilis song traits may be conveyed to transformants by the D.virilis nonA gene, but the large differences between the D.virilis and D.melanogaster lovesongs are probably due to the influence of many other genes.
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CHAPTER 1

INTRODUCTION
The ideas of inheritance, particularly of human behavioural characteristics, have engaged human thought for hundreds of years. However it is only in the last century that the basics of what we now understand as the genetics of behaviour, were founded. Francis Galton, whose thinking was profoundly influenced by that of his cousin, Charles Darwin, was the first to address the question of whether human behaviour contained a hereditary component. By using statistical methods, he analysed different physical and mental characteristics in large numbers of families, including the first behavioural studies of twins (Galton, 1869). His conclusions stressed that nature prevails over nurture, yet neither he nor Darwin, were aware of the particulate theory of inheritance which had been published earlier by Mendel. If they had, both could have supported considerably their respective theories, on the inheritance of human mental capacity on one hand, and evolution by natural selection on the other.

The debate concerning the relative contributions of nature and nurture, took a downward turn in the early years of the 20th century, especially in the United States, where by the 1920's the extreme environmentalism of J.B. Watson, totally dominated Psychology. Small pockets of resistance remained, particularly in the laboratory of Tolman, who performed the first selection experiment on 'smart' and 'dumb' rats. His student, Tryon extended this early work, and generated the famous maize bright and maize dull strains of rats (Tryon; 1940). However, the work was not fully published until 20 years after the experiment, such was the prevailing atmosphere. A breath of fresh air finally came from Europe, where the students of the ethologists Lorenz and Tinbergen, began to experiment with the genetic analysis of mice and fruit-flies (reviewed in Manning, 1975). The latter species was particularly important given the enormous genetic literature that Morgan and his students had been amassing on *Drosophila* from about 1915 onwards. So by the early 1960's, the first papers on behavioural genetic analysis had appeared, and there was no turning back.

Welcome as this new development was, most behavioural genetic research focused on studying behavioural traits that could only be described as quantitative. It was unusual that any study, whether it was on the mouse, the rat or the fly, implicated single genes in determining a significant proportion of behavioural variation. This was
to change in 1967, when Benzer advocated mutagenesis as an approach to dissecting nervous system function in *Drosophila* (Benzer, 1967; 1971).

His students enthusiastically took up the challenge, and using clever screening procedures, isolated behavioural mutants which spanned the range from simple reflexes like flight, to more complex phenotypes, such as learning. With other genetic methods, such as mosaic mapping, the anatomical foci of mutant genes could be roughly localised (Benzer, 1973). This procedure, which was originally based on gynandromorphs, could also be extended to map sex-specific behaviour in both male and female flies (Hotta and Benzer, 1976).

By the early to mid-1980's, *Drosophila* behaviour genetics had far surpassed that of its main rival, the mouse. Yet it was only with the advent of transformation technology (Rubin and Spradling, 1982), that the first behavioural gene was cloned (Reddy *et al.*, 1984; Bargiello and Young., 1984), and then demonstrated to encode a behavioural protein by rescue of the mutation (Zehring *et al.*, 1984; Bargiello *et al.*, 1984). The gene in question was *period (per)*, which remains the 'flagship' for the single gene dissection of behaviour, and which determines both circadian locomotor cycles (Konopka and Benzer, 1971), as well as fast one-minute oscillations in the male's courtship song (Kyriacou and Hall, 1980; 1989). Thus the mutate, clone and transform school had its successes, and many genes, initially identified as behavioural mutants, have provided major stories, not necessarily in the study of behaviour, but also in development (ie *sevenless*, Rubin, 1988; Simon *et al.*, 1989), and channel physiology (eg *Shaker*, Wu and Ganetzky, 1992).

More recently, reverse genetics has been brought to bear on behaviour in the fly (Sentry *et al.*, 1994). The second generation enhancer trap using the yeast transcriptional activator GAL4 allows the simultaneous visualisation of expression patterns, and the misexpression of any gene that can be placed under GAL4 control, in the tissue of interest (Kaiser, 1993). The recent misexpression of the *transformer (tra)* gene to feminize male brain structures provides an elegant example of this approach (Ferveur *et al.*, 1995, O'Dell *et al.*, 1995). As can be seen from this brief review into the history of behavioural genetics, the fruitfly looms large in the story, and the sexual
behaviour of *Drosophila*, has been particularly prominent as a model phenotype (Hall, 1994). I shall limit the rest of my introduction to a discussion of sexual behaviour in the fly, and its relevance to behavioural genetics and speciation.

**Neurogenetics of sexual behaviour in Drosophila.**

Courtship in *Drosophila* involves a series of interactive behaviours performed by both male and female. However the male takes the more active part of the courtship which is extensively documented (figure 1.1; for a review, see Hall, 1994). When a mature *Drosophila* male perceives a female, he orients towards her and starts tapping the female's abdomen using his foreleg. If the female moves, he follows her, extends one wing and vibrates it. The wing vibration produces the "lovesong", an acoustic signal which constitutes the most effective and species-specific component of each fly's courtship. Depending on the female's receptiveness, he may go back and repeat the same actions several times. Finally, the male will extend his proboscis and lick the female's genitalia. This is always followed by his first copulation attempt. If the female rejects him, he might stop courting for few seconds. When he resumes courting, the male invariably goes back to orientation and wing vibration stages, and repeats the behavioural sequence. When a sexually mature female has been courted sufficiently, she may accept the male's copulation attempt and allow copulation which has a species-specific duration (in *D.melanogaster* it lasts about 20 minutes).

Courtship behaviours are innate: males isolated as eggs and kept singly until maturity are able to recognize a potential mate and perform effectively all the courtship steps described above (O'Dell and Kaiser, 1997). However, this does not mean that courtship is a completely stereotyped series of actions that cannot be modified. For example, previously mated females elicit relatively little courtship compared to virgin females, and block the male's copulatory attempts by extruding their ovipositor (Connoly and Cook, 1973). Males that have courted fertilized females show reduced levels of courtship when they are subsequently placed in the presence of virgin females (Siegel and Hall, 1979). It seems that the males are not inhibited by rejection
Figure 1.1. The principal elements of courtship in *D. melanogaster*. Original drawings by Barrie Burnet.
behaviour itself, but by aversive olfactory cues emitted by the female (Ackerman and Siegel, 1986). The decreased courtship towards virgin females after exposure to mated females results from counterconditioning, in which the male associates otherwise excitatory pheromones with aversive chemicals simultaneously emitted by the mated female (Zawistowski, 1988). Thus after exposing the male to a fertilized female, appetitive stimuli (female pheromones) either acquire aversive qualities or lose their excitatory capacity.

In a similar experiment, males were presented to virgin females in the presence of quinine, a repellent used as a negative reinforcer for conditioning of *D. melanogaster* (Ackerman and Siegel, 1986). The males exposed to virgin females in the presence of quinine revealed reduced courtship levels in a subsequent test with virgin females in the absence of quinine. However, males placed alone, without virgin females, in the presence of quinine, did not show decreased courtship in subsequent tests with virgin females, in the absence of quinine. This suggests that the exposure to quinine itself has no after-effects, and the observed decline in courtship was due to counter-conditioning by exposure to aversive (quinine) and excitatory (virgin female pheromones) stimuli together.

Studies with learning and memory mutants (for a review, see Davis, 1996) tend to support this scenario. *dunce* (*dnc*, Davis and Kiger, 1981; Byers *et al.*, 1981) and *amnesiac* (*amn*, Folkers, 1982; Feani and Quinn, 1995) mutant males, showed no reduction in courtship levels after simultaneous exposure to virgin females and quinine (Ackerman and Siegel, 1986). Mutant males such as, *amnesiac* and *rutabaga* (Gailey *et al.*, 1984; Dudai, 1985; Livingstone, 1985; Tully and Gergen, 1986; Levin *et al.*, 1992) perform inappropriately vigorous courtship immediately after being trained in the presence of mated females (Hall, 1994). The same is observed in transgenic flies carrying a specific inhibitor of the calcium-calmodulin-dependent protein kinase II (CaMKII), an enzyme known to be involved in the neuronal mechanism of learning and memory (Griffith *et al.*, 1993). Males carrying this construct could be trained in the presence of mated females, in that after one hour of exposure to mated females, they showed the same decrements in courtship shown by wild-type flies. However, the
transformant males failed to retain the effects of the training: when subsequently presented with virgin females, instead of showing inhibition of courtship, they courted vigorously.

Another kind of experience-dependent modification of behaviour can be observed in *Drosophila*. One-day old or younger males tend to elicit vigorous courtship from mature males since they produce an aphrodisiac pheromone which is different from that of virgin females (Jallon, 1984, Vaias et al., 1993). However, when a male is placed together with an immature male, its courting behaviour tends to decrease after a while. Successively, the mature male will completely avoid further courtship of young males (Gailey et al., 1982). It seems that habituation is the key to this type of courtship-related learning, and it is sufficient to expose mature males to the sex-stimulating substances extracted from young males to repress their attraction towards young males (Gailey et al., 1986). All the learning mutants mentioned above are aberrant in this regard, and keep courting young males regardless of their previous experiences with them (Hall, 1986, Hall, 1994).

Apart from stressing the importance of learning and memory as important aspects of courtship behaviours, the experiments described above also underline the fundamental role that pheromones play in mate stimulation and discrimination. It is not surprising then that mutant males that cannot smell normally, such as *smellblind* (poor responses to a variety of volatile compounds) and *olfactoryC* (poor responses to acetates and some alcohols), are also subnormal in their courtship performance since they probably do not sense the females' sex-inducing odours (Tompkins et al., 1980; Tompkins and Hall, 1981).

Male-to-female chemical communication also plays an important role during courtship. The mutation *nerd* (*nrd*, Ferveur and Jallon, 1993) dramatically reduces the levels of 7-tricosene (7-T, the male predominant sex pheromone, Ferveur and Jallon, 1993) synthesized by males. *nerd* males are subnormal in their courtship behaviour and reproductive success. It seems that females find *nerd* male's courtship insufficiently stimulating, and fail to emit the next stimulus in the courtship sequence.
Consequently, *nerd* affects the probability of a given mutant to reach the more advanced stages of courtship behaviour.

Also visual stimuli are extremely important during courtship, especially for the male, which has to be able to track the female. Blind males such as *no-receptor-potentialA* (lacking light-elicited photoreceptor potentials) and *glass* (lacking photoreceptors and having reduced compound eyes) are not able to orient properly towards the female and consequently spend less time courting than wild-type males (Tompkins, 1984). *Optomotor-blind* (*omb*, defective in their optomotor responses, Pflugfelder et al., 1990; Pflugfelder et al., 1992), which are insensitive to horizontally moving stimuli, behave during courtship as if they were totally blind, and court subnormally (Tompkins, 1986, Hall, 1994). Also *white* (*w*) mutants court poorly, since they lack screening pigments that shield the eye from being flooded by too much light and as a result they do not respond properly to moving stimuli (Hall, 1994). If only visual and olfactory cues provided by the females were responsible for stimulating males to court, then males that cannot see or smell would be expected not to be able to court at all. However, double mutants *smellblind; glass* display courtship if they can touch the female, showing that tactile stimuli are also important during courtship (Tompkins, 1984). Since a mature male courts vigorously virgin females, but is not attracted by other mature males, olfactory, visual and tactile stimuli are undoubtedly sex-specific.

Our understanding of the genetic control of sexual differentiation and sexual behaviour in *Drosophila* has progressed enormously since the first studies on sex-determination, started almost 90 years ago by Morgan and his colleagues (see Belote, 1992; Baker, 1989; Hodgkin, 1989; Hodgkin, 1990; Burtis, 1993; McKeown, 1994). The primary determinant of sex in *Drosophila* is the ratio of X chromosomes to sets of autosomes (X:A ratio). One of the first responses to the X:A ratio in females (X:A=1), is the activation of the *Sex-lethal* (*Sxl*) locus (Cline, 1984). *Sxl* autoregulates the splicing of its own pre-mRNA (Bell et al., 1991) and activates *transformer* (*tra*) by regulating the *tra* messenger splicing (Inoue et al., 1990; Boggs et al., 1987; Belote et al., 1989; Sosnowski et al., 1989). The products of the *tra* and
the transformer-2 (tra-2, Belote and Baker, 1982; Mattox and Baker, 1991; Amrein et al., 1988; 1990 Goralski et al., 1989) genes then regulate the splicing of the premRNAs of the doublesex (dsx, Baker and Wolfner, 1988; Hedley and Maniatis, 1991) gene, leading to the production of a female-specific form of the Dsx protein (DsxF) (Tian and Maniatis, 1993; Hoshijima et al., 1991; Hedley and Maniatis, 1991; Burtis and Baker, 1989; Ryner and Baker, 1991). In males (X:A=0.5) Sxl and tra do not make functional products, and dsx pre-mRNA is spliced in its default manner, resulting in a male-specific Dsx protein (DsxM) which differs from the female-specific protein at its carboxy-terminus. The DsxF and DsxM proteins function as sex-specific transcription factors leading respectively to female and male differentiation.

Using temperature-sensitive diplo-X mutants (Sxlmut1,fm#3/Sxlfm#7,M#1), which externally looked like males, it was shown that the Sxl gene must function normally during the pupal period or the first few days of adult life for a fly to elicit vigorous courtship (Tompkins, 1984b; 1986), suggesting that the female-specific Sxl transcript is essential for the production of the female pheromone, 7,11-heptacosadiene (HCD) (Tompkins et al., 1980; Tompkins et al., 1981; Jallon and Cobb, 1987; Ferveur and Sureau, 1996; Ferveur et al., 1996), and the repression of the male inhibitory pheromone 7-tricosene (7T) synthesis (Jallon, 1984; Ferveur and Sureau, 1996; Ferveur and Jallon, 1996). Haplo-X flies carrying a gain-of-function Sxl mutation (which leads to ectopic expression of female-specific Sxl functions) elicit anomalously high levels of interest from normal males, and also perform less courtship than wild-type males, supporting the idea that Sxl controls the production of female pheromones (Tompkins and McRobert, 1989).

The synthesis of female-specific pheromones is regulated by Sxl via tra and tra-2, since tra and tra-2 mutations make diplo-X flies unattractive to normal males (Baker and Ridge, 1980, McRobert and Tompkins, 1985). In males tra is not functional, and it seems that the gene intersex (ix) may play a role in the production of sex-specific pheromones. In fact ix mutations, although they have no morphological effects on flies, make haplo-X flies attractive to normal males but not as attractive as females, indicating that these mutants are either producing some HCD (stimulating pheromone)
or not producing enough 7T (inhibitory pheromone) (Baker and Ridge, 1980; McRobert and Tompkins, 1985; Jallon, 1984; Scott, 1986; Scott et al., 1988; Scott and Jackson, 1988).

There are no doubts that Sxl, tra and tra-2 control all aspects of somatic, neuronal and behavioural sex-specific differentiations. XX flies homozygous for tra mutations not only look like males but they also behave like males, being able to court females vigorously and to sing to them in an essentially normal manner (Kyriacou and Hall, 1980; Kulkarni et al., 1988; Rendahl et al., 1992; Bernstein et al., 1992). A conditional mutation of tra-2 allowed Belote and Baker (1987) to "turn off" the gene at different stages of the flies' life cycle simply by raising the temperature. It was found that inactivation of the gene in diplo-X pupae or even young adults could lead to the appearance of male courtship behaviour. This result suggested that the product of the tra-2 gene must be functional in adults in order to maintain a female-specific behavioural programme. In addition, it indicated that the adult nervous system retains some functional plasticity in respect to the sex-specificity of courtship (Belote and Baker, 1987).

In contrast, dsx does not control all aspects of sex-specific somatic differentiation. For example, tra and tra-2, but not dsx, control the development of a pair of bilaterally symmetrical male-specific abdominal muscles called Muscles of Lawrence (MOLs, Lawrence and Johnston, 1986; Taylor, 1992). It seems that the development of these muscles depends on the sex of the neurons that innervate them (Lawrence and Johnston, 1986; Currie and Bate, 1995). In addition, constitutive expression of the male form of the Dsx protein transforms females to morphological males, but these flies do not court (Taylor et al., 1994). These data suggest that a previously unknown branch of the sex-determination hierarchy immediately downstream of tra and tra-2 may control sex-specific neuronal development and behaviour. It has to be noted however that dsx is involved in the development of one set of neurons (Taylor and Truman, 1992) and one particular aspect of the courtship song (Villella and Hall, 1996).
The best candidate for acting in this new neuro-behavioural branch of the sex-determination pathway seems to be the fruitless (fru) gene (Hall, 1978; Gailey et al., 1991; Taylor, 1992). The original mutation at the fru locus caused males to court both females and males and to be courted by wild-type males. The most striking behaviour of the fru mutant males is that they form long "courtship-chains" in which every individual is simultaneously courting and being courted (Gill, 1963; Gailey and Hall, 1989). fru mutants are never able to copulate (Hall, 1978), and, in addition, they fail to develop a normal MOL (Gailey et al., 1991; Taylor and Knittel, 1995). Ito et al. (1996) and Ryner et al. (1996) have recently cloned fru. Both groups found that fru is a large and transcriptionally complex locus, carrying at its 5' untranslated region three copies of the Tra-Tra-2 recognition motif. The predicted translation products include a male-specific and a female-specific form and show similarities to the BTB-ZF family of transcription factors (Albagli et al., 1995). These results, and the fact that fru is expressed in a population of brain cells, including those of the antennal lobe, which have been suggested to be involved in determination of sexual orientation (see later), support an intriguing scenario in flies in which sexual orientation is controlled by the same hierarchy of genes that govern all other aspects of sex.

A second tra-dependent but dsx-independent locus controlling sexual behaviour has been recently isolated (Finley et al., 1997). Male mutants of this gene, named dissatisfaction (dsf), are bisexual and although they attempt to copulate, they do so with difficulty due to a defect in their abdominal curling mechanism. Interestingly, dsf mutations cause abnormal female courtship in that they resist males and fail to lay mature eggs. The defective abdominal curling in males and the inability to lay eggs in females seems to be due to abnormal differentiation of sex-specific abdominal neurons. Because of several features similar to those observed in fru mutants (homosexual courtship, abdominal curling defects) dsf could be the first fru-dependent gene in the putative neuro-behavioural sexual cascade. However fundamental behavioural and neurological differences exist between dsf and fru, such as the courtship and fertility abnormalities observed in dsf females, and the fact the dsf males produce normal MOLs. Therefore Finley et al. (1997) suggest that dsf and fru might be acting in two separate pathways.
Various areas of the nervous system have been identified as foci for determined aspects of courtship behaviour (reviewed in Greenspan, 1995 and Hall, 1994). In the 1970s, J.C. Hall was one of the first researchers that used gynandromorphs (part female-part male) flies to try to find the areas in the brain that need to be either male or female in order to achieve a particular step in the courtship pattern. He found that, in order to be able to perform the early steps of courtship, such as orientation, tapping, following and wing exention, at least one side of the dorsal posterior brain has to be haplo-X (Hall, 1977; 1979). For two of the more advanced behaviours, such as wing vibration and attempted copulation, male tissue is required in the thoracic ganglia. In particular, only the left or right ventral nervous system needs to be male for normal courtship song output from both wings (Schilcher and Hall, 1979). Another advanced courtship behaviour, licking, has its focus in the dorsal posterior brain, but in this case the focus is submissive, since both sides of this part of the brain must be haplo-X in order to achieve proboscis extension (Hall, 1979). Tompkins and Hall (1983) also identified the brain sites controlling female receptivity. They found a group of cells in the dorsal anterior brain that have to be bilaterally diplo-X in order to ensure receptivity to copulation.

More recently, Ferveur et al. (1995) have found the neural foci that control sexual orientation in Drosophila. Using the Gal4 system (Brand and Perrimon, 1993), they drove ectopic expression of the tra gene in order to feminize determined regions of the male brain. They discovered that males that were feminized in a portion of the antennal lobes, or in a subset of the corpora peduncolata (mushroom bodies), courted males and females indiscriminately. These two regions of the brain are known to be involved in the detection and processing of olfactory stimuli (Carlson, 1991; Stocker, 1994; Stocker and Gendre, 1989; Heisenberg et al., 1985; de Belle and Heisenberg, 1994). Therefore these partially feminized strains might be lacking male-specific centres that enable them to detect antiaphrodisiac pheromones produced by normal males. Alternatively, the feminization of the above structures might enable the affected flies to detect sex-stimulating male pheromones. However, since these flies' brains are still mainly male, their behavioural output remains typically male-like.
An experiment similar to that of Ferveur et al. (1995) involved ectopic expression of tra in well defined subsets of cells within the mushroom bodies (known as Kenyon cells), and mapped with even greater accuracy several sites controlling sexual orientation (O'Dell et al., 1995). Sexual orientation has also been altered by the ectopic general expression of the white (w) gene. Activation of a \textit{hsp-70/miniwhite} transgene in mature males resulted in a drastic change in their sexual preference since they courted vigorously other males and mainly ignored females. The cause of this unexpected phenomenon is not known, however it could have something to do with lower levels of serotonin induced by the misexpression of the \textit{w} gene (Zhang and Odenwald, 1995). Interestingly, it was also noted that wild-type mature males placed in a vigorous homosexual courtship scenario started to court other males too, demonstrating that also environmental factors can influence sexual orientation in flies (Zhang and Odenwald, 1995).

The role of courtship song.

Since it was noted that the wing-vibration sounds emitted by the male during courtship are species-specific (Waldron, 1964), the courtship songs of more than a hundred species of \textit{Drosophila} have been characterized (Ewing and Bennet-Clark, 1968; 1969; Ewing, 1970; 1979; Miller et al., 1975; Chang and Miller, 1978; Lakovaara and Hoikkala, 1979; Ikeda et al., 1980; Cowling and Burnet, 1981; Ewing and Miyan, 1986).

The \textit{D.melanogaster} courtship song consists of two main components: pulse song and sine song (figure 1.2). The former is composed by trains of sound pulses ranging from a minimum of two to a maximum of 50 pulses per train. Each pulse usually consists of one to three cycles (Kulkami and Hall, 1987) and lasts about 5-10 ms. Successive pulses are separated by intervals (interpulse-intervals, IPIs) with a mean value of 35 ms (Ewing and Bennet-Clark, 1968). In 1980, Kyriacou and Hall noted that, in \textit{D.melanogaster}, the IPI values oscillate sinusoidally between 28 ms and 40 ms, with a period of 55-60 s. Finally, the sine song is composed of humming
Figure 1.2. Burst of *D. melanogaster* courtship song, including sine (hum) and pulse components.
sounds of variable length with fundamental frequencies of about 160-170 Hz (Cowling and Burnet, 1981; Wheeler et al., 1989). A continuous period of song constitute what Cowling and Burnet (1981) called a phrase. Phrases, which are of variable length, usually contain both pulse and sine song in no fixed order, however phrases containing only one or other of these components are often observed (Wilson et al., 1976).

Cowling and Burnet (1981) analyzed and compared the courtship behaviour and songs of six sibling species in the *D. melanogaster* subgroup: *D. melanogaster, D. mauritiana, D. simulans, D. yakuba, D. teissieri* and *D. erecta*. They found clear differences both in behaviour and lovesongs characteristics between these six species, with the only exception being *D. mauritiana*. The courtship song produced by this species in fact has a mean sine song frequency (SSF) and intrapulse frequency (IPF) close to that of *D. simulans*, while its IPI values resemble those of *D. melanogaster*. The ambiguity of the *D. mauritiana* song can be explained assuming that species-specific song differences have a function in allowing recognition of conspecific flies and thus maintain sexual isolation between different species. Sexual isolation mechanisms need to be stronger when two species have the potential to interbreed, for example when they are sympatric (Fisher, 1930; Dobzhansky, 1940; Welbergen et al., 1987). Reinforcement is the process by which natural selection strengthens sexual isolation between incipient species, reducing the frequency of maladaptive hybridization and hence completing reproductive isolation (Noor, 1995). Since *D. mauritiana* is found only on Mauritius, where no other species of the *D. melanogaster* subgroup are found, there is no need for its song to be very different from that of the other species. On the contrary, *D. melanogaster* and *D. simulans* live in sympatry, therefore their songs may differ considerably in order to maintain sexual isolation between the two species. Cowling and Burnet found that *D. simulans* has a pulse song with a mean IPI value of 55 ms. However, other workers have reported a value of 48 ms (Ewing and Bennet-Clark, 1968). It seems that the IPI values of *D. simulans* are in general more variable than those of *D. melanogaster* (Kawanishi and Watanabe, 1980). The IPI rhythm in this species was found to be 35-40 s (Kyriacou and Hall, 1980; 1986).
In the *D.virilis* group species it seems that differences in courtship songs might also contribute to maintaining sexual isolation between sympatric species (Hoikkala *et al.*, 1982). The *montana* phylad species (*D.kanekoi, D.littoralis, D.ezoana, D.borealis, D.flavomontana, D.lacicola, D.montana montana*, and *D.montana ovivorum*) live in sympatry and are very effectively sexually isolated (Patterston and Stone, 1952; Stone *et al.*, 1960; Throckmorton, 1982). As expected, these species emit very distinct courtship sounds. In contrast, the *virilis* phylad species (*D.americana americana, D.americana texana, D.novamericana, D.lummei, and D.virilis*) are not very effectively sexually isolated and are usually allopatric. The songs of these species are very similar to each other and consist of dense pulse trains with very short IPIs, the pulse trains of *D.virilis* being shorter than those of the rest of the species (Hoikkala *et al.*, 1982).

In their study, Cowling and Burnet (1981) tried to map the song differences found between the six species of the *D.melanogaster* subgroup by generating interspecific hybrids (when possible) between any pair of the species analyzed. The characteristics of the hybrid songs suggested that the genes controlling the IPI were located on the autosomes. Sine song frequency seems to be determined mainly by autosomal factors, but the ability to generate sine song appears to be sex-linked. *D.yakuba* males do not produce any sine component in their lovesongs, and interspecific hybrid males from a cross *D.yakuba* females x *D.mauritiana* males also show absence of this element in their courtship songs (Cowling and Burnet, 1981), while hybrid males from the reciprocal cross (females *D.mauritiana* x males *D.yakuba*) always produce sine song (Demetriades, 1997). Finally, the control of the mean intrapulse frequency in the six sibling species of the *D.melanogaster* subgroup seems to be largely autosomal (Cowling and Burnet, 1981).

Similar studies performed on the songs of *D.persimilis* and *D.pseudoobscura* revealed that the genes controlling the qualitative aspects of the songs, that is whether the overall pattern is the *pseudoobscura* or *persimilis* type, are located on the X chromosome. However in these two species the IPI length seems to be controlled by autosomal genes (Ewing, 1969). In the *D.virilis* phylad, differences in several traits of
the lovesong (such as pulse train length (PTL), number of pulses per train (PN), IPI length, pulse length (PL), number of cycles in a pulse (CPP), and length of a cycle (CL)) appear to be determined by autosomal genes (Hoikkala and Lumme, 1987). In crosses between *D. virilis* and *D. littoralis* or *D. flavomontana* (both species belonging to the *montana* phylad) reciprocal hybrids differed from each other in PTL, IPI, PL and CPP, indicating a sex-linked or cytoplasmic inheritance. In the backcrosses between *D. virilis* and *D. littoralis* the involvement of the X chromosome was found to be decisive. Hoikkala and Lumme (1987) propose therefore that a major change on the X-chromosome allowing variation in IPI occurred during the separation of the two *virilis* group phylads (*virilis* and *montana*), the long IPI allowing variation also in PL and CN. The evolution of the sounds in the *virilis* phylad seems to have gone towards longer and denser pulse trains, whereas in the *montana* phylad, the sounds have evolved in various directions.

In addition to its function as a species-discriminator, the courtship song has an undoubted role in stimulating the female to mate. Von Schilcher (1976a; b) found that *D. melanogaster* females prestimulated with sine song, before being placed with males, showed reduced copulation latencies compared to females which had not been pre-stimulated. However, a pulse song with a constant IPI of 34 ms did not have the same effect in similar experiments. Nevertheless, the same pulse song could speed mating latencies of females in the presence of mute, wingless males. Von Schilcher suggested that females might summate sine songs, increasing their receptivity to a threshold level. On the contrary, pulse song might act immediately as a trigger to mating.

Kyriacou and Hall (1982; 1986) studied the receptivity of *D. melanogaster* and *D. simulans* females which had been stimulated with artificially generated pulse songs. The pulse songs they tested were more elaborate than those used by Von Schilcher, since they incorporated all the possible combinations of species-specific 35 ms IPI, 48 ms IPI, 55 s and 35 s oscillation periods. It was found that *D. melanogaster* females responded best to stimulation with a song with *melanogaster*-like characteristics, 35 ms IPIs with a superimposed rhythm of 55 s, while *D. simulans* females preferred a *simulans*-like song with 48 ms IPI and a 35 s rhythm. Therefore pulse song can
stimulate females when both the IPI oscillation and IPI values are species-specific. Greenacre et al. (1993) subsequently observed that *D.melanogaster* females actively discriminated against the *D.simulans* heterospecific song: they did not necessarily prefer a 55 s song cycle, since a 80 s rhythm stimulated them as effectively. However they did reject a 35s cycle *simulans*-like song.

*D.melanogaster* females can also be pre-stimulated by a species-specific 55s rhythmic-35 ms IPI pulse song (Kyriacou and Hall, 1984). Pre-stimulation effects seem to involve mechanism of information storage and retrivial of information (the conspecific IPI cycles). In fact memory and learning mutants such as *dunce, rutabaga, amnesiac* and CAM-kinase depleted females show minimal or short lived effects of song pre-stimulation (Kyriacou and Hall, 1984; Griffith et al., 1993).

The stimulation and pre-stimulation experiments (Kyriacou and Hall, 1982; 1984; 1986) highlight the importance of the IPI cycle component of the pulse song as a means of discriminating conspecific mates. However, it has been argued that courtship encounters in the wild might be very brief, not allowing enough time for the females to "calculate" the IPI oscillations (Ewing and Ewing, 1987). Nevertheless it is possible that females recognize the rate of variation of the IPI around a certain value. Indeed, Alt et al. (1998) have recently suggested that females may recognize the "nature" of a cycle from the IPIs generated in less than a complete 60 s oscillation. Alternatively, females might possess a preferred IPI value: by rhythmically changing its IPI values, a male increases his chances of producing the right IPI values for stimulating a large number of females (Kyriacou and Hall, 1982).

Thus IPI seems to be one of the most important song parameters involved in sexual stimulation and conspecific recognition during courtship. Evidence from a study of the *D.auraria* complex supports this idea. The species of the *auraria* complex are mostly sympatric and show strong premating isolation. Species-specific courtship song and failure in copulating displayed by "mute" males suggest that the courtship song is particularly important in maintaining sexual isolation in this complex. Interestingly, the IPI is the only song character which is consistent species-specific
among the several courship elements studied in these species (Tomaru and Oguma, 1994a; 1994b).

Analysis of the songs produced by reciprocal *D.melanogaster/D.simulans* interspecific hybrid males allowed Kyriacou and Hall (1986) to show that mean IPI values are controlled by autosomes, whereas IPI cycles are determined by the X chromosome. This was consistent with the X chromosome *per* locus determining species-specific lovesong cycle variation, because Kyriacou and Hall (1980; 1989) noticed that three mutations of the *period (per)* locus which shortened (*per*\(^s\)), lengthened (*per*\(^L\)) and obliterated (*per*\(^0\)) the circadian rhythms of *D.melanogaster* (Konopka and Benzer, 1971), had parallel effects on the song IPI cycles. The *per*\(^s\) mutant had an IPI rhythm of \(\sim 40\) s, *per*\(^L\) had a cycle of \(\sim 80\) s, and *per*\(^0\) seemed not to have any evident IPI cycles. Furthermore, deletion of a repetitive Threonine-Glycine (TG) region from an otherwise wild-type *D.melanogaster per* transgene produced IPI cycles of \(- 40\) s in *per*\(^0\) transformant flies (Yu *et al.*, 1987). These results supported the idea that the *per* locus, and more specifically its TG region, could carry species-specific information related to the IPI cycling. This was confirmed by the work of Wheeler *et al.* (1991), who generated several chimeric constructs in which the TG repeat, together with some flanking DNA, were interchanged between *D.simulans* and *D.melanogaster per* genes. Transformant *per*\(^0\) males carrying the *D.melanogaster per* gene, but a *D.simulans* TG region, produced songs with the characteristic *D.simulans* \(- 40\) s IPI cycles. Transformant males carrying the *D.simulans per* gene, but a TG region from *D.melanogaster*, sang with *melanogaster*-like IPI cycles of \(\sim 60\) s.

**Courtship song mutants.**

Only a few loci have been identified, along with *per*, which alter the male's courtship song. The *cacophony (cac)* mutant was isolated through mutagenesis, by identifying flies which took too long to mate (von Schilcher, 1977; Kulkarni and Hall, 1987). The characteristics of a *cac* courtship song are polycyclic pulses of high amplitude and longer than normal IPIs (on average 45 ms instead of the wild-type 35
ms; figure 1.3). In spite of such grossly different pulse song patterns, the sine song produced by *cac* mutants is essentially indistinguishable from that of wild-type males. *cac* males and females are observed to fly normally with normal wing-beat frequency, thus the mutation appears to affect only the physiology of pulse song generation (Kulkarni and Hall, 1987). However it must be noted that *cac* males are less successful in mating than wild-type flies, even after wing amputation, which suggest that *cac* mutants bear some additional defects that reduce their mating ability (Kulkarni and Hall, 1987). Such defects were subsequently mapped to a separate locus after it was noted that recombinant *sn^3 cac* males, in spite of their abnormal polycyclic song, initiated mating much more quickly than did males of the original *cac* stock (Kulkarni and Hall, 1987).

Another genuine courtship song mutation identified, the *dissonance* mutation (Kulkarni et al., 1988), turned out to be an allele of the *no-on-transientA (nonA)* gene (Rendahl et al., 1992), and will be described in detail in the next section.

Yokokura et al. (1995) identified *croaker (cro)* mutants which initiate courtship after a long delay, if compared to wild-type males, and produce polycyclic pulse songs with prolonged IPIs, similar to those of *cac* males. Because flying ability is significantly impaired in *cro* mutants, a general defect in the motor system was suspected. However this possibility was excluded after several tests that confirmed that muscle function is normal in these mutants. Additionally, electroretinogram (ERG) recordings (see later) and gustatory behaviour essays of *cro* mutants appeared normal. These findings support the notion that *cro* flies have a defect confined to the CNS, leaving the sensory input and the motor output systems intact. The fact that the *cro* mutation also reduces female attractiveness without changing pheromonal production (D.Yamamoto and J-M Jallon, unpublished results) supports the idea that it causes a central defect.

It should be mentioned that the two mutants *fru* and *dsx*, known for their effects on sexual behaviour and sexual determination, also generate defective courtship songs. The original *fru^1* mutants (Gill, 1963) and the secondly isolated *fru^2* (Moses, 1989), have been observed to produce a pulse song with aberrantly long IPIs (Wheeler
Figure 1.3. Pulse song of mutant and normal males. A: a wild-type (Canton S) male's typical bout of pulse song. B: a moderately aberrant bout produced by a nonA^{lss} male; note that this train starts with wild-type-like pulses. C: wildly aberrant bursts from the same male as in B, though only the latter portion of this pulse train is shown (i.e. the arrow points to pulse No. 11; this train, like that in B, commenced with normal-appearing pulses). D: a typical cac pulse train. The total time for each plot is ca. 400 ms. Picture taken from Kulkarni et al., (1988).
et al., 1989, Villella et al., 1997). Moreover, the other two fru mutations discovered so far (fru$^3$ and fru$^4$, Castrillon et al., 1993) have been shown to completely suppress pulse song (Villella et al., 1997). dsx mutant males were found to produce essentially normal pulse song bouts, but to completely lack the sine song component (Villella and Hall, 1996).

The no-on-transientA (nonA) gene.

The sex-linked no-on-transientA\textsuperscript{dissonance} (nonA\textsuperscript{dis}) mutation was found directly by screening for flies that produced aberrant songs (Kulkarni et al, 1988). As in the case of per, and the Dmca1A-encoding locus (of which cac is an allele, Smith et al., 1996), it was found that the nonA locus actually affected more than one aspect of the fly's behaviour (Kulkarni et al, 1988; Smith et al., 1996; Rendahl et al., 1992; Smith et al., in press). nonA is involved in at least two distinct and apparently unrelated behaviours: the fly's vision and the male's courtship song.

The first nonA mutants were isolated by the group lead by Benzer (Hotta and Benzer, 1970; Pak et al., 1970), in their screens for flies defective in phototaxis (the tendency to follow the light). The same mutants also produced an altered electroretinogram (ERG), which is an extracellular measure of the light-generated currents in the eyes and brain of the insect (Heisenberg, 1971b; Pak, 1975). The ERG of nonA mutants lacked both light-on and light-off transient spikes, whereas the maintained depolarization which is produced by the sum of the photoreceptors generation potentials, was apparently normal (figure 1.4). The on and off-transients are believed to arise from the large monopolar cells L1 and L2 in the lamina, which is the first optic ganglion postsynaptic to the photoreceptor cells (Pak, 1975; Coombe, 1986). Thus the lack of transient spikes could mean a defect in the transmission from the photoreceptors to the neurons of the lamina, or the inability of the the neurons of the lamina to respond to information from the photoreceptors. Furthermore nonA mutants show several other visual defects which suggest an even more central defect in the nervous system. For example, nonA mutants are impaired in their optomotor
Figure 1.4. A: typical ERG of a wild-type fly. B: ERG recorded from a mutant nonA<sup>dist</sup> fly; note the complete absence of light-on and light-off transient spikes. The vertical scale bar represents 5mV, the horizontal indicates 3s. On the right, a schematic representation of the dipteran compound eye and optic ganglia.
response (Heisenberg, 1972; Heisenberg and Gotz, 1975). The optomotor test in *Drosophila* measures the tendency of the fly to follow movement in its environment. If a fly is located in the middle of a plexiglass rotating drum which has vertical black and white stripes, the insect tends to follow the direction of the stripes. *nonA* mutants show different degrees of impairment in this test (Heisenberg, 1972). In particular, *nonA*\(^{H2}\) flies seem to respond better to regressive motion (back to front) than to progressive motion (front-to-back, Heisenberg, 1972).

The *nonA\(^{disS}\)* mutation was isolated, as mentioned above, for its effect on the male's courtship song (Kulkarni *et al.*, 1988). In *nonA\(^{disS}\)* the pulse song is specifically affected in that pulse trains start out normally, but as the train proceeds, pulses become increasingly polycyclic and very high in amplitude (figure 1.3). However, short trains in *nonA\(^{disS}\)* (containing less than 6 pulses) are usually completely normal. An extreme phenotype is displayed by *tra* mutant diplo-X flies (which look and behave as males) carrying the *nonA\(^{disS}\)* mutation over a deletion of the *nonA* locus: the pulses produced by these flies can bear up to 100 cycles per pulse (Rendahl *et al.*, 1992). Further analysis of *nonA\(^{disS}\)* mutants revealed that these flies are impaired also in their visual behaviour, and they display exactly the same defects found in all the other *nonA* visual-only mutations (Kulkarni *et al.*, 1988). As a consequence of their visual impairment, *nonA\(^{disS}\)* males frequently lose contact with the females they are courting, and sometimes are observed to sing to empty spaces. However they do not court without the presence of a virgin female, since they do not sing if isolated in fresh unused courtship chambers, nor do they produce any wing vibration when placed in mating chambers that have previously hosted virgin females (Kulkarni *et al.*, 1988). The aberrant song of *nonA\(^{disS}\)* males is not related to the fact that they continuously lose contact with the female, since blind males (such as *norpA*\(^{P24}\)) produce perfectly normal lovesongs (Kulkarni *et al.*, 1988).

*nonA* alleles are all recessive and display a complex pattern of complementation. *nonA\(^{disS}\)*, when heterozygous with any of the visual-only mutations (*nonA*\(^{H2}\), *nonA*\(^{P14}\), *nonA*\(^{P49}\), *nonA*\(^{P60}\)), in a *tra* genetic background (to generate diplo-X males), leads to absence of ERG transient spikes and lack of optomotor responses.
Only two of the visual-only alleles \((\text{non}A^{H2} \text{ and } \text{non}A^{P49})\) can clearly complement \(\text{non}A^{\text{diss}}\) song defects. Heterozygous \(\text{non}A^{\text{diss}}/\text{non}A^{P14}\) "pseudomales" (\(\text{tra}\) diplo-X flies) produce songs intermediate between the wild-type and \(\text{non}A^{\text{diss}}\) (Rendahl et al., 1992). On the other hand all the \(\text{non}A\) alleles, except \(\text{non}A^{\text{diss}}\), produce perfectly normal courtship songs (Kulkarni et al., 1988). The fact that \(\text{non}A^{P14}\) males are closer to wild-type in their singing behaviour than diplo-X \(\text{non}A^{\text{diss}}/\text{non}A^{P14}\) heterozygotes is probably due to dosage compensation of the expression levels for X-linked genes (Jaffe and Laird, 1986). Thus one dose of the \(\text{non}A^{P14}\) allele in a haplo-X fly would produce higher level of gene product than one dose of the same allele in a diplo-X individual. This observation also suggests that \(\text{non}A^{\text{diss}}\) is, in terms of biological functions, more similar to a null mutation than is \(\text{non}A^{P14}\).

The \(\text{non}A\) locus has been mapped to the cytological position 14C1-2 by \textit{in-situ} hybridization (Jones and Rubin, 1990, Besser et al., 1990). The gene was cloned independently and almost contemporaneously by two groups (Jones and Rubin, 1990, Besser et al., 1990). Jones and Rubin (1990) used P-element mediated transformation to show that two overlapping 11 and 16 Kb fragments of genomic DNA can rescue all the \(\text{non}A\) visual defects. Subsequently, Rendahl et al. (1992) demonstrated that the same two fragments can also rescue all the phenotypes caused by the \textit{diss} \((\text{non}A^{\text{diss}})\) mutation, thus undoubtedly confirming that \textit{diss} is an allele of the \(\text{non}A\) locus. Besser et al. (1990) identified the NONA protein among proteins extracted from polytene chromosomes of \textit{Drosophila} larvae, and used an antibody against it to detect a \(\text{non}A\) cDNA from an expression library.

The 9 Kb DNA fragment encoding the \(\text{non}A\) gene has been sequenced (Jones and Rubin, 1990, Besser et al., 1990). Conceptual translation of the locus predicts a single open reading frame of 700 amino acids. The NONA protein presents some interesting features (see figure 1.5): its N-terminal third is very rich in glycine, asparagine and glutamine. The C-terminal two thirds of the protein is rich in charged residues (Jones and Rubin, 1990, Besser et al., 1990). The central segment of the protein contains two tandemly repeated 80 amino acid motifs common to a family of proteins known for their ability to bind RNA. This 80 amino acid sequence is named
Figure 1.5. Schematic representation of the putative functional domains of the NONA, NONO, PSF and p54 proteins. RRM = RNA-binding domains (RRM1 and RRM2). +/- = charged domain. HTH = helix-turn-helix motif. G, N, Q = Glycine, Asparagine and Glutamine-rich region. G = poly-Glycine repeat. Q = Glutamine-rich region. P = Proline-rich region. P, Q = Proline and Glutamine-rich region.
RRM (RNA recognition motif, known also as RNP or RBD motif, Siomi and Dreyfuss, 1997) and is moderately conserved from yeast to man (Bandziulis, 1989). The most conserved segments within such domain are one octapeptide (RNP-1) and one hexapeptide (RNP-2). The family containing the RNP motif includes small nuclear ribonucleoproteins (snRNPs) known for their role in RNA splicing, heterogeneous ribonucleoproteins (hnRNPs), and several Drosophila gene products involved in the fly's development, such as embryonic lethal abnormal visual system (elav, Robinow et al., 1988), or in sex determination, such as Sex-lethal (Sxl, Bell et al., 1988) and transformer-2 (tra-2, reviewed in Baker, 1989).

Three mammalian proteins are related to the product of nonA in a region containing the two RRM domains plus a downstream ~ 110 amino acid peptide sequence: the product of non-O (Yang et al., 1993), the splicing factor PSF (Patton et al., 1993), and the HeLa protein p54nrβ (Dong et al., 1993). The NONO protein is known to be able to bind specifically to an octamer of double-stranded DNA through two helix-turn-helix domains, and RNA or single-stranded DNA via its RNP domains. The sequence of NONO presents two other regions of interest: a glutamine rich region and a proline rich one (figure 1.5). These regions, like the glycine rich domain found in NONA, are believed to be functionally important in protein-protein interactions (Patton et al., 1993). So far, three different forms of NONO have been isolated: an ubiquitous form expressed in all the mammalian tissues examined, a lymphoid-specific form, and a retinal/hypothalamic-specific one (Yang et al., 1993). The mammalian splicing factor PSF is known to be required for the early stages of spliceosome formation in vitro. This protein displays a marked preference for uridine stretches at the branch-point of mammalian introns (Patton et al., 1993). The function of the HeLa protein p54nrβ is still unknown, however, preliminary analyses suggest that this protein could bind to RNA in vitro (Dong et al., 1993). Interestingly, both PSF and p54nrβ are unusually rich in Gln and Pro at their N- termini. The ~310 amino acid sequence homologous between nonA and these three mammalian proteins has been named the DBHS domain, which stands for Drosophila behaviour and human splicing domain (Dong et al., 1993). More recently, an additional gene product has been found to share homology with nonA in a region overlapping the DBHS domain. This protein is
the predicted translation product of a novel Drosophila P1 clone (M. Palazzolo, unpublished material), which shares a 99% identity with nonA in the DBHS region (Rendahl et al., 1996).

RNA-binding proteins can perform a broad variety of functions, such as capping, polyadenylation, constitutive or alternative splicing of pre-mRNAs, transport in the cytoplasm or stabilization of mRNAs (Rendahl et al., 1996; Graumann and Marahiel, 1996). Several RRM-containing proteins of Drosophila have been identified and their role in mRNA splicing actually demonstrated (review: Rio, 1993). Many of these proteins are hypothesized to be specifically required for mRNA splicing in the nervous system (reviewed by Yao et al., 1993). Examples are the product of the gene elav, which is expressed at early stages of neuronal differentiation (Robinow and White, 1988; Robinow and White, 1991); couch potato (cpo), expressed in precursor and differentiated cells of the PNS (Bellen, et al., 1992); musashi (msi), which is expressed in neurons of the PNS and CNS and is involved in adult organ sensory development (Nakamura et al., 1994). Because of its homology to such proteins, and given its pleiotropic behavioural functions, it is believed that the product of nonA might be implicated in some aspects of processing of nervous system-specific mRNAs (Rendahl et al., 1996).

Pre-mRNAs known to undergo regulated splicing in the Drosophila nervous system, such as those transcribed from the sodium, potassium, and calcium channel genes, and pre-mRNAs from loci required for wild-type visual or courtship behaviours, are all possible targets of NONA splicing functions. For example, alternatively spliced transcripts obtained from the Shaker (Sh) potassium channel locus are localized in the dorsal longitudinal muscles (DLMs) and retina of late pupae and adults (Mottes et al., 1993). Several mRNAs encoding voltage-gated sodium-channel polypeptides are transcribed from the para gene (Thackeray and Ganetzky, 1994). The histidine-decarboxylase-encoding hdc gene, which is essential in the production of ERG transients, also gives rise to multiple transcripts which are found in various regions of the central brain and the thoracic ganglion (Burg et al., 1993). Finally, the transcript of the Dmca1A-encoding locus, which gives rise to a voltage-gated calcium channel
protein, and displays a complicated pattern of alternative splicing, is a particularly appealing candidate for being regulated by NONA (Smith et al., 1996). The alleles of the Dmca1A-encoding gene, in fact, produce an array of courtship song, visual behaviour, and viability defects similar to, although distinguishable from, those produced by nonA alleles. The cac mutation specifically affects, like the diss mutation, the pulse portion of the male's courtship song, rendering the pulses visibly polycyclic and high in amplitude. However cac mutant songs can be distinguished from diss since in cac all the pulses, independently from their position in a given train, are highly polycyclic (Kulkami and Hall, 1987). The night-blindA mutation, another allele of the Dmca1A-encoding locus (Smith et al., 1996) affects the optomotor responses and produces reduced transient spikes in ERG recordings. Finally, mutant alleles of l(1)L13 (which also map to the Dmca1A-encoding locus Smith et al., 1996), cause late embryonic lethality (Kulkami and Hall, 1987). The diversity of the Dmca1A transcripts suggests that the complicated complementation interactions observed between cac, nbA, and l(1)L13 mutations (Kulkami and Hall, 1987) could be due to isoform-specific lesions (Smith et al., 1996).

Besser et al. (1990) showed that the NONA protein is localized in several puffs on polytene chromosomes of Drosophila larvae, and its binding seemed correlated with active transcription within these puffs (Frash and Saumweber, 1989). This finding is not in disagreement with the idea that NONA could be a splicing factor. In fact there is evidence that splicing can occur on nascent transcripts (Dong et al., 1993; Beyer and Osheim, 1988). Moreover, in Drosophila there are two other examples of splicing factors which can be found on polytene chromosome active puffs: the proteins B52 (Champling et al., 1991) and RBP1 (Kim et al., 1992). Both proteins are members of the SR family of nuclear phosphoproteins, which appear to function in constitutive and alternative splicing (Mayeda et al., 1992; Zahler et al., 1992; Kim et al., 1992; Fu et al., 1992).

The nonA^{P14}, nonA^{H2} and nonA^{diss} mutant alleles have been sequenced by Rendahl et al. (1996). They are all caused by single point mutations: nonA^{P14} codes for an asparagine instead of a glycine at amino acid position 400 (based on the nonA
sequence from Jones and Rubin, 1990); \textit{nonA}^{H2} has an arginine instead of a glycine at position 463, while in \textit{nonA}^{dis} a cysteine is found at position 548 instead of the wild-type arginine. The mutations in \textit{nonA}^{P14} and \textit{nonA}^{H2} are respectively in the centre of and just outside the second RRM repeat. The latter is close enough to this RRM that it could affect ligand binding (Kenan \textit{et al.}, 1991). \textit{nonA}^{dis}, which is the most severe mutation, surprisingly does not lie in the RRM domains, but it is found in the proximal charged region. Simplistically it could then be inferred that NONA visual functions are particularly sensitive to alterations in the RRM domains, with courtship song being also affected whenever the charged region is mutated. An alternative interpretation would be that the array of phenotypes observed in \textit{nonA} mutants results from increasingly severe, non-specific alterations of the protein. Therefore, relatively mild \textit{nonA} mutations would affect visual behaviour only, while more severe protein alterations would affect courtship song as well.

The levels of protein produced by each mutant allele were examined in Western-blots by Rendahl \textit{et al.} (1992). It was found that all the mutants, with the exception of \textit{nonA}^{P14}, produced normal amounts of full-length NONA protein. The fact that \textit{nonA}^{P14} has lower protein levels is intriguing since this allele affects only vision. This observation, taken together with the fact that a \textit{nonA} encoding transgene which yields subnormal levels of protein is still able to rescue all \textit{nonA}-mutant phenotypes, suggest that protein levels might not be a primary factor in determining the mutant phenotypes (Stanewsky \textit{et al.}, 1996).

The phenotypic effects caused by an amorphic allele of the \textit{nonA} locus have been studied by generating an X-linked deletion (\textit{T(1;4)9e2-10}) which uncovered both \textit{nonA} and a distal, partially overlapping, lethal locus (\textit{l(1)i19e}, Jones and Rubin, 1990). Viable \textit{nonA}(\textit{\textdagger}) mutants were then generated by transforming flies carrying the (\textit{T(1;4)9e2-10}) deletion with a clone encoding exclusively for the lethal gene's product (Stanewsky \textit{et al.}, 1993). In hemizygous males, deletion of the \textit{nonA} gene causes semilethality: only 10-30\% of the expected number eclose from metamorphosis. If kept at a temperature below 25°C no \textit{nonA}(\textit{\textdagger}) males emerge. \textit{nonA}(\textit{\textdagger}) individuals are deficient in locomotor activity and flight and less than one-third of them survive to
reproduce. In visual and courship behaviour, nonA-deficient males show similar, although more severe defects to those of nonA\textsuperscript{disS} mutants, thus confirming that the nonA\textsuperscript{disS} mutation is indeed the allele closest to an amorphic mutation.

Several other nonA mutations have been generated \textit{in vitro} in order to assess the behavioural repercussions of altering highly conserved and fuctionally important amino acid residues within the RNA-binding domains and the C-terminal charged region (Rendahl \textit{et al}., 1996; Stanewsky \textit{et al}., 1996). The results of these studies reveal that the first RRM domain (RRM1) in nonA is absolutely necessary for all the known functions of NONA. Mutations in this region, not only cause severe defects in both visual and song phenotypes, but also invariably reduce the viability of the affected flies. In particular, mutations replacing functionally important residues in the RNP2 and RNP1 of RRM1 resulted in "worse-than-nothing" phenotypes: mutant flies were in fact much more subviable and feeble than nonA\textsuperscript{(/)} flies. This suggests that these mutations could be antimorphic, in a way that the mutated form of NONA might be actually interfering with the functioning of other, similarly acting, polypeptides. In nonA\textsuperscript{(/)}, other nucleic-acid-binding complexes could be partially compensating for the gene's product absence, without the interference of qualitatively altered NONA molecules. The RRM2 domain, on the contrary, seems to be much less important for the functioning of the nonA product. Two mutations in the RNP2 of RRM2 left unaltered all the visual, courtship and viability phenotypes tested. Other mutations at the RRM2 produced defects of the visual behaviour only, leaving the song and viability phenotypes generally unaltered. Therefore it has been suggested that RRM2 might be primarily involved in visual behaviour. Given its lower level of conservation, compared to that of RRM1, the second RRM of NONA could be specifically processing transcripts required in the nervous system for normal vision in a stringent sequence-specific recognition manner. The high stringency of such NONA-ligand interactions for normal visual output is consistent with the observation that, whenever a mutation affects NONA functions, the visual phenotype is the first to be affected. Increasingly severe mutations (such as those affecting the RRM1 of NONA) would lead to subsequent song and, finally, viability defects.
Interestingly, all the in vitro RRM mutated forms of NONA bind to chromatin in an essentially normal manner (Stanewsky et al., 1996). This implies that domains important for chromatin binding might be different from those interacting with RNA. In previous experiments, Stanewsky et al. (1993) had also shown that two different C terminal-truncated forms of NONA could bind to chromatin, suggesting that this part of the protein is not responsible for NONA's presence on polytene chromosomes. Furthermore, both two C-terminal truncated NONA types were unable to rescue any behavioural mutant phenotype, indicating that all the biological functions of NONA are independent from its binding to chromatin.

The distribution of nonA RNA and protein during embryonic development has been studied (Rendahl et al., 1992). The nonA transcript is already present in the fertilized egg. At a more advanced developmental stage its expression seems to decrease in all tissues except in the central nervous system (CNS) where it starts to decline only towards the end of embryonic development. The expression pattern of the NONA protein is similar, but not identical, to that of its RNA. The protein seems to be present ubiquitously in the developing oocyte and the embryonic, larval, pupal and adult stages. Interestingly, the protein is localized in the cytoplasm of the fertilized egg, but it enters the nuclei at the cellular blastoderm stage (Frash and Saumweber, 1989). It then remains nuclear in all tissues during embryonic development. For the protein there is no apparent central nervous system-specific expression as observed for the nonA RNA. In the adult flies, NONA is localized in the nuclei of cells of most tissues analyzed. Its expression was detected in many non-neural tissues as well as in the nervous system (Rendahl et al., 1992). The expression pattern of nonA RNA and protein has been studied in all the nonA chemically-induced mutants and compared to that of wild-type flies. All the mutants show the same amount and distribution of RNA and protein as seen in the wild-type. The only exception is nonAP14 which presents a significant decrease in antibody-staining levels, consistent with the results obtained with Western Blots for this allele (Rendahl et al., 1992).

It is rather surprising that a protein which is ubiquitously expressed throughout all stages of development and adulthood, would be involved in determining specific
and distinct aspects of behaviour such as vision and courtship. However, NONA is not a unique case in this respect. For example, the products of the per and dnc genes are found in most Drosophila tissues analyzed during all developmental and adult stages. In spite of this, deletions of per or dnc are fully viable, but mutations at either of these two loci cause multiple but specific defects in the fly's behaviour (circadian and courtship behaviour in the case of per, reviewed by Hall and Kyriacou, 1990; learning behaviour in the case of dnc, reviewed by Quinn and Greenspan, 1984; Hall, 1986; Tully, 1987).

The significance of the temporal expression of nonA has been dissected by Rendahl and Hall (1996) with the use of a heat-shock-inducible nonA construct. An hsp-nonA* (cDNA) transgene was placed in a nonA<sup>dis</sup> genetic background and activated either during development only, or during the first five days of adulthood only, or during both development and imaginal stages. Surprisingly, it was found that any of these three temporal patterns of nonA expression could produce full rescue of the visual and courtship singing defects of nonA<sup>dis</sup> mutants. The fact that the conditional nonA construct was tested on a nonA<sup>dis</sup> genetic background instead of a nonA<sup>-</sup>, did not allow full appreciation of nonA requirements during development in terms of viability and general vigour. nonA does have a role in development, since nonA<sup>-</sup> flies often do not eclose and the gene product is known to be essential for the female germ line (Stanewsky <i>et al.</i>, 1993). The finding that nonA expression during development only is enough to guarantee full rescue of behaviours in the adult, could be explained if the NONA protein was very stable. In support of this, it has been noted that, in wild-type, NONA protein levels are still high in embryonic tissues 6 hours after the nonA mRNA levels have started to decline (Stanewsky <i>et al.</i>, 1993). Alternatively, nonA expression during development could turn on a cascade of downstream genes whose products then keep acting for a very long time. nonA is not the only case of a gene whose expression in adulthood only is enough to obtain rescue of the behaviours it controls. Expression of the per gene (using a hsp-per construct) exclusively in adulthood or even only during behavioural testing, assured rescue of the rhythmic phenotypes associated with the locus. However for per, continuous expression of the gene is required to obtain the relevant rhythmic behaviours, owing to the fact that PER
protein has a very short half life (Ewer *et al.*, 1988; 1990). Thus expression of *per* in pre-adult stages only is not enough to restore rhythmicity once the flies are adults.

**Aims of the project.**

We have seen how mutations of the *nonA* gene can affect the expression of the male's love song. Sound pulses produced by *nonA* <sup>dis</sup> mutants are reminiscent of the types of pulses produced by males of other *Drosophila* species. One of these species is *D. virilis* whose song pulses are in fact polycyclic (Hoikkala *et al.*., 1982; Hoikkala and Lumme, 1987). This similarity between a mutant phenotype in *D. melanogaster* and a wild-type phenotype in another species is analogous to the case of the *per*<sup>S</sup> mutation of *D. melanogaster*. This mutation not only shortens the circadian locomotor activity cycles of the fly (Konopka and Benzer, 1971), but it also shortens the ~60 s male's love song cycles to about 40 s (Kyriacou and Hall, 1980; 1981). Males of *D. simulans* sing with love song cycles of about 40 s (Kyriacou and Hall, 1980; 1986) and we have seen how species-specific changes in the *period* gene around the Threonine-Glycine repeat region convey the species-specific characteristics of the love song rhythms (Yu *et al.*, 1987; Wheeler *et al.*, 1991). Could it possibly be that a corresponding situation exists in the case of the *nonA* gene, i.e. that the *nonA* gene conveys species-specific courtship song information?

The *per* gene not only conveys species-specific lovesong information, but also can transmit species-specific patterns of locomotor activity (Petersen *et al.*, 1988). *per* has therefore been called a "speciation" gene by Coyne (1992) because it transmits species-specific information for at least two characters which may help to maintain species boundaries, and consequently may have important implications for the speciation process itself. How the love song may prevent interspecific matings has already been covered earlier, but locomotor activity patterns could be equally, or even more important as a pre-mating interspecific barrier, by restricting flies' movements to different phases of the day. In the case of the *nonA* gene, could it be that species-specific characteristics of the love song, particularly of the sound pulses, which are
clearly affected in nonA\textsuperscript{diss} mutants, are controlled by this single gene? A simple way to answer this question is to isolate the nonA gene from a different species and transform it into \textit{D.melanogaster} flies carrying the nonA\textsuperscript{diss} mutation or a deletion of the locus (nonA\textsuperscript{c}).

So far \textit{per} is the only behavioural gene that has restored species-specific characters from a donor species to the host (Petersen \textit{et al}., 1988; Wheeler \textit{et al}., 1991). In both phenotypes discussed above, locomotor activity and courtship song, the transfer of the behaviour by the donor species \textit{per} gene was complete. This in itself is surprising, because speciation theories predict that changes in behaviour, or in any other phenotype such as morphology, occur by the accumulation of many small changes at multiple loci (Coyne, 1992). The \textit{per} gene clearly contravenes this elementary law of traditional speciation thinking. Could it be that the nonA gene also presents us with an example of such a complete transfer of behaviour from one species to another? The precedents have been set by the \textit{per} gene, and my project seeks to examine whether the nonA gene is similarly implicated in the control of species-specific behaviour. To investigate this possibility, I have cloned the nonA gene from a species whose courtship song was as different from that of \textit{D.melanogaster} as possible. All the species of the \textit{D.melanogaster} subgroup have pulse songs that are very similar to those of \textit{D.melanogaster}, so clearly these were not appropriate for my purposes (Cowling and Burnet, 1981). A great deal of work has been done on the courtship song of \textit{D.virilis} (Hoikkala \textit{et al}., 1982; Hoikkala \textit{et al}., 1987 Hoikkala and Lumme, 1984), which is very different from that of \textit{D.melanogaster}. Given the use of \textit{D.virilis} as a comparative model system, particularly in developmental studies (Michael \textit{et al}., 1990; Heberlein and Rubin, 1990; O'Neil and Belote, 1992), and given the availability of genomic libraries of this species, I decided to begin my investigations by attempting to identify a nonA homologue in \textit{D.virilis}. By examining the rescue of the visual optomotor behaviour and love song characteristics in a number of independent transformants carrying the \textit{D.virilis} nonA homologue, I was able to assess whether in fact the nonA gene can be considered to be a "speciation gene" in the same way as \textit{per}. 
CHAPTER 2

MATERIALS & METHODS
BASIC PROCEDURES

Basic techniques such as the purification of nucleic acids with phenol-chlorophorm, ethanol precipitation, and DNA quantitation, were performed following Sambrook et al. (1989).

NUCLEIC ACIDS ISOLATION

Genomic DNA isolation:

\[ \text{TE} = 10 \, \text{mM Tris-HCl (pH 8.0); 1 mM EDTA (pH 8.0).} \]

Fly genomic DNA to be used in PCR reactions was prepared using the method of Gloor and Engels (1990). Single males were ground in 50 µl of buffer (10 mM Tris-HCl pH 8.2; 1 mM EDTA; 25 mM NaCl, 20 µg/ml Proteinase K), incubated at 37 °C for 30 min, and then heated at ≥ 92°C for at least 2 min. 1µl of DNA was sufficient for each 10 µl of PCR reaction.

When better quality DNA was required (i.e. for Southern blotting), 10-50 flies were homogenized in 200-500 µl of a solution containing 0.1 M Tris-HCl (pH 9.0); 0.1 M EDTA; 1% SDS and 0.5-1% DEPC (added directly before use) and heated at 70°C for 20-30 min. The samples were incubated on ice for 30 min after adding 14 µl of Potassium Acetate for each 100 µl of homogenate. The DNA was recovered in the supernatant after spinning at 4°C for 30 min. 5 µl of RNaseA were added and the samples incubated at 37° for at least 30 min. The DNA was purified with phenol/chloroform and precipitated by adding 0.5 volumes of isopropanol and spinning for 15 min at room temperature. The DNA was then washed with 70% ethanol, dried and resuspended in 10-20 µl of TE.
Plasmid DNA isolation:

Small scale:

Solution A = 50 mM Tris-HCl (pH 8.0); 4% Triton X-100; 2.5 M LiCl; 62.5 mM EDTA.

The method by Goode and Feinstein (1992) was used to rapidly prepare plasmid DNA to be digested. 15 ml of an overnight bacterial culture was centrifuged at 13000 rpm for 30 s, and the pellet resuspended in 100 μl of solution A. The DNA was then extracted in 100 μl of phenol/chlorophorm and precipitated in 200 μl of cold 100% ethanol. The pellet was washed with 1 ml of 70% ethanol, dried, and then resuspended in 10 μl of TE + 100 μg/ml RNaseA.

Midi prep:

Up to 200 μg DNA from a high copy plasmid vector were isolated using the Hybaid Recovery™ Plasmid Midi Prep Kit. This kit usually provided high quality DNA to be used as a template in sequencing reactions.

Large scale:

Large scale plasmid DNA was obtained using the Magic™ Megapreps DNA Purification System by Promega or the Qiagen Plasmid Maxi Kit. Culture volumes and procedures followed the instructions provided by each kit.

Total RNA extractions:

Total RNA was extracted from D. virilis using the RNAzol™ method from Biotecx Laboratories, Inc., Houston, Texas. For long term storage the RNA was kept in isopropanol at -80°C in order to minimize degradation.
POLYMERASE CHAIN REACTION (PCR)

7.4X PCR mix = 167 µl of 2 M Tris-HCl
83 µl of 1 M Ammonium Sulphate
33.5 µl of 1 M MgCl_2
3.6 µl of 2-mercaptoethanol
3.4 µl of 10 mM EDTA (pH 8)
75 µl of 100 mM dATP
75 µl of 100 mM dCTP
75 µl of 100 mM dGTP
75 µl of 100 mM dTTP
85 µl of 10mg/ml DNase free BSA.

The Polymerase Chain Reaction (PCR) was performed according to Jeffreys et al. (1988). Each 10 µl of reaction contained 1.35 µl of 7.4x PCR mix, 1 µl of each 10 µM primer, 0.15 µl of Taq Polymerase (5 units/µl), and H_2O to volume. Taq Polymerases from different sources were used, such as Amplitaq Polymerase from Perkin-Elmer-Cetus, Taq polymerase from Amersham, or Thermostable DNA Polymerase from Advanced Biotechnologies. The reactions were carried out in a Perkin-Elmer-Cetus thermal cycler or in a Techne Programmable PHC-1 and PHC-3. The thermal profile mostly used was 1 min at 95°C (denaturation); 1 min at 65°C (annealing); 1 min at 70°C (extension); for 30-40 cycles.

RT-PCR

Reverse transcription from total RNA was performed by adding to 1 µg of RNA to the following "RT mix":

- 4 µl 5X First Strand Buffer (250 mM Tris-Hcl pH 8.3; 375 mM KCl; 15 mM MgCl_2)
- 0.8 µl dNTPs (25 mM each)
- 1 µl specific primer (10 µM)
- 1 µl Reverse Transcriptase (M-MLV RT from GIBCOBRL; 200 U/µl))
- 0.6 µl rRNasin (Ribonuclease Inhibitor from Promega; 38 U/µl)
• DEPC H₂O to a total volume of 20 µl.

**DEPC H₂O** = 1 ml of DEPC (diethyl pyrocarbonate, Sigma) was added to each litre of H₂O and left overnight before autoclaving.

The "RT mix" was incubated at room temperature for 15 min, then at 37°C for 65 min, and finally heated at 95°C for 10 min. 5 µl of the "RT mix" were then used as a template DNA for each PCR reaction which contained 2.7 µl of 7.4X PCR mix; 1 µl of 5' primer (10 µM); 1 µl of 3' primer (10 µM); 0.3 µl of Taq Polymerase; and 11 µl of H₂O.

**GEL ELECTOPHORESIS AND DNA RECOVERY FROM GEL BANDS**

1X **TAE** = 0.04 M Tris-acetate; 0.001 M EDTA.

6X **Gel-loading buffer** = 0.25% bromophenol blue; 0.25% xylene cyanol FF; 30% gycerol in water.

Agarose gel electrophoresis followed the methods of Sambrook *et al.* (1989). Gels were usually prepared at concentrations 0.7%-1.5%, according to the size of the DNA fragments to be resolved. 0.5 mg/ml of ethidium bromide was added to the melted gel before pouring it, and once set, the gel was run in 1X TAE buffer at voltages which ranged between 20-120 V. DNA markers, to be run alongside the samples, were either λ cut with *HindIII* (for fragments between 2 and 23 Kb), or φ X174 cut with *HaeIII* (for fragments from 70 bp to 1.3 Kb). 1/10 volume of gel loading buffer was added to increase the density of each DNA sample and to indicate the progression of the electrophoresis. The DNA was then visualized on the gel by UV fluorescence and photographed using a video imaging system (GDS2000, UVP International). For a better quality photograph, a Polaroid MP-4 camera and Kodak negative films (T-max Professional 4052) were used. The films were then processed with Kodak LX24 developer, FX40 fixer and HX40 hardener.
DNA was recovered from agarose bands using the QIAEX and QIAEXII gel purification kits (QIAGEN Inc.). In case of very large bands (weight above 250 mg), the DNA was extracted by placing the band in a dialysis bag filled with 2 ml of TE buffer and performing electrophoresis on the bag submerged in TAE. The DNA migrated from the gel band to the TE buffer which was then transferred in microcentrifuge tubes and purified with one phenol/chlorophorm and one chlorophorm/isoamylalcohol extraction. The DNA was then ethanol-precipitated and resuspended in an appropriate volume of H₂O.

DNA SEQUENCING

Direct sequencing of PCR products:

Double stranded direct sequencing of PCR products was performed according to Bachmann et al. (1990) using the dideoxy chain-termination method of Sanger et al. (1977). The amplified DNA to be sequenced was usually recovered from an agarose gel band using the QIAEX gel purification kit (QIAGEN). The template was annealed to the specific primer by mixing 6 μl of DNA with 2 μl of sequencing buffer 5X (200 mM Tris-HCl pH 7.5; 100 mM MgCl₂; 250 mM NaCl), 1.5 μl of the specific primer (10 μM) and 0.5 μl of 10% Np40. The tube containing the mixture was then boiled for 3 min and rapidly cooled in dry ice. The sequencing reactions were performed using the Sequenase Version 2.0 Kit from United States Biochemicals and then run on a 5% denaturing polyacrilamide gel using a Sequi-Gen (Bio-Rad) nucleic acid sequencing cell. At the end of the run the gel was fixed for 15 min in a solution 10% acetic acid and 10% methanol and placed on a sheet of 3MM paper. The gel was then vacuum dried at 80°C and placed in an autoradiography cassette with a Fuji RX100 X-ray film.

Sequencing double stranded plasmid DNA:

1.5-3 μg of plasmid DNA were sequenced using the T7 Sequencing Kit from Pharmacia. The annealing of primer to the double stranded DNA was performed
following the standard protocol suggested in the kit instruction booklet.

**SOUTHERN BLOTTING OF DNA** (Southern, 1975).

**Denaturing solution** = 1.5M NaCl; 0.5 M NaOH.

**Neutralizing solution** = 1.5 NaCl; 1M Tris-HCl (pH 8.0).

20X **SSC** = 3 M NaCl; 0.3 M Na₃ citrate (pH 7.0).

The gel to be blotted was photographed after placing a ruler next to it in order to have a record of the position of the DNA markers. When large fragments of DNA (>10 Kb) needed to be transferred, the gel was depurinated for 10 min in a solution of 0.25 M HCl, rinsed with water, then placed in the denaturing solution for 30 min, and finally neutralized for 30 min. The gel was then placed on the blotting apparatus (as described in Sambrook *et al.*, 1989) and surrounded by cling film. A precut, prewet (3X SSC) piece of Hybond-N filter (Amersham International) was carefully placed on the gel, followed by one wet (3X SSC), one dry pieces of 3MM paper, and a thick pad of absorbent paper towels. DNA transfer was usually allowed to occur overnight. The filter was then rinsed in 3X SSC, air dried and fixed with a 45 s UV light exposure on a transilluminator.

**Preparation of radiolabeled DNA probe:**

**10X Random priming buffer (RPB)** = 500 mM Tris-HCl (pH 6.6); 100 mM MgCl₂; 10 mM DTT; 500 μg/ml BSA (DNase and RNase free, Sigma).

3 **dNTPs** = 20 mM each of dATP, dGTP and dTTP in H₂O.

A modification of the method of primer extension with random primers by Keller *et al.* (1989) was used to generate nucleic-acid probes with specific activities of 10⁹ cpm/μg. 50-200 ng of double-stranded DNA in 10 μl of H₂O were heat-denatured
by boiling for 5 min and placed immediately on ice for at least 5 min. The denatured DNA was mixed with:

- 5 µl of 10X RPB
- 5 µl of p(dN)_6 (2 mg/ml, Pharmacia)
- 1 µl of 3 dNTPs
- 2 µl of [α\textsuperscript{32}P]dCTP (>3000 Ci/mmole)
- 1 µl of Klenow (5 U/µl)
- 26 µl of H₂O.

The reaction was incubated at 37°C for 30 min, and then terminated with 5 µl of 0.5M EDTA. The unincorporated nucleotides were removed by ethanol precipitation adding 1 µl of tRNA (1 µg/µl), 1/10 volume of 3 M NaAc pH 5.2, and 2 volumes of ethanol 100%. The probe was denatured by heating at 100°C for 5 min and immediately cooled on ice for 5 min before mixing to the filter-hybridization solution (see later).

Filters hybridization with \textsuperscript{32}P probes:

Prehybridization:

At least 2 hours before adding the radiolabeled probe, the blotted filters were incubated at 65°C in a pre-hybridization solution composed of:

- 1 mg/ml ficoll
- 1 mg/ml polyvinyl pyrrolidone
- 5mg/ml BSA fraction V
- 0.5% SDS
- 2 mg/ml Marvel dried milk
- 6X SSC.
Hybridization:

The pre-hybridization buffer was changed with some identical fresh solution (pre-heated at 65°C) just before adding the radiolabeled probe. The filter was then incubated at 65°C overnight.

Filters washing:

Hybridized filters were washed briefly at room temperature in a solution of 4X SSC, 0.1% SDS. This wash was repeated at 65°C for 30 min. Then a series of washes of increasing stringency were performed (2X SSC, 0.1% SDS; 1X SSC, 0.1% SDS; 0.5X SSC, 0.1% SDS; 0.1X SSC, 0.1% SDS), until the filters emitted almost background levels of radioactivity. Every wash was done for 30 min at 65°C. Filters were then air dried (not completely if the filter had to be re-probed), wrapped in SaranWrap, and placed in an autoradiography cassette containing an intensifying screen and a Fuji RX100 X-ray film.

CLONING PROCEDURES

Dephosphorylation of linearized plasmid DNA:

1) Using calf intestinal alkaline phosphatase (CIP):

10X CIP buffer = 0.5 M Tris-HCl (pH 9.0); 10 mM MgCl₂; 1 mM ZnCl₂; 10 mM spermidine.

10X STE = 100 mM Tri-HCl (pH 8.0); 1 M NaCl; 10 mM EDTA.

0.01 u/pmol ends of CIP (GIBCO BRL) were added to a 48 µl reaction containing the linearized DNA and 5 µl of 10X CIP buffer. To dephosphorylate protruding 5' termini, the mixture was incubated at 37 °C for 30 min, and for an additional 30 min after adding a second aliquot of the enzyme. To dephosphorylate DNA with blunt ends or recessed 5' termini, the reaction was incubated for 15 min at 37°C, and 15 min at 56°C, and the two incubations repeated after adding a second
aliquot of CIP. The reaction was then stopped by adding 10 µl of 10X STE, and 5 µl of 10% SDS and heating at 68°C for 15 min. The DNA was then extracted twice with phenol/chlorophorm, and twice with chlorophorm.

2) Using Shrimp Alkaline Phosphatase (SAP):

10x SAP buffer = 200 mM Tris-HCl (pH 8.0); 100 mM MgCl₂.

Dephosphorylations were performed following the instructions accompanying the SAP enzyme (United States Biochemicals). The reactions were prepared in 20 µl total volume containing 2 µl of 10X SAP buffer, linearized DNA, and suggested amounts of SAP. The mixture was incubated at 37°C for 1 hour, and the reaction stopped by heating at 65°C for 15 min. After heating, the enzyme was completely inactive, and the DNA was ready to use in ligation reactions.

Ligation reactions:

Ligations were performed following Sambrook et al. (1989). Insert and vector DNA were usually mixed in 3:1 ratios in a 15 µl total volume of reaction containing T4 Ligase Buffer (from GIBCO BRL) at working concentration (1X), and 1 unit of T4 DNA ligase (GIBCO BRL). The ligation reactions were incubated overnight at 16°C.

Plasmid transformation (chemical method):

Preparation of competent cells:

XL1-Blue = recA1 endA1 gyrA96 thi-1 hsdR17 supE44 relA1 lac [F proAB lacI ΔM15 Tn10 (TetR)].

LB (Luria-Bertani) medium (or Luria broth) = 1% (w/v) Bacto-tryptone (Difco); 0.5% (w/v) Bacto-yeast extract (Difco); 1% (w/v) NaCl.

Tfb I = 30 mM KAc; 100 mM RbCl₂; 10 mM CaCl₂; 50 mM MnCl₂; 15% (v/v) glycerol; pH 5.8.

TfbII = 10 mM MOPS; 75 mM CaCl₂; 10 mM RbCl₂; 15% (v/v) glycerol; pH 6.5.
Bacterial cells (E. coli strain XL1-Blue, Stratagene) were grown at 37°C in 500 ml of LB broth. When the early log phase was reached (OD<sub>550</sub> = 0.5-0.55), the culture was chilled on ice for 5 min, and then spun in a Sorvall centrifuge (6,000 r.p.m. for 5 min at 4°C). The cells were resuspended in 200 ml of TfbI (2/5 original volume), left on ice for 5 min and centrifuged again at 6,000 r.p.m. for 5 min (at 4°C). The pellet was resuspended in 8ml of TfbII (1/25 volume), and left on ice for 15 min. Suitable volumes of cells (usually 100-200 μl) were then aliquoted into microcentrifuge tubes, using a prechilled pipette and tubes. The tubes were snap freezed in dry ice and stored at -80°C.

**Transformation procedure:**

**LB agar plates =** LB medium + 15 g/liter Bacto-agar.

100 μl of competent cells were added to 10 μl of ligation and left on ice for 20 min. The DNA-cells mixture was heat shocked at 42°C for 45 s and returned to ice for 1-2 min. The transformed cells were grown at 37° for 1 hour in 1ml of Luria broth and then plated (usually in 100 μl aliquots) on LB agar plates containing appropriate antibiotics. When blue-white selection was required, the plates were made adding X-gal and IPTG according to Sambrook *et al.* (1989).

**Transformation by electroporation:**

**Cells for electroporation:**

XL1-Blue cells were harvested in Luria broth at an A<sub>600</sub> of about 0.55. A series of washes was performed in order to concentrate the cells into a dense suspension and to wash away the large amounts of electrolytes in the culture medium. The washing solutions used (either sterilized H<sub>2</sub>O or 10% glycerol) were pre-chilled at 4°C. The washing protocol was the following:

1 L of culture ⇒ pellet (4,000 r.p.m., 15 min, 4°C) ⇒ resuspend in 1 L of H<sub>2</sub>O

⇒ pellet (4,000 r.p.m., 15 min, 4°C) ⇒ resuspend in 1/2 L of H<sub>2</sub>O
pellet (4,000 r.p.m., 15 min, 4°C) → resuspend in 20 ml of 10% glycerol
pellet (4,000 r.p.m., 15 min, 4°C) → resuspend in 2 ml of 10% glycerol.

The cells were then aliquoted (50 µl in each tube) and stored at -80°C.

Electroporation procedure:

Ligations were ethanol precipitated and resuspended in 5 µl of TE or H2O. 1-5 µl of transforming DNA was mixed with 40 µl of washed cells (prepared as above). The mixture was transferred on a cuvette chilled on ice and placed between the electrodes of the electroporation unit. An electrical pulse of 15 KV was delivered, and the cells immediately resuspended in 1 ml of Luria broth and gently shaken at 37°C for 1 hour before plating them onto LB agar plates (see above).

Bacterial cracking:

This method was used to rapidly identify recombinant clones when blue-white selection was not possible. 1ml of overnight culture from a single colony was spun down and resuspended in 200 µl of 1X gel loading buffer. After adding 200 µl of phenol/chloroform, the cells were vortexed and spun for 2 min. 20 µl of the top layer were then loaded on an agarose gel (usually 1%) and run at 100-120 V. Insert-containing plasmid DNA runs considerably slower than non-recombinant and could be easily detected.

GENOMIC DNA LIBRARY SCREENING

A D.virilis genomic DNA library constructed by Ron Blackman (Harvard University, Cambridge, MA) was kindly provided by John Belote (Syracuse University, NY). The library had been prepared with genomic DNA from adult flies partially digested with the enzyme MboI. Size selected fragments (15-20 Kb) had been inserted
between the Bam HI sites of the lambda (λ) EMBL3 phage (Frischauf et al., 1983) and packaged in vitro.

**Plating and titration of the library:**

**Preparation of plating bacteria:**

A culture of the Y1090 bacterial strain was inoculated overnight at 37°C in 200 μl of Luria broth containing 0.2% maltose and ampicillin (50 mg). The cells were then spun down and resuspended in 10 mM MgSO₄ to an appropriate density (OD₆₀₀ = 0.5).

**Plating bacteriophage λ:**

**SM buffer** = 50mM Tris-HCl (pH 7.5); 100 mM NaCl; 8 mM MgSO₄; 0.01% gelatin.

**LB top agarose** = 1% (w/v) Bacto-trypptone; 0.5% (w/v) Bacto-yeast extract; 0.5% (w/v); 0.6% (w/v) agarose.

Serial dilutions of bacteriophages were prepared in 50 μl of SM buffer. 0.2 ml of plating bacteria were then added to each dilution, and incubated at 37°C for 15 min, allowing the cells to adsorb the bacteriophage particles. 3 ml of molten (47°C) LB top agarose were added, and each preparation plated on pre-set LB agar plates. The plates were incubated at 37°C until plaques appeared. The number of plaques given by each dilution was an indication of the library titre.

**In situ hybridization of bacteriophage λ plaques:**

**Preparation of the filters:**

**Denaturing solution** = 1.5 M NaCl; 0.5 M NaOH.

**Neutralizing solution** = 1.5 M NaCl; 1 M Tris-HCl (pH 7.6).

Appropriate dilutions of the library were plated in a total number of 30 LB plates and incubated at 37°C overnight. When the plaques became almost confluent, the plates were cooled at 4°C for 1 hour. One marked Hybond-N (Amersham) circular
filter was placed on each plate for 5 min. The filters were lifted carefully from the plates and placed for 10 min in the denaturing solution, for 1 min in 1 M Tris-HCl (pH 7.6), for 10 min the neutralizing solution, and then rinsed in 3X SSC. When completely dry (usually after 1 hour at room temperature), the filters were sandwiched in SaranWrap, and the DNA was fixed by UV exposure for 45 s.

**Prehybridization, hybridization and washing of the filters:**

These techniques were performed as already described above for Southern blotting filters.

**Isolation of positives and rescreening:**

The position of the possible positives on the developed film was located on the plates. A "plug" of top agarose containing the plaque of interest was then lifted using a pipette tip, and placed in a microcentrifuge tube containing 500 µl of SM buffer and 20 µl of chlorophorm. The suspension was left at room temperature for 4-6 hours to allow the bacteriophage particles to diffuse out of the agarose. Several dilutions of the suspension were then plated and rescreened as described above. The procedure of positive picking, replating and rescreening was continued until all the plaques on a plate were positives.

**Phage lysate preparation: plate method:**

For each plate, about $1 \times 10^5$ pfu of bacteriophage (or 60-100 µl of a phage "plug" left to elute 1 hour at room temperature in 1 ml of SM buffer) were plated and incubated at 37°C until the plaques became confluent. The plates were then overlaid with 2-3 ml of SM buffer, and the top agar scraped with a spatula into a high speed 30 ml centrifuge tube (Corex). After incubating at room temperature for 30 min, the suspension was centrifuged at 10,000 r.p.m. for 10 min at 4°C. The supernatant was transferred in a fresh tube, and 0.3% of chloroform was added for long term storage.
**Bacteriophage DNA isolation:**

Bacteriophage λ DNA was extracted from 10 ml of phage lysate using the Magic™ Lambda Preps DNA Purification System.

**P-ELEMENT TRANSFORMATION OF DROSOPHILA**

P element mediated transformation was performed according to Spradling and Rubin (1982). The transformation vector used was *pw8* (Klemenz et al., 1987) which carries *white* as a marker. Embryos microinjected were either *w; Δ2-3 Sb/TM6* (Robertson et al., 1988), or *w118* (Pastink et al., 1987). When using the latter injectees, transposase was provided by co-transformation of a helper plasmid *PUChsp* Δ2-3 (a gift from J.M. Dura).

**Embryos collection and dechorionation:**

4-7 day old females were encouraged to lay eggs by placing them on a fruit juice-based food. Eggs were collected at intervals no longer than 60 minutes to assure injection prior the time of pole cell formation. Embryos dechorionation was performed manually by placing eggs over a piece of double-sided sticky tape on a microscope slide and gently rolling them until the chorion was ruptured. The dechorionated embryos were then aligned on one side of the tape with the posterior end just hanging off the tape. About 10 minutes after dechorionation the eggs were covered with Voltalef oil (Grade 10S) to prevent further dessication.
Microinjection of embryos:

Microinjection was performed using a Nicon TMS inverted microscope and a Narishige micromanipulator. Needles were prepared by pulling microcapillary tubes (GC100TF-15, Clark Electromedical Instruments) using a Flaming Brown micropipette puller (Model P80/PC, Sutter Instruments Co.) and placed on a microelectrode holder from World Precision Instruments, Inc. A constant flow of DNA from the needle was obtained thanks to a PV830 Pneumatic Pico Pump (World Precision Instruments, Inc). Embryos were injected, according to Spradling and Rubin (1982), into their posterior tip, and the amount of DNA introduced depended on the degree of dessication of the embryo.

Post-injection care of embryos:

All improperly injected or aged embryos were destroyed, and the coverslip with the remaining embryos was placed on a petri dish containing standard fly food dyed with a red food colouring. The red colour allowed easy detection of the developing embryos. More oil was added over the embryos to protect them from dessication. The plate was incubated at 18°C. Hatched larvae were removed from the plates after a couple of days, and placed in numbers of 5-10 into sugar food vials. Further development was accelerated by placing the larvae at 25°C. Adults (G₀ generation) were individually mated to w¹¹¹⁸ flies, and their progeny (G₁ generation) screened for transformants (red-eyed flies).
BEHAVIOUR

FLY STOCKS and BALANCERS

D. melanogaster:

_Canton S_ strain had been kept in the laboratory for several years.

* _w*¹⁺: see Lindsley and Zimm, 1992.

* _w; e Δ 2-3 Sb/TM6_: transposase providing strain (Laski et al., 1986; Robertson et al., 1988).


* _T(1;4)9e2-10/FM7_: Stanewsky et al. (1993). Strain with reciprocal translocation _T(1;4), uncovering the essential l(1)i9e locus and the nonA gene._


* _P[(ry) 235R11-SacII]5_: Stanewsky et al. (1993). Transformant stock carrying the _nonA_ gene with a frame-shift mutation introduced at the _SacII_ site in the first exon.

* _nonA*⁻¹ / _FM7a; ry5 0 6 / _MKRS_: donated by Jeff Hall.

* _w, FM7_: balancer for the first chromosome (see Lindsley and Zimm, 1992).

* _w; TM2/MKRS_: balancer for the third chromosome (see Lindsley and Zimm, 1992).

* _w; T 2,3 ap²/a2/SM5 TM3_: balancer for both second and third chromosome (see Lindsley and Zimm, 1992).

All balancer stocks were obtained from American or European _Drosophila_ stock centres.

_D. virilis_ (strain 1411, Finland) was donated by Anneli Hoikkala, University of Oulu, Finland.

STOCK MAINTENANCE

Fly stocks were reared on sugar/agar medium (46.3 g of sucrose, 46.3 g of yeast, 7.1 g of agar and 2 g of Nipagen in 1 L of water) in one-third pint milk bottles or glass vials (10 cm height x 2.2 cm diameter) The flies were kept in temperature-controlled rooms at either 18°C or 25 (± 1)°C.
RECORDING AND ANALYSING DROSOPHILA LOVESONGS

Recording *D.melanogaster* songs:

Newly eclosed males were isolated and placed individually into vials at 25°C. At the age of 2-4 days their song was recorded by placing one male and one or two virgin females (maximum 1 day old), into a small rectangular (1.5cm x 1.1cm large, 0.6cm high) perspex cell with a nylon mesh at the bottom. The flies were inserted in the cell through a small hole cut in one side of the cell. The cell was placed, mesh down, on top of a condenser microphone (custom-made in Leicester) contained in a foam box. The box could be closed to function as an anechoic chamber, and the movements of the flies observed through a magnifying view lens on top of the box. A light inside the box assured visibility. The box also contained an amplifier and filters, and the song was recorded onto a Revox 477 reel to reel tape recorder at 7.5 ips. The chamber temperature was monitored continuously with a digital thermometer, and recordings were carried out between 25-26°C.

Recording *D.virilis* songs:

Males and females were collected immediately after eclosion and kept in sugar food vials at 25°C. Males were kept individually, while females were grouped in numbers of 5 per vial. At about 15 days of age, one male and one female were placed in a perspex cell identical to that used for *D.melanogaster* flies, but which had a bigger hole for inserting the larger *virilis* flies. The recording conditions and equipment were the same as those used for *D.melanogaster*.

Each male's song was recorded for 10-15 minutes using a Revox reel-to-reel tape recorder. Long silences were common, and were avoided by switching the tape recorder off when males were not singing. The recording was filtered below 1000Hz and above 100Hz using an EF5-03 LP/HP filter (Barr & Stroud, Cambridge, U.K.), and then played into a CED 1401 (Cambridge Electronic Design) analogue-to-digital converter, in order to get a graphic representation of the song signal. The signals
could then be displayed on the screen of an IBM-PC 486, and stored on the hard disk. The SPIKE2 software developed by CED was used for primary analysis of the signals. This program allows simultaneous display and processing of waveform data and events.

**Song characters analysed:**

**Cycles per pulse (CPP):**

Figure 2.1 shows a small portion of one of the songs recorded, as it was displayed using the Spike2 LEVELS script. The LEVELS script can be used to detect events such as peaks and troughs from waveform channels. Channel 1 contains the song waveform. Above the waveform there is a second channel (channel 2) displaying the song events (shown as bars) which in this case are the peaks and the troughs found in the song's pulses. Peaks and troughs were automatically marked by the Spike2 program. For the peaks, a positive threshold rising above the song's background noise was first selected so that Spike2 marked every single event above it. All the peak events were stored and displayed on channel 2 above the song waveform. To automatically mark all the troughs in the song, a negative threshold was chosen below the background. All the troughs were recorded and displayed in channel 2 together with all the peaks. At this point the song was manually edited by moving along the song waveform and removing or adding events from channel 2 as needed. The editing mainly involved removing events caused by loud noises, and adding or removing some events from pulses in which the exact number of peaks and/or troughs had not been marked properly in the initial automatic analysis.

When the manual editing was completed, the events stored in channel 2 were imported into the Microsoft Excel 5.0 program. Each datum was expressed as the real time, in milliseconds, in which the event had occurred. A macro written by Martin Couchman permitted calculation of the number of cycles in each pulse and the regression for CPP during song bursts. For each pulse, the number of cycles was calculated as: \(((\text{peaks}+\text{troughs})-1) / 2\). Instructions were given that allowed the
Figure 2.1. Portion of song displayed using the Spike2 LEVELS program. Channel 1 (song waveform), channel 2 (with marks for pulse peaks and troughs) and channel 3 (with marks for sine song) are indicated. Time scale in seconds.
computer to recognize events of the same pulse from events of the next pulse and pulses of one train from pulses of the next train. The minimum gap allowed between a peak and a trough in each pulse was 0.6 ms. The maximum gap between a peak and a trough in a pulse was set at 5 ms: if an event occurred more than 5 ms from the previous one then it was not be considered part of the same pulse. The minimum gap from one pulse to the next pulse of the same train was 10 ms, so an event more than 10 ms apart from the previous one was considered part of the next pulse. The minimum distance between two pulse trains was set at 70 ms: if an event was more than 70 ms apart from the previous one, then it was considered part of the first pulse in the next pulse train. CPPs were calculated only for trains that contained at least 5 pulses or more, shorter trains were not included in the analysis. In order to make sure that the macro was properly calculating the CPPs, an entire song was manually analysed and the results compared with those obtained with the macro.

For each song, the macro calculated the average CPP scores for the pulses found in each consecutive position along the pulse trains, i.e. it calculated the average CPP for all the pulses in the first position, the second position, the third position and so on. A least squares regression was performed using these averages. Averages calculated for pulse positions containing less than 5 data points were not included in the regression analysis.

**Pulse frequency (PF):**

From the CPP analysis, the macro calculated the average frequency of each pulse by dividing the time between the first and the last event in the pulse by the number of cycles in it and converting into Hz. A least square regression was performed using the average frequency values of pulses at consecutive positions in the pulse trains. Again, only averages calculated from at least 5 frequency values were used for the regression.

**Pulse song (PS):**

The macro automatically calculated the total amount, in seconds, of pulse song. For each train of pulses the time between the first event in the first pulse and the last
event in the last pulse, gave the total length, in seconds, of the train. The total amount of pulse song was calculated as the sum of all the pulse trains' lengths in the song. The length of short pulse trains (containing less than 5 pulses) was not included in the calculation.

**Sine song (SS):**

In every song, the beginning and the end of each burst of sine song was marked manually and the events stored on channel 3 displayed above the the peaks and troughs channel (figure 2.1). The events in channel 3 were imported into Microsoft Excel 5.0 together with the events from channel 2. The macro automatically calculated the length, in seconds, of each sine song burst by using the time points that marked the beginning and the end of the burst. The total amount of sine song was calculated for each song by summing the lengths of all the sine song bursts found.

**Sine song frequency (SSF):**

Ten sine song bursts were selected from the terminal part of each song (which is usually more rich in sine song) in order to calculate the frequency. The Fast Fourier analysis of Spike2 was used to obtain the frequency of each selected sine song burst. The SSF for each song was calculated as the mean of the 10 selected sine song burst frequencies.

**Sine song proportion (SSP):**

This character was calculated automatically as the amount of sine song over the total amount of song recorded (SS/PS+SS).

**Interpulse intervals (IPIs):**

The mean IPI of each song was calculated as: PS/(Npulses-Nbursts) where PS is the total amount, in ms, of pulse song recorded, Npulses is the total number of pulses in the song and Nbursts is the number of bursts.

An example of a typical page output obtained after applying the macro to an individual song is shown in figure 2.2. Both PF and CPP regressions are accompanied
Regression of pulse frequency using data in first 22 columns:

\[
\begin{array}{lr}
\text{slope (m)} & -5.7751 \\
\text{s. error of m} & 0.5429 \\
\text{y intercept (b)} & 335.6106 \\
\text{s. error of b} & 7.1299 \\
\text{coeff. of determ} & 0.8498 \\
\text{F statistic} & 113.1718 \\
\text{degrees of freedom} & 20.0000 \\
\text{regression sum o} & 29532.9174 \\
\text{residual sum of squares} & 5219.1297 \\
\end{array}
\]

\[y = -5.7751x + 335.6106\]

Data file: Waveform

Regression of cycles per pulse using data in first 22 columns:

\[
\begin{array}{lr}
\text{slope (m)} & 0.0426 \\
\text{s. error of m} & 0.0035 \\
\text{y intercept (b)} & 1.8998 \\
\text{s. error of b} & 0.0457 \\
\text{coeff. of determ} & 0.8826 \\
\text{F statistic} & 150.3363 \\
\text{degrees of freedom} & 20.0000 \\
\text{regression sum o} & 1.6087 \\
\text{residual sum of squares} & 0.2140 \\
\end{array}
\]

\[y = 0.0426x + 1.8998\]

Data file: Waveform

Figure 2.2.

Typical page output of the macro written by Martin Couchman.
by the appropriate F-ratio which indicates the significance of the fit. Their slope and y-
axes intercepts are calculated and illustrated graphically. The amount of sine song and
pulse song and the proportion of sine song are also indicated at the bottom of the page.

WALKING OPTOMOTOR TEST

The optomotor response of walking flies was tested following the method
described by Burnet et al. (1968). 3-8 day old flies were dark-adapted for about 4
hours in food vials. Each fly was tested individually for its turning behaviour in a
moving visual field. This was created using a Plexiglass drum (diameter, 8 cm; height,
9 cm) rotating around a vertical axis, which had alternating black and white vertical
stripes.

Testing D.melanogaster:

Each fly was placed in the middle arm of a T-shaped glass tube. This arm was
painted black, so that the fly was forced to walk out of this opaque tube, into a choice
point where it could turn into the right or the left arm. To test whether the turning
behaviour corresponded to the moving environment, the tube was placed in the middle
of the rotating drum. A fly produced a correct response every time it turned out of the
opaque arm in the same direction of the rotating stripes. Each fly was given 20 trials,
and each time the rotating direction of the stripes was changed. Ten flies per genotype
were tested. The test was performed twice using different stripe widths. “Small”
stripes subtended a small angle of 13.3° from the centre of the drum; “large” stripes
subtended a larger angle of 36°. The drum was constantly rotated at 46 rpm. A desk
lamp (60 watts) was placed above the drum in order to illuminate in a uniform way the
centre of the cylindre. All tests were performed at room temperature (20-25°).

Testing D.virilis:

Given the larger size of D.virilis flies, they could not move freely in the T-
shaped tube used for testing D.melanogaster flies. Each D.virilis fly was therefore
placed in an empty glass vial which was positioned at the centre of the rotating
cylinder. The test was started once the fly had reached the top of the tube (given its negative geotropism). From there, the movement of the flies were scored as the drum was rotating. A correct optomotor response was obtained each time a fly performed at least five rotations in the same direction as the moving stripes. Again, two sets of stripes ("small" and "large", see above) were used, and each fly tested 20 times in each set of stripes.

STATISTICAL ANALYSES

All statistical analyses, such as one or two-way ANOVAs, $\chi^2$ tests, planned comparisons and Newman-Keuls $a$ posteriori tests (Winer, 1971), were performed using the Statistica Microsoft for Windows Software (StatSoft, Inc., 2325 East 13th Street, Tulsa, Oklahoma 74104, U.S.A.). Multivariate analysis was performed using the Systat Software (Version 7.0 for Windows, 1997 by SPSS Inc.).

COMPUTER ANALYSIS OF DNA SEQUENCES

All the computer analysis of DNA sequences were performed using the programs of the GCG package for molecular biology (Version 7; University of Wisconsin Genetics Computer Group, Madison, WI, USA; Devereux et al., 1984). The various programs used will be mentioned in the relevant chapter.
CHAPTER 3

The *D. virilis nonA* gene
INTRODUCTION

The nonA gene was cloned and sequenced by Jones and Rubin (1990). By P element-mediated transformation with different genomic clones from a 20 kb interval within the 14C1,2 region of the X chromosome, they identified a XbaI-EcoRI 9Kb genomic DNA fragment necessary and sufficient to rescue all the nonA mutant phenotypes (see also Rendahl et al., 1992). Simultaneously, Besser et al. (1990) cloned the BJ6 gene whose product they found associated with transcriptionally active puffs on polytene chromosomes of Drosophila larvae. When they compared their sequence of the BJ6 gene to that of nonA, they found that the two sequences were identical.

The nonA gene is organized in five exons separated by four introns (figure 3.1). Based on their cDNA data, Jones and Rubin (1990) suggested the existence of two differently spliced forms of the NONA protein generated by alternative splicing of the last intron. In the form I of NONA the fourth intron is spliced out and the fifth exon encodes the last 35 amino acids of the protein. In form II, the fourth intron encodes the last 33 amino acids of nonA, so form I and form II NONAs only differ in their carboxy terminus. However, the results of successives studies have argued against the existence of a form II protein. Firstly, only form I has been shown to carry all the nonA related biological functions (Jones and Rubin, 1990; Rendhal et al., 1992). Secondly, attempts to identify the form II NONA in Western blots have never succeeded (Stanewsky et al, 1993).
Figure 3.1. Schematic diagram of the two putative forms originating from the nonA gene (Jones and Rubin, 1990). Black boxes indicate the exons. Introns are indicated by thin lines. The thick line after the last exon represents the 3' untranslated region with the two polyadenylation signals and the polyA tail (Besser et al., 1990).

Two mRNA transcripts of ~3 and ~4 Kb have been identified in Northern blots using probes for the nonA gene. The existence of two polyadenylation sites, the first found 673 nucleotides after the stop codon, the second about 1 Kb after the first, could explain the occurrence of the shorter transcript as a result of alternative termination (Besser et al., 1990). A third 2.8 Kb transcript was sporadically resolved from the 3 Kb species by Jones and Rubin (1990) and Rendahl et al. (1992).

The 700 amino acid putative NONA protein is basic and has an N-terminus rich in glycine, asparagine and glutamine, often arranged in repeats. The C-terminal domain is rich in charged amino acids, while the last 100 carboxy terminal residues again contain clusters of glycine and asparagine. Hydrophobicity analysis indicated that the protein does not contain a membrane-spanning segment and is unlikely to have an N-terminal signal sequence (Jones and Rubin, 1990; Besser et al., 1990).

As mentioned in chapter 1, the central domain of the protein has two tandemly repeated motifs of 80 amino acids each, bearing homology to the RNA recognition motifs (RRMs). These motifs are found in a large family of more than 30 RNA-binding proteins including snRNPs, hnRNPs, and several proteins encoded by
Drosophila genes such as embryonic lethal abnormal visual system (elav) (Robinow et al., 1988), Sex-lethal (Sxl) (Bell et al., 1988) and transformer-2 (tra-2) (Amrein et al., 1988; Bandziulis et al., 1989; Besser et al., 1990; Rendhal et al., 1992). The most conserved RRM feature is an eight-amino acid sequence called RNP1 (Kenan et al., 1991). Some RNA-binding proteins contain an additional hexapeptide known as RNP2 (Kenan et al., 1991; Dong et al., 1993). The RNP1 and RNP2 elements contain aromatic and basic residues, implicated in interactions between proteins and single-stranded nucleic acids (Kenan et al., 1991).

Each of the two RRMs found in NONA possesses an RNP1 and an RNP2 (Rendahl et al., 1992; 1996). The more conserved RRM of NONA, situated at amino acid position 295-373 from the N-terminus (Jones and Rubin, 1990), contains 6 out of 6 amino acids of the RNP-2 consensus sequence (Bandziulis et al., 1989) and 4 out of 8 amino acids of the RNP-1 consensus sequence (Rendahl et al., 1992). Figure 3.2, taken and corrected from Rendahl et al. (1992) shows an alignment of the two RRM sequences (NONA 1 and NONA 2 in figure 3.2) of NONA with those of the yeast PABP (polyadenylate binding protein, Adam et al., 1986; Sachs et al., 1986), SXL (Bell et al., 1988) and TRA-2 (Amrein et al., 1988).

Auxiliary domains, hypothesized to mediate RNA-protein or protein-protein interactions, are also commonly found in RRM family members. These include homopolymeric glycine rich regions (Dreyfuss et al., 1993), arginine-serine repeats (RS; Zamore et al., 1992) and arginine-glycine-glycine repeats (RGG; Kiledjian and Dreyfuss, 1992). A glycine-rich region, present in many hnRNPs polypeptides and thought to have protein-protein annealing activity (Dreyfuss et al., 1993), is found near the N-terminus of nonA (see figure 3.3).
Figure 3.2.

Sequence similarities involving NONA's putative RNA binding motifs. The two RNA-binding domains of NONA (1 and 2) are aligned with the four domains of the yeast polyadenylate binding protein (PABP), the two domains of Drosophila Sex-lethal (SXL) and the single domain of transformer-2 (TRA-2). Below each block of comparisons is the "consensus" sequence, referring to the larger family of more than 30 proteins (Bandziulis et al., 1989). The RNP-1 and RNP-2 motifs are indicated in bold. Within the NONA RNPs, conserved aromatic amino acids of RNP-1 and RNP-2 are underlined, and characteristic hydrophobic residues of the RNP domains (Kenan, et al., 1991) are marked (* or *), indicating the presence of hydrophobic amino acids in one or both NONA domains.

<table>
<thead>
<tr>
<th></th>
<th>RNP-2</th>
<th>CONSENSUS</th>
<th>RNP-1</th>
<th>CONSENSUS</th>
</tr>
</thead>
<tbody>
<tr>
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<td>SQSVENSSAS</td>
<td><strong>LYVDGL-</strong>-EPS</td>
<td>VSEAHLYDIF</td>
<td>SP-IGSVSSI</td>
</tr>
<tr>
<td>PABP 2</td>
<td>PSLRKGSQSN</td>
<td><strong>IFIKRL-</strong>-HPD</td>
<td>IDNKAUYDF</td>
<td>SV-FPGLSS</td>
</tr>
<tr>
<td>PABP 3</td>
<td>LEETKATYTN</td>
<td><strong>LIVRMI-</strong>-NSE</td>
<td>TTDECFQELF</td>
<td>AK-FPGLVA</td>
</tr>
<tr>
<td>PABP 4</td>
<td>EKRMKYQGNV</td>
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<td>VDEKLEEF</td>
<td>AP-VGPTISA</td>
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<tr>
<td>SXL 1</td>
<td>MNDPRASMTN</td>
<td><strong>LILVNYL-</strong>-PDQ</td>
<td>MTDFEYALF</td>
<td>RA-IPINTC</td>
</tr>
<tr>
<td>SXL 2</td>
<td>PGESEIKDTN</td>
<td><strong>LYVNLN-</strong>-PRK</td>
<td>ITTFDQLDTF</td>
<td>GK-YGTVQK</td>
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<td><strong>LGVPFL-</strong>-NTN</td>
<td>TSHKTVRELF</td>
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<tr>
<td>NONA 1</td>
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<td><strong>LYVNLN-</strong>-TNT</td>
<td>ITTDLREMF</td>
<td>KP-YGEISI</td>
</tr>
<tr>
<td>NONA 2</td>
<td>--------NATI</td>
<td>LRVSNL-**-TPF</td>
<td>VSNELEYKSF</td>
<td>EI-FPGLER</td>
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**CONSENSUS**

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<tbody>
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<td>PABP 2</td>
<td>S-KGPFVHF</td>
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<td>EKHDAVKAV</td>
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<td>S-KGPFVCFV</td>
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<td>S-FYGAFVF</td>
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<td>SXL 2</td>
<td>P-RGFVAVVR</td>
<td>NKRREEAQEI</td>
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<td>TRA-2</td>
<td>S-RGFCFYYP</td>
<td>EKLSDARAAK</td>
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<tr>
<td>NONA 1</td>
<td>D-KMETFLEKV</td>
<td>DYHPNAEKAIA</td>
</tr>
<tr>
<td>NONA 2</td>
<td>H-MQGTVHEF</td>
<td>AKXSASSACGL</td>
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**CONSENSUS**

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</thead>
<tbody>
<tr>
<td>PABP 1</td>
<td>R</td>
<td>YA</td>
<td>Y</td>
</tr>
</tbody>
</table>
Figure 3.3. Schematic representation of the principal protein domains in NONA. Amino acids are represented by a one-letter code. The two RRM domains are indicated, and the darker bars within represent the RNP2 and RNP1 motifs. +/- indicates the charged domain. The black box shows the position of the Glycine repeat stretch.

The more recent alignment of NONA with novel sequences from the database (Rendahl et al., 1996) resulted in a larger region of homology than previously identified. A conserved region of about 320 amino acids (called DBHS, Dong et al., 1993) was found between NONA, the human splicing factor PSF (Patton et al., 1993), the HeLa protein p54\textsuperscript{nb} (Dong et al., 1993), the murine NON-POU containing Octomer binding protein or NONO (Yang et al., 1993), and nAhomo (M. Palazzolo, unpublished material). Figure 3.4, from Rendahl et al. (1996), presents an alignment between NONA and the four DBHS domain-containing proteins isolated so far. Table 3.1 reports the percent identity between each pair of proteins (Rendahl et al., 1996).

Table 3.1. Percent identities among NONA and related DBHS containing proteins. These were derived from the 311-amino acid region that spans the two RRMs and the adjacent region of relatively high charge density. From Rendahl et al. (1996)

<table>
<thead>
<tr>
<th></th>
<th>dr-NONA</th>
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<th>hu-p54\textsuperscript{nb}</th>
<th>dr-nAhomo</th>
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<td>71</td>
<td>72</td>
<td>42</td>
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<tr>
<td>dr-NONA</td>
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<td>43</td>
<td>80</td>
<td>41</td>
</tr>
<tr>
<td>mu-NONO</td>
<td></td>
<td>99</td>
<td>41</td>
<td></td>
</tr>
<tr>
<td>hu-p54\textsuperscript{nb}</td>
<td></td>
<td></td>
<td>41</td>
<td></td>
</tr>
</tbody>
</table>
Figure 3.4. Amino acid sequence alignment of NONA and related RRM-containing proteins. Bold lines above the alignment denote the two RRMs shared by this subfamily. The boxes within the bold lines indicate the RNPI and RNPII motifs. The consensus line was formed by placing a letter under any column in which at least 4 out of 5 residues are identical. Identical residues in a given column are blanked out, whereas nonconsensus residues are in lower case.
This chapter describes the identification, the cloning and sequencing of the *D.virilis nonA* homologue. The complete coding sequence of the gene was obtained, plus an additional ~2 kb of 5' upstream promoter sequence. Comparisons between the *D.virilis* and the *D.melanogaster* sequences was performed. Given the approximate 40 (Moriyama, 1987) to 60 (Beverley and Wilson, 1984) million years divergence between *D.virilis* and *D.melanogaster*, sequence conservation should highlight regions of possible functional importance.

**METHODS**

*D.virilis* genomic library.

A *D.virilis* genomic library constructed by Ron Blackman was kindly donated by John Belote. Genomic DNA had been isolated from adult flies and partially digested with *MboI*. Size-selected fragments (15-20Kb) had then been inserted into the *BamHI* sites of the EMBL3 phage vector and packaged *in vitro*.

*D.virilis* PCR fragment

A small fragment of the *D.virilis nonA* gene was amplified by PCR after trying various combinations of primers all based on the second exon sequence of the *D.melanogaster nonA* gene (Jones and Rubin, 1990). Table 3.2 shows the sequence and position (relative to the *D.melanogaster nonA* sequence) of these primers. Only primers 5'C and 3'A worked after lowering the annealing temperature of the PCR to 50°C. PCR reactions were performed as described in chapter 2.
Table 3.2. Primers based on the second exon sequence of the *D.melanogaster nonA* gene. Their position based on the *nonA* sequence from Jones and Rubin (1990) is indicated.

<table>
<thead>
<tr>
<th>PRIMER</th>
<th>SEQUENCE</th>
<th>POSITION</th>
</tr>
</thead>
<tbody>
<tr>
<td>5'A</td>
<td>5'-AAGAACAATGAACCGGCAAC-3'</td>
<td>3617-3636</td>
</tr>
<tr>
<td>5'B</td>
<td>5'-CGATGAACAGCACCAATATG-3'</td>
<td>3948-3967</td>
</tr>
<tr>
<td>5'C</td>
<td>5'-CGCGAGATGTTCAAGCCATA-3'</td>
<td>4163-4182</td>
</tr>
<tr>
<td>3'A</td>
<td>5'-GCCCTCTCGATGGGACCAAA-3'</td>
<td>4422-4403</td>
</tr>
<tr>
<td>3'B</td>
<td>5'-TCCTGGTTGAAGTCGGGCAT-3'</td>
<td>4644-4625</td>
</tr>
<tr>
<td>3'C</td>
<td>5'-CCTTGACGCAACAGTTTCAGT-3'</td>
<td>4839-4820</td>
</tr>
</tbody>
</table>

The *D.virilis* 261 bp PCR fragment was gel purified using the QIAEX gel purification kit (QUIAGEN Inc.) and directly sequenced with the dideoxy-chain-termination method (see chapter 2). Analysis of the sequence strongly suggested that the fragment was part of the putative *D. virilis nonA* second exon. Figure 3.5 shows an alignment between the *D.virilis* PCR fragment and the homologous fragment from the *D.melanogaster nonA* sequence. 20 ng of the *D.virilis* PCR product were labelled with $^{32}$P by random priming (as described in chapter 2), and used as a probe to screen the *D.virilis* genomic library.

**Library screen.**

The *D.virilis* library was plated using the *E.coli* strain Y1090. About 3x10^5 plaques were screened at high stringency (0.1% SDS; 0.5X SSC at 65°C) isolating 8 putative positive clones. Only two of them proved to be positive after successive rescreening. DNA was obtained from one of them (2.2b1) using the Promega Magic Lambda Preps DNA purification System Kit (see chapter 2).
Figure 3.5. Sequence of the *D. virilis* PCR fragment aligned to the homologous fragment of the *D. melanogaster nonA* gene. The primers used to amplify the *D. virilis* DNA fragment are indicated by the arrows. The nucleotide numbers shown for the *D. melanogaster* sequence are equivalent to those of the *nonA* sequence in Jones and Rubin (1990).
Subcloning and sequencing.

Clone 2.2b1 was cut into 5 fragments using the restriction enzyme SphI (data not shown), run on a 0.8% agarose gel and Southern blotted. When a 9Kb XbaI-EcoRI DNA fragment containing the *D.melanogaster nonA* gene (Jones and Rubin, 1990) was used as a probe, only two of the five SphI fragments hybridized to it. Of these two, the larger one (~6.5 Kb) hybridized very strongly to the *melanogaster* gene, while the smaller (~6 Kb) produced only a very faint band in the autoradiograph (figure 3.6). Both fragments were subcloned into the vector pUC 18 (Vieira and Messing, 1982) at the SphI site and were respectively named pUC39 (the 6.5 Kb fragment in pUC18) and pUC7 (the 6 Kb fragment in pUC18).

The Universal and Reverse primers, annealing respectively to the 5' and 3' side of the pUC18 polylinker, were used to sequence both ends of the pUC7 and pUC39 clones. It was found that pUC7 contained the first half of the putative *D.virilis nonA* gene, up to roughly a third of the second exon. The orientation of this insert was the same as the pUC vector. The larger pUC39 clone contained the remaining half of the gene in opposite orientation to pUC. New primers were designed based on the sequence obtained from the second exon in order to obtain its complete sequence. These primers were designed in a way to allow sequencing of overlapping regions of the exon thus assuring repeated sequencings of both DNA strands (see figure 3.7).

A strategy of cutting pUC7 and pUC39 with different enzymes, and religating the vector containing a shorter fragment of the insert was used in an attempt to obtain sequence from any of the other exons. A restriction map of the putative *D.virilis nonA* gene was also generated (figure 3.8). After cutting pUC7 with EcoRI, three fragments were produced with the approximate sizes of 1 Kb, 1.5 Kb and 6.5 Kb. The 6.5 Kb fragment was composed of the 3 Kb pUC+a 3.5 Kb insert. This 6.5 Kb fragment was recircularized using the T4 Ligase enzyme (for ligation techniques see chapter 2) and used to transform *E.coli* cells (XL1-Blue, Stratagene) as described in chapter 2. Plasmid DNA was isolated as described in chapter 2. The Reverse primer, annealing in 3' of the pUC polylinker was used to sequence the 3' end of the 3.5 Kb insert in pUC. The sequence obtained was compared to the *D.melanogaster nonA* sequence and
Figure 3.6.

Southern blot with the DNA from the positive λ clone 2.2b1 hybridized to the ~9 Kb D. melanogaster nonA gene. Each lane represents different digestions of the DNA and the enzymes used are indicated. The ~6.5 Kb and ~6 Kb SphI fragments which were subcloned into pUC18 are indicated by the arrows. Markers were not included in the blot, so the approximate positions of the λ HindIII fragments were drawn.
Figure 3.7.
Scheme illustrating the overlapping DNA fragments sequenced in order to obtain the complete sequence of the *D. virilis nonA* gene. Exons are indicated by the filled boxes and appropriately numbered. Introns (which are not to scale) are indicated by the thin lines. Each long arrow represents the average 200 bp sequenced using each primer. The primer name is written on top of each arrow. Small red arrows under the exons represent the position of the primers (whose names are indicated) used to amplify and sequence *D. virilis nonA* cDNA fragments in order to establish the exact exon-intron boundaries.
Figure 3.8.
Restriction map of the putative *D. virilis* nonA gene. Filled boxes represent the exons. Horizontal lines indicate the introns and 5' and 3' untranslated sequences. The long vertical line in the middle of the figure represents the *SphI* site which divides the gene into the two 6 Kb and 6.5 Kb fragments cloned into pUC (pUC 7 and pUC 39, see text). The sizes of the first two introns is indicated in bold.

**pUC7**

**pUC39**

*S = SphI

E = *EcoRI*

H = *HindIII*

B = *BamHI*

X = *XbaI*

Sal = *SalI*
showed homology in a region towards the end of the nonA gene first exon. New primers extending the sequence in 5' and 3' were designed in order to complete the sequence of the putative D.virilis nonA first exon.

Sequence from the D.virilis nonA putative fifth exon was obtained simply by using a primer based on the sequence from the beginning of exon five of D.melanogaster nonA at position 7106-7125 (Jones and Rubin, 1990) (5'5: 5'-AGGACTCCTCGCTTTTGGA-3'). The sequence of this exon was subsequently confirmed several times as described below.

To obtain sequence from the third and fourth exons without having to sequence also the introns, I opted for an approach which employed RT-PCR (see chapter 2). RNA was isolated from D.virilis flies using the RNAzol™B method (see chapter 2) and reverse transcribed using a 3' primer binding to to the middle of the fifth exon (3'SphI: 3' GCGGCATGCTGCCCTTTGATCATAT 5', see figure 3.7). A 510 bp cDNA fragment was then amplified using the 3'SphI primer and a 5' primer annealing to the end of the second exon(5'S/>M: 5' TATGCATGCAGATGGAGTATGCTCGCT 3'). Both primers had an SphI site (underlined) allowing cloning of the 510 bp PCR product into PUC18. The Universal and Reverse Primers, in conjunction with the primers used for the cDNA amplification, were used to sequence the third, the fourth, and part of the fifth exons. These sequences were then reconfirmed after designing new primers annealing to the beginning and the end of the third exon (5'3EXa, 3'3EXa, and 5'3EXb, see figure 3.7, and appendix 3.1) and to the beginning and the end of the fifth exon (5'5EXa, 3'5EXa, figure 3.7, appendix 3.1) and using the genomic clone as a template DNA (pUC39). Comparison between the genomic and the cDNA sequences allowed the localization of the intron-exon boundaries from the third exon to the end of the gene.

The exact intron-exon boundaries between the first and the second exons were identified using again RT-PCR. D.virilis RNA was isolated, and cDNA produced using a 3' primer annealing to the the beginning of the second exon (5'SphI: 5'-TATGCATGCTGCCCTGGCTTTGGAAGTTTGGA-3', see figure 3.7). A 225 bp cDNA fragment was then amplified using the above 3' primer and a 5' primer annealing to the
end of the first exon (E15'SphI: 5'-TATGCATGCGCGAACAAGGGTGGATT-3', see figure 3.7). Thanks to the SphI sites designed in each primer (underlined in the primer sequence), the PCR product could be cloned into the SphI site of pUC18. The Universal, Reverse, E15'SphI and E23'SphI primers were used to sequence the PCR fragment. The sequence obtained from the cDNA was then compared to the one obtained from the genomic clone in order to identify the exact intron-exon junctions.

**Sequence of the D.virilis nonA promoter region.**

A ~ 2 Kb DNA sequence upstream of the putative first codon of the D.virilis nonA gene was obtained by automatic sequencing performed at the University of Padova, Italy. The pUC7 clone was used as the genomic DNA template and the sequence was progressively extended starting from the beginning of the putative D.virilis nonA first exon.

**Computer analyses.**

All sequence analyses were performed with the programs of the GCG package for molecular biology (Version 7; University of Wisconsin Genetics Computer Group, Madison, WI, USA; Devereux et al., 1984).

Sequence alignments were generated either with the Bestfit or the Gap GCG programs. The first was used for aligning protein sequences, while the second produced the DNA sequences alignments. Dot matrix analysis were carried out with the Compare and Dotplot programs. The secondary structure of proteins was predicted using the programs Pepplot, Peptidestructure, and Plotstructure. Codon usage patterns were studied with the Codonfrequency program. Sequence simplicity analysis was performed by J. Hancock using SIMPLE34 (Hancock and Armstrong, 1994), a modified version of Tautz et al.'s (1986) SIMPLE program. The TF
SEARCH program (from the Internet, http://pdap1.trc.rwcp.or.jp/htbin/nph-tf; Akiyama, 1995) was used to search for functional motifs in the 5' regulatory region of nonA. Possible protein motif were investigated with the PSITE program (Version 1, Solovyev and Kolchanov, 1994) from the Internet (http://dot.imgen.bcm.tmc.edu:9331/pssprediction/Help/psite.html).

RESULTS & DISCUSSION

Structure of the *D.virilis* nonA gene

The five-exon-four-intron organization found in the *D.melanogaster* nonA gene is conserved in the *D.virilis* homologue (figure 3.8). The intron-exon boundaries are also conserved, as revealed by the cDNA and genomic DNA comparison analysis (see methods). Although the sequence of the four introns of the *D.virilis* nonA gene was not obtained, the approximate lengths of the first and second introns were calculated. This was done by measuring the length of PCR products obtained using primers annealing to the exon boundaries. Both introns are considerably longer than their *D.melanogaster* counterparts. The first, being ~2 kb long, is about twice the size of the first *D.melanogaster* nonA intron, the second, also measuring ~2 Kb, is about four times the size of the *D.melanogaster*'s. The exact length of the *D.virilis* nonA intron three was not directly calculated, however this intron appears to be as short as that of *D.melanogaster*. In fact a sequencing reaction using a primer annealing to the end of the third exon (5'3Exb, see figure 3.7) produced a sequence of the last 40 bases of intron three. The upstream intron sequence could not be read being too close to the 5'3Exb primer. Assuming that on average the first ~30 bases from a given sequencing primer cannot be read, I calculated the length of the third intron to be no more than 70 bases. The length of the fourth intron is unknown.
Jones and Rubin (1990) hypothesized the existence of a second splice form of NONA which would result from the translation of the beginning of the fourth intron and would add an extra 33 amino acids at the end of the fourth exon (see figure 3.1, FormII). Although the existence of this form II of NONA is doubtful (see introduction), I decided to check whether translation of the beginning of the fourth exon in D. virilis nonA could produce an extra stretch of amino acids as in D. melanogaster. The sequence of the first ~200 bases of the fourth D. virilis nonA intron was therefore obtained and translation in the three possible frames produced by the program Map of the GCG packet. The sequence is presented in appendix 3.2. 14 amino acids can be translated from the beginning of the fourth intron of D. virilis in the same reading frame of the end of the fourth exon. This stretch of amino acids is half the size of that predicted in D. melanogaster and the sequence, apart for the first two residues, is completely different. Although RNA studies, possibly via RT-PCR, could be done in order to check whether a second splice form with this extra 14 amino acid sequence at the end is actually produced from nonA in D. virilis, it seems unlikely.

The 5' regulatory region of nonA.

Figure 3.9 presents an alignment of the ~2 Kb 5' DNA sequence containing the putative promoter region of the D. virilis nonA gene aligned with the corresponding region of D. melanogaster nonA. The alignment was performed with the Gap program of the GCG packet (see methods), which finds the alignment of two complete sequences that maximizes the number of matches and minimizes the number of gaps. According to this alignment the two sequences share a 55.9 % identity.

A dot matrix analysis (window: 21; stringency:14) was performed between the D. virilis and the D. melanogaster sequences to graphically highlight the regions of homology (figure 3.10a). The areas of good conservation between the two sequences should reflect functional importance, given the long time of divergence (40 to 60 Myr) that separates D. virilis and D. melanogaster (Beverley and Wilson, 1984; Moriyama, 1987). The dotplot reveals seven regions of moderate-to-good conservation which are
Figure 3.9. Alignment of the 5' regulatory regions of *D. virilis* and *D. melanogaster nonA*. The putative transcription factor binding sites are highlighted in colour.
transcription start (Stanewsky, 1993)
Figure 3.10. Dot plot comparisons of the nonA promoter region. A: \textit{D. virilis} versus \textit{D. melanogaster}. B: \textit{D. virilis} versus itself. C: \textit{D. melanogaster} versus itself.
interspaced by regions of complete divergence between the two sequences. In addition, the terminal third of the sequences appears to be very diverged.

The dot plot analysis was repeated by plotting each of the two sequences against itself (figure 3.10b and 3.10c), this time using a lower stringency (window: 35, stringency: 19). This gives an idea of the internal structure, in terms of repeatability, of each sequence. From the dotplot results it seem that repetitive DNA does not play a major role in the sequence composition of the *D.melanogaster* promoter region (figure 3.10c). However, the promoter of *D.virilis nonA* seems to have a considerable amount of repetitive DNA (figure 3.10b) which is concentrated in the regions which mostly differ from the *D.melanogaster* sequence (figure 3.10a).

When homologous genes from different species are compared, it is not rare to observe that areas of of high divergence are often associated with regions of highly repetitive DNA. This phenomenon has been noted not only in coding sequences, for example the coding sequence of the *hunchback (hb)* gene (Treier *et. al.*, 1989) and the Threonine-Glycine region of the *per* gene (Peixoto *et. al.*, 1993), but also in regulatory regions such as the promoter an spacer regions of the ribosomal RNA genes (Tautz *et. al.*, 1986, Tautz *et. al.*, 1987; Hancock and Dover, 1988). The promoter region of *hb* has been recently characterized in four different species (*Musca domestica, D.melanogaster, D.virilis* and *Tribolium castaneum*) and high levels of short DNA motif repeats (simple DNA, Tautz *et. al.*, 1986) were noticed in all the four species' promoter sequences (Hancock *et. al.*, 1996). Events such as slippage and gene conversion seem to be the main molecular processes generating the rapid turnover of such redundant DNA sequences (Dover *et. al.*, 1982; Dover, 1993; Tautz *et. al.*, 1986).

The TF SEARCH program (Akiyama, 1995) was used to analyze the 5' regulatory region of the *D.melanogaster* and *D.virilis nonA* genes. This program highlights small nucleotide motifs homologous to known transcription factor binding sites from the TFMATRIX transcription factor binding site profile database of Wingender *et. al.* (1997). Each motif is presented with a score of identity to known motifs in the database. Short motifs (< 10 nucleotides) were considered in the analysis only if scoring > 90, while longer motifs (> 10 nucleotides) were considered if they
scored > 85. The motifs found in each sequence and the number of times they occurred in the sequence are listed in table 3.3. Figure 3.9 highlights the position of the motifs found in each of the two aligned sequences.

Table 3.3. Transcription factor-binding motifs in the *D.virilis* and *D.melanogaster* promoter region.

<table>
<thead>
<tr>
<th></th>
<th>HSF</th>
<th>DF/D</th>
<th>HB</th>
<th>DL</th>
<th>BRC-Z</th>
<th>CF2</th>
<th>ABD-B</th>
<th>SU(H)</th>
<th>FTZ</th>
<th>KR</th>
<th>SN</th>
<th>BCD</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>vir</em></td>
<td>23</td>
<td>14</td>
<td>10</td>
<td>1</td>
<td>9</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td><em>mel</em></td>
<td>27</td>
<td>9</td>
<td>2</td>
<td>4</td>
<td>4</td>
<td>3</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
</tbody>
</table>

Both sequences have high numbers of motifs homologous to the heat shock factor binding motif (HSF). These are distributed throughout the whole sequence of *D.virilis* and *D.melanogaster* regulatory regions in both orientations. In *Drosophila* the HSF binding element is only five bases long and its consensus sequence is NGAAN, where the first nucleotide (N1) is usually an A and the three nucleotides GAA are highly conserved (Fernandes et. al., 1994). A typical heat shock factor binding signal is constituted by several HSF units with alternate orientations. It has been shown *in vitro* that at least two adjacent HSF units are necessary to obtain a stable binding with the heat shock transcription factor. The distance between adjacent HSF units seems to vary in different genes (Fernandes et. al., 1994). In *D.melanogaster* there are five pairs of HSF units (with a score > 90) in which the two HSF motifs are separated by less than 15 nucleotides. Similarly, in *D.virilis*, five pairs of nearby (< 15 nucleotides apart) HSF motifs are found. The presence of these signals could be sufficient to promote binding of heat shock transcription factors in the promoter regions of both nonA genes. This would assure production of NONA protein during stress conditions. It has to be noted, however, that, given the small size of the HSF motifs, homology between these motifs and regions in the examined sequences could be purely coincidental.
Nine motifs with homology (score > 85) to the homeotic protein Deformed binding motif were found in the *D.melanogaster* 5' sequence, and 14 were found in *D.virilis*. Within each sequence these motifs are found at regular intervals, and interestingly, on three occasions they are found in homologous positions within the two aligned sequences (see figure 3.9). The conservation of these motifs, whose sequence is 17 nucleotides long, may be more than coincidence, and could have functional implications. The same could be said about a *hb* product putative binding domain (score > 85) which was found at exactly the same position in the two *D.melanogaster* and *D.virilis* sequences (figure 3.9). However, it has to be noted that these homologies in the position of functional motifs all depend on the alignment of the two sequences. In fact, comparing the position of functional motifs acquires significance only if we assume that the alignment of the two examined sequences with the Gap program is indeed correct.

One other *hb* binding domain is found in the *D.melanogaster* sequence, and nine others in *D.virilis*. In the anterior part of the *Drosophila* embryo, levels of the *hb* gene product control the transcription of the gap gene *Krüppel* (by repressing it when at high concentrations, and activating it at lower concentrations) and simultaneously repress the transcription of the abdominal structures-forming gap genes *knirps* and *giant* (Rivera-Pomar and Jackle, 1996).

Four putative dorsal (*dl*) binding motifs with an identity score > 85 (the last two of them with score > 90) are present in the sequence of *D.melanogaster* (see figure 3.9) while only one (score > 85) is found in the *D.virilis* sequence in a region that is not alignable with the *D.melanogaster* sequence. In *Drosophila*, dorsal (*dl*) contributes to determining the destiny of the different embryonic regions along the dorso-ventral axis (Ray and Schüpbach, 1996). The product of *dl* is localized in the nucleus only on the ventral side of the embryo. Dorsal both activates and represses gene expression. It activates the genes *twist* (*twi*) and *snail* (*sna*), which are required for the development of ventral structures, and it represses the genes *decapentaplegic* (*dpp*) and *zerknüllt* (*zen*), which are required for the development of dorsal structures (Steward and Govind, 1993; Rusch and Levine, 1996).
Four motifs in the *D. melanogaster* sequence and nine in *D. virilis* are homologous (score > 85) to the sequence recognized by the transcription factors produced by the *Broad-Complex (BR-C)* genes. The transcription factors produced by the *BR-C* are known to bind and activate transcription of the glue protein gene *Sgs-4* in third instar larvae (Kalm *et al.*, 1994). The NONA protein also binds to the promoter of the *Sgs-4* gene when this is being actively transcribed (Saumweber, 1990; Besser *et al.*, 1990). It has been hypothesised that NONA could be processing the nascent RNA as it is being transcribed from the glue protein gene (Besser *et al.*, 1990). Could it be that the *BR-C* transcription factors activate the *Sgs-4* gene and, by promoting expression of the *nonA* gene as well, also ensure that the transcripts of the glue protein gene are properly processed?

Several other transcription factors binding motifs were found in both *D. melanogaster* and *D. virilis* sequences (such as the motifs for binding of the products of the genes: *Chorion factor 2 (Cf2), fushi tarazu (ftz), Abdominal B (Abd-B)* and *Suppressor of Hairless (Su(H)),* see table 3.3). However in all these cases the motifs were found in very different locations within the two sequences (see figure 3.9). It is not possible to speculate on the possible functional significance of these motifs. However it has to be mentioned that, in few instances, certain motifs scored very highly (> 90) for identity with known binding motifs. For example, the putative *ftz* binding motif found in the beginning of the *D. melanogaster* sequence (figure 3.9) had a score of 94.3. Similarly, the *Su(H)* binding domain identified in the sequence of *D. virilis* scored 90.4. Given that the *ftz* binding motif is 12 nucleotides long, and the *Su(H)* motif measures 15 nucleotides, their presence in these promoters could reflect some functional importance, however protein-DNA mobility assays should be used to shed light on these possibilities (e.g. Guille and Kneale, 1997).

Finally, several binding motifs were found exclusively in the *D. melanogaster* sequence (those for the gene products of *Kruppel (Kr), singed (sn) and bicoid (bcd)*). In particular, one domain homologous to the *bcd* binding motif was assigned a very high score (96.2), as was a *Kr* putative binding motif (score = 93.5). Again, *in vivo*
and in vitro approaches could identify whether any of these binding sites are functional (Guille and Kneale, 1997; St Johnston and Nüsslein-Volhard, 1992).

Lastly, several sequences homologous to the TATA box consensus were localized by the TF SEARCH program. Only those with the appropriate orientation for the nonA gene are presented in figure 3.9. In the D.virilis promoter three putative TATA boxes are recognized, however the last one presents the highest homology (score = 88) to the consensus sequence (Brethnach and Chambon, 1981). Since the position of the transcriptional start site is not known in D.virilis nonA, it is not possible to speculate on the position of the TATA box element. In the sequence of D.melanogaster the only TATA box found by the program does not correspond to the one predicted by Besser et. al. (1990) and does not fit with the position of the transcription start site localized by S1 mapping by Stanewsky et al.(1993) (nucleotide 1828 in the genomic sequence published by Jones and Rubin, 1990; indicated in figure 3.9). Approximately 30 bp upstream of the start of transcription (Stanewsky et al., 1993), there is however a TATAAATA motif (highlighted in figure 3.9), that the TF SEARCH program fails to detect, and which is likely to be the real TATA box in D.melanogaster. The program Findpattern of the GCG packet was therefore used as an additional means to localize sequences homologous to the TATA box consensus motif (Brethnach and Chambon, 1981) in the D.virilis promoter. All sequences homologous to the TATA A/T A/T A/T motif were searched with the aid of this program. Several homologous motifs were found, however only four, located in the terminal part of the sequence (after nucleotide 1000), were considered (respective positions: 1105, 1216, 1725, 1877, boxed in figure 6.9). Of these, the motif at position 1725 seems particularly interesting, since it was recognized also by the TF SEARCH program and was assigned the highest score among the possible TATA box motifs (see above).

The nonA region is genetically complex. An essential gene called l(1)I19e is associated with the nonA gene and partially overlaps with it so that the two genes cannot be molecularly separated (Stanewsky et al., 1993). At present, only point mutations or small deletions 3' of a Sall site in the middle of the nonA second exon can
be assumed to affect the nonA gene only (Stanewsky et al., 1993). This means that all the sequence upstream of the middle of the second exon of nonA must be shared between the two genes. Therefore the 5' upstream DNA region of nonA that has been studied here could also be a functional part of the l(1)19e gene. If in D.virilis the same overlap exists between nonA and the l(1)19e gene as in D.melanogaster, then the homology between the D.melanogaster and the D.virilis 5' regulatory regions could be reflecting functional importance not only for the nonA gene, but also for l(1)19e.

The coding sequence of the D.virilis nonA gene

Figure 3.11 shows the nucleotide and amino acid sequence of the translated region of the D.virilis nonA gene. Arrows indicate the position of the introns. A dot matrix comparison (window: 21, stringency: 14) between the D.virilis and the D.melanogaster nonA coding sequences is presented in figure 3.12a and reveals an area of considerable divergence covering approximately the first half of the gene. This is followed by a region of very high homology between the two genes which extends for almost all the remaining length of the sequences. The Gap program was used to align the two sequences (appendix 3.3). After alignment, the two sequences shared 72.4% identity. Not surprisingly, the area of high homology corresponds to the sequence for the RNA-binding domains (RRM1 and RRM2) and the charged domain (see Introduction). In addition, the sequence at the end of the nonA gene seems to be quite well conserved between D.virilis and D.melanogaster. However, a number of parallel shifts of the plot line indicate regions of gaps in the alignment of the two DNA sequences. Regions of very densely packed dots are also visible; these occur in several regions at the beginning and at the end of each of the two sequences, and indicate repetitive regions that cause numerous shifts in the plot line. Note that repetitive regions at the beginning of the D.virilis nonA gene align with sequences not only at the beginning, but also in the middle and at the end of the D.melanogaster sequence, and regions at the end of the D.virilis gene are similar to sequences found approximately in the middle of the D.melanogaster gene.
Figure 3.11. DNA and amino acid sequence of the *D. virilis nonA* gene. Vertical bars indicate the position of the introns. The sequence highlighted in yellow is homologous to the DBHS domain (see introduction). The two RNA-binding domains (RRM1 and RRM2) are underlined. The RNP1 motifs are shown in blue and the RNP2 in red.

```
1  ATGGAAAATTCTGTTAAAATGGACAATTCTGGCAATTCAACACCGCTGCCACAGCGCCAA  60
M E N S V K M D N S G N S T P L P Q R Q  
61  CGAAGAGCTTAATATCAACCCGAATATATATCTGGCACAACCTGGCCACCAGAGCCAAAT  120
R R A N N Q P N K I G K L G P Q K Q N  
121  GAAGGTGCACCCGCGGACGCGGACGCGGACGCGGCGGCGGCGGCGGCGGCGGCGGCGG  240
N G G G G S V G G G G G G G G G G G G G Q  
181  AATCTAGGTGCCGCGGATTCGCGCCTGCTGCTGCGCCTGCTGCTGCTGCGCCTGCTGCTG  300
N Q N K N F A N K G G F G G G G N R N R  
241  AATCTAGGTGCCGCGGACGCGGACGCGGACGCGGCGGCGGCGGCGGCGGCGGCGGCGG  360
N R G G N Q N R S F Q N Q N Q N Q S K S  
301  ACAACGATGCCCCAACACGCGAGGGGGAATCTGACAAATCTAACGAACCTTTGGCCCAT  420
T T D A P K A D G G N L N D K S N E A N  
361  ACAACGATGCCCCAACACGCGAGGGGGAATCTGACAAATCTAACGAACCTTTGGCCCAT  480
N A N Q S N S N S A A Q A Q A Q L Q A Q  
421  GCCCAACGCTCATCCCCGCCAGCCAGCCAGCCAGCCAGCCAGCCAGCCAGCCAGCCAG  540
A Q A H A Q A Q A Q A Q A Q A H A Q A Q  
481  GCCCAACGCTCATCCCCGCCAGCCAGCCAGCCAGCCAGCCAGCCAGCCAGCCAGCCAG  600
A Q A H A H A Q N Q A F R A R G G G G  
541  GCCCAACGCTCATCCCCGCCAGCCAGCCAGCCAGCCAGCCAGCCAGCCAGCCAGCCAG  660
G G G G G G G G G G G G G G G G G G G G  
601  GGCCGTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTG  720
G G G G R D  R N P D R R G G G G G G G G Q  
661  GGAGGAGGCGGACGCTAGTAGGGAATCCGGATCGGCGTGGTGGTGGTGGTGGTGGTGGTGG  780
A Q A H A H A Q N Q A F R A R G G G G  
721  GGAGGAGGCGGACGCTAGTAGGGAATCCGGATCGGCGTGGTGGTGGTGGTGGTGGTGGTGG  840
A Q A H A H A Q N Q A F R A R G G G G  
781  CGACAGATGCTCTGCTCCAGACAGATGCTCTGCTCCAGACAGATGCTCTGCTCCAGACAGA  900
R S I S G T H E L P P I E V A Q E T K  
841  TTCTCTGCTGGACCAATCCATCCATCCATCCATCCATCCATCCATCCATCCATCCATCCAT  960
F S G R N R L Y V G N L T N D I T D E E  
901  TTGCGGAAATTTAAGCTACGCGTAAAATAGCCGAGATATTTCCTACGCGTAAAATAGCCGAG  1020
L R E M F K P Y G E I G E I F S N L E K  
961  AACTTACATTCCCATTAAGGTTAGCTATATATATATATATATATATATATATATATATATAT  1080
N F T F L K V D Y H I N A E K A K R P L  
1021  GACCGTCTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTG  1140
D G S M R K G R H V R V R F A P N A T I  
1081  TTGCGGGTACATCCATCCCATCCATCCATCCATCCATCCATCCATCCATCCATCCATCCAT  1200
L R V S N L T P F V S N E L L Y K S F E  
```
Figure 3.12. Dot plot comparisons of the nonA gene coding sequence. A: *D. virilis* versus *D. melanogaster*. B: *D. virilis* versus itself. C: *D. melanogaster* versus itself.
The dot matrix analysis was repeated after lowering the stringency (window: 35, stringency: 19) and plotting each of the two nonA sequences against itself. The output for the D.virilis gene (figure 3.12b) clearly shows numerous and large regions of repeated DNA, clustered especially at the beginning and the end of the gene. The D.melanogaster nonA gene (figure 3.12c) seems much less repetitive than its virilis homologue, however regions of repeats are still evident especially within the first third of the gene, and towards the end of it.

The frequent occurrence of several repeated short motifs in close proximity (cryptic simplicity) seems to be a widespread and important component of eukaryotic genomes (Tautz et. al., 1986). To have a more accurate idea of the extent of the redundancy in each nonA gene (either of D.melanogaster or D.virilis) and to discover which sequence motifs are involved, a cryptic simplicity analysis was performed on the D.melanogaster and the D.virilis nonA gene coding sequences. The cryptic simplicity test was carried out by John Hancock, using the SIMPLE34 program (Hancock and Armstrong, 1994). The SIMPLE34 program, which is a modified version of Tautz et. al.'s (1986) SIMPLE program, analyzes a given sequence by moving a 64 bp window along it. For each window, it finds three or four base pair motifs at its centre, and searches for duplicates of those motifs within the window. It awards one point for each duplicate of the three base motif, and an extra three points if the four base motif is duplicated. It then calculates an average score for the entire sequence based on all the window scores, which is the "Simplicity Factor" (SF). It then repeats the same analysis for ten random sequences with the same base doublet frequency as the input sequence and the same length. This generates a "Randomized Simplicity Factor" (RSF). The "Relative Simplicity Factor" (RelSF) is the ratio of SF over RSF, and indicates how much more simplicity the input sequence contains than the random sequences.

The SF of the D.melanogaster nonA coding sequence was 3.034, and as the RSF was 1.954, the RelSF of D.melanogaster nonA was calculated as 1.552. The SF of the D.virilis nonA gene was 3.687, the RSF 1.943, and the RelSF 1.898. From these values it therefore appears that the D.virilis nonA sequence is more repetitive than that of D.melanogaster. Figure 3.13a and 3.13b are the graphic representation
Figure 3.13. Results of the cryptic simplicity test for the coding sequence of nonA. A: D. virilis. B: D. melanogaster. The highest peaks represent motifs (whose sequence is indicated) found to be concentrated in the sequence to a degree unlikely to occur by chance.
for the results of the cryptic simplicity test. The high peaks found in both displays, indicate that both nonA genes have regions which are cryptically simple through the repetition of one short motif or another. The highest peaks represent the most statistically significant simple motifs (termed SSMs) in the sequence. These correspond to the motifs found to be concentrated in the sequence to a degree unlikely to occur by chance. The base compositions of these short motifs is indicated in the graphic display for each of the two sequences analyzed. Both D.melanogaster and D.virilis nonA genes present areas of simplicity especially towards the first third of the gene, even though D.virilis does so to a much larger extent. In both nonA genes there is also an additional repetitive region towards the end of the sequence, however, again, this is more pronounced in the D.virilis gene. Moreover, the very high peak towards the end of the D.virilis sequence indicates that a single motif constitutes a significantly large part of the repeats in this part of sequence. In the terminal part of the D.melanogaster sequence instead, although many repeats are found, no motif in particular is significantly over-represented.

Recent comparison of the promoter region of the hb gene (Hancock J., pers. comm.), has revealed not only that a large number of SSM motifs exists in this region of D.melanogaster, D.virilis, and Musca domestica, but also that many of these highly represented motifs are shared between the sequences of these different species. Similarly, in the coding region of D.melanogaster and D.virilis nonA some SSMs were found to be shared by the two sequences (GGTG and GTGG, see figure 3.13). However, several other SSMs were found in the D.virilis sequence but not in the D.melanogaster. A CAAT motif which is highly repeated at the beginning of nonA in D.virilis, was also found in the nonA sequence of D.melanogaster. In the D.melanogaster gene this motif is localized towards the end of the sequence and it is represented to a much lower extent. Most of the repeated motifs in nonA are constituted by cyclic permutations of the short sequence GGT or GGC. Note that GGT and GGC both code for Gly which constitutes a very large repeated region in the protein of the D.virilis nonA gene and a slightly smaller one in the protein of D.melanogaster.
Conceptual translational start of the *D. virilis nonA* gene

Figure 3.14 shows a partial sequence from the *D. virilis nonA* gene indicating the ATG motif which is believed to initiate translation. Conceptual translation of the sequence starting from this ATG produces an open reading frame with homology to that of the *D. melanogaster nonA* gene (see figure 3.15). Few other ATG motifs can be found upstream of the one highlighted in figure 3.14. However conceptual translation from these other ATGs contains multiple termination codons in all three reading frames. Moreover, the predicted ATG starting codon is in agreement with the most stringent requirements dictated by Kozak's consensus translation initiation sequence (GCCRCCATGG) (Kozak, 1987; 1991). In fact, in this case, a purine (G) is present three bases before the ATG motif, and a G is immediately following it.

Protein comparison

Figure 3.15 shows an alignment of the amino acid sequence of *D. virilis* and *D. melanogaster* NONA proteins, created with the Bestfit program of the GCG packet. This program finds optimal alignments by inserting gaps to maximize the number of matches using the algorithm of Smith and Waterman (1981). The percent of overall identity between the two is just over 75%, while the similarity is about 83%. Previous comparisons between several *D. virilis* and *D. melanogaster* homologous gene products have shown protein identities ranging between 50 and 83%. Table 3.4 shows the percent similarity and identity shared by several *D. virilis* and *D. melanogaster* proteins, calculated with the algorithm of Smith and Waterman (1981). Note that, in general, developmental gene products seems to be most conserved in these two species.

**Table 3.4.** Percent similarity and identity shared by 10 *D. melanogaster* and *D. virilis* homologous proteins.

<table>
<thead>
<tr>
<th>NONA</th>
<th>EN</th>
<th>HB</th>
<th>KNIRPS</th>
<th>NOS</th>
<th>OSK</th>
<th>PER</th>
<th>RUNT</th>
<th>SEV</th>
<th>TRA</th>
</tr>
</thead>
<tbody>
<tr>
<td>% sim</td>
<td>82.8</td>
<td>87.8</td>
<td>88.5</td>
<td>83</td>
<td>77.7</td>
<td>74.3</td>
<td>78.4</td>
<td>86.9</td>
<td>77.9</td>
</tr>
<tr>
<td>% id</td>
<td>75.1</td>
<td>82.9</td>
<td>81.2</td>
<td>74.8</td>
<td>64.2</td>
<td>59.4</td>
<td>67.6</td>
<td>80.9</td>
<td>64.5</td>
</tr>
</tbody>
</table>
Figure 3.14.
Conceptual translational start of the *D. virilis* nonA gene. a, b, and c are the three reading frames. The ATG motif believed to initiate translation is boxed in red. The first three amino acids of the *D. virilis* NONA protein are underlined in red. Other ATGs (boxed in white) are found in the ~400 nucleotides upstream, however, translation from these produces stop codons in all three reading frames.

```
AGTTTAAACTAAAATAAGTGPACCATATTTCAACAAGTTAGCTAGTTTTATTTAATTAG
a  SLN*A INVYPYSTS*AVVYLNN*-
b  V*TK*MYHIIQQVKL*FI*IS-
c  FKLNKCTIFNKLSCSLFKLV-

TTTTATTCTCTTGGGTGATTTTTTTTGGCTTAGTCTTAATTCAATTGCGAAAACTTTCT
a  FYSLSLG*FFFFLSSLNSIAKLSS-
b  FISLWVDFFGLVLIQLRNFCL-
c  LFLEGIFLASA*SFCETFC-

TTTTAAGTCTTTTATATATATATCATTTCAACACATATTTATTTATACATAGAG
a  FLYFLG*FFFWLSSLNSIAKLSS-
b  FISLWVDFGLVLIQLRNFCL-
c  LFLEGIFLASA*SFCETFC-

TTTTATCGTTAAGGAAAATATACCTGATGCTGTTTCACATCTATTTATATCGATAG
a  FIS*RENSNAFHIITLEYR*-b  SFRLGKISKIRMRFSTLSFYIDR-
c  HFVKGKYLECVSHTHYFISID-

ATCGATGCAGAAAATCAACCTGTGCAGATATAGAGTTTTTGTGTTGATGACAAACATATTG
a  IDATKSTLTLTDIELFRSHNC-
b  SMRQNPQP*IQC*NCFVPTIIV-
c  RCDKINPDRCRTVSFQPSLS-
```

```
CAATTCTACGCCTACATTTCAACGGGCTGTCAGTTGCTTTGTAATTGATTTTTTTTTT
a  HFILQFLYAGAGRLCN*FFFF-
b  ISLRFSGCTGGLDCVIDFEL-
c  FHYASVSRGWTVV*LIFF*
```

```
AAAAACATTATTAAATATCTGATTACGAAATT-
a  KTIILQSCNGK-
b  KQFYFNLVMEN-
c  NNFTSIL*WIKI-
```
The first half of the protein is clearly the most diverged (figure 3.15), with the first exon of nonA showing only 60% identity, as is the case for the first half of the second exon. Note that these regions of divergence are constituted in large part by stretches of repeats. In the first exon stretches of poly-Gly are found especially in the D.virilis protein. In the beginning of the second exon, a QN and a degenerate GNQGGX repeat found in the D.melanogaster gene have been replaced by a QA and a very long poly-Gly repeat. A long stretch of 29 Gly residues is the most striking feature of the nonA virilis gene. Several known RNA-binding proteins, such as the hnRNP proteins A1 and A2, and the nucleolar pre-rRNA-binding protein, nuclein, have auxiliary domains constituted by Glycine rich regions (Bandziulis et al., 1989). These auxiliary domains are believed to function primarily in protein-protein interactions, since all of these proteins are components of multiprotein ribonucleoprotein complexes. However, the auxiliary domains may also influence the RNA-binding activity of a protein. A comparison between the binding of A1 to single-stranded (ss) DNA and RNA with that of UP1, the truncated auxiliary domain-less derivative of A1, suggested that the auxiliary domain may modify the polynucleotide binding properties of RNA-binding domains (Bandziulis et al., 1989).

Given the functional importance of the RNA-binding domain (RRM1+RRM2) and the adjacent charged domain of NONA (Rendahl et al., 1996, Stanewsky et al., 1996), it is not surprising that these regions of the protein are very well conserved between D.virilis and D.melanogaster. Only 5 non-conservative changes are in fact found in this protein region that spans a total of 311 amino acids. The amino acid sequence encoded by the second part of exon three is also very well conserved, except for a region in which a small poly-G repeat found in D.melanogaster seems to have expanded into a much larger one in D.virilis. The small exon four encodes for a very diverged part of the protein where alignment between the D.virilis and D.melanogaster amino acid sequences is possible only in regions containing N repeats (see figure 3.15).

Perhaps more interesting is the high degree of conservation found in the amino acid sequence encoded by the fifth exon. This part of the protein in fact does not
Figure 3.15. *D. virilis* and *D. melanogaster* NONA alignment. Breaks (vertical lines) in the sequence represent intron positions. The amino acid sequence in a yellow background represents the DBHS domain (see Introduction). The two RNA-binding domains (RRM1 and RRM2) are underlined, and the RNP1 and RNP2 motifs are boxed. Putative post-translational modification sites are highlighted in colour as follows:

- Red: N-glycosylation (only homologous sites in the two proteins are shown)
- Orange: cAMP-, cGMP-dependent protein kinase phosphorylation
- Green: Protein kinase C phosphorylation
- Blue: Casein kinase II phosphorylation
- Pink: Tyrosine kinase phosphorylation
appear to have any recognizable functional motif. Translation of the fifth exon appears however to be indispensable for the functioning of the NONA protein: nonA mutant flies (either \textit{nonA} or \textit{nonA} \textit{dis}) carrying a \textit{nonA} transgene, in which an \textit{in vitro} generated stop codon prevents translation of the fifth exon, do not show any rescue of the mutant phenotypes (Jones and Rubin, 1990; Rendahl \textit{et al.}, 1992; Stanewsky \textit{et al.}, 1993).

\textbf{Protein motifs in NONA.}

The PSITE program (Version 1, Solovyev and Kolchanov, 1994) searches for functional motifs in a protein using the PROSITE database (Bairoch and Bucher, 1994). The results of the program for the \textit{D.virilis} and \textit{D.melanogaster} NONA proteins are graphically illustrated in figure 3.15.

Eight possible N-glycosylation sites are found in the \textit{D.virilis} NONA sequence, and fourteen in the \textit{D.melanogaster}. Five of these are found at homologous positions in the two aligned proteins (see figure 3.15). Interestingly, a site found at position 645 of the \textit{D.virilis} amino acid sequence (figure 3.15), is also present in the \textit{D.melanogaster} sequence in spite of being located in a poorly conserved region of the protein (the region encoded by exon four). It would seem however improbable that NONA is extensively glycosylated, since from Western Blot studies with both monoclonal and polyclonal antibodies (Stanewsky \textit{et al.}, 1993; Rendahl \textit{et al.}, 1996) the protein does not appear to be highly modified and the band it produces in the blots is of the predicted size, based on the primary sequence.

Two putative cAMP- and cGMP-dependent protein kinase phosphorylation sites are present in the \textit{D.virilis} protein, one of which is also found in the \textit{D.melanogaster} protein. Interestingly, the site found only in the \textit{D.virilis} amino acid sequence, owes its presence to a S residue which is found at amino acid position 534 of the \textit{D.virilis} protein instead of a V residue (as in the \textit{D.melanogaster} NONA). The same S residue in the \textit{D.virilis} protein creates also a possible casein kinase II
phosphorylation site which is not found in the NONA protein of \textit{D.melanogaster}. It is intriguing that the V-S substitution occurs in the \textit{D.virilis} amino acid sequence within the highly conserved RRMs + acidic domains and is one of the very few non-conservative changes in this region of the protein. This allows speculation that the V-S substitution could have some functional importance for the \textit{D.virilis} protein. Another possibly interesting site is found at position 250 of the \textit{D.virilis} amino acid sequence. It is a site for tyrosine kinase phosphorylation and it is not found in the \textit{D.melanogaster} amino acid sequence. Finally, five protein kinase C phosphorylation sites are found at homologous positions within the two aligned NONA proteins, while three more are found only in the \textit{D.virilis} protein in a region of high divergence (see figure 3.15).

\textbf{NONA secondary structure}

The secondary structure of the NONA protein of \textit{D.virilis} and \textit{D.melanogaster} was predicted using the GCG programs Pepplot, Peptide structure and Plotstructure. The output of Pepplot is shown in figure 3.16 and 3.17. The first panel of the Pepplot picture shows a schematic representation of the amino acid sequence. Each residue is represented by a line at the position where it occurs in the sequence. The lengths and colours of the lines are used to show chemically similar groups of amino acids as follows: RED= hydrophilic, charged (down=acidic, up=basic); YELLOW= hydrophilic, uncharged (short=amides, long=alcohols); GREEN= hydrophobic (short=aliphatic, long=aromatic); BLACK= Proline; UNMARKED= Alanine, Glycine, Cysteine. The second panel displays the residues which are beta-sheet forming and breaking, as defined by Chou and Fasman (1978). To produce beta-structures, there should be at least three beta-forming residues and not more than one breaking residue within a window of five. The third panel of the plot shows the Chou and Fasman (1978) propensity measures for alpha-helix and beta-sheet. As each curve rises above the threshold for its colour, it satisfies one criterion for propagation of an alpha-helix or beta-sheet structure. If the curves for alpha and beta propagation drop below the black threshold and if there is at least one breaking residue in four, then the structure
Figure 3.16. Pepplot of the *D. virilis* NONA protein.
Figure 3.17. Pepplot of the *D. melanogaster* NONA protein.
may terminate. Both curves represent the average of a residue-specific attribute over a window of four. The fourth panel displays the residues that are alpha-helix forming and breaking, as defined by Chou and Fasman (1978). For alpha helices to be produced, there should be four or more alpha-forming residues and no more than one breaking residue within six residues. The fifth panel shows regions of the sequence typically found in turns (Chou and Fasman, 1978), while the last panel shows the average hydrophobicity given by the Kyte and Doolittle hydropathy measure curve (1982). This curve is the average of a residue-specific hydrophobicity index over a window of nine residues. When the line is in the upper half of the frame, it indicates a hydrophobic region, and when it is in the lower half, a hydrophilic region.

The two plots for the *D.virilis* and the *D.melanogaster* NONA proteins show a region of clear divergence between residues at position ~150 and ~250. In this region the *D.virilis* protein shows a long array of alpha-helix-forming residues which give rise to a long region of alpha-helix conformation. This conformation is generated by the very long QA repeat that characterizes the N-terminal part of the *D.virilis* protein. This alpha-helix region is immediately followed by a large ininterrupted domain of turns, which corresponds to the long Glycine-repetitive region found in NONA of *D.virilis*. In the corresponding region of *D.melanogaster* NONA (amino acids 150-250) no large alpha helix domain can be found, and there are only short and frequently interrupted regions rich in turns. The remaining parts of the two proteins are more similar to each other: small islands with alpha-helix or beta-sheet conformation are found in both NONA proteins between residue 280 to approximately amino acid 450. Finally, from amino acid 450 to amino acid 600 there is in both proteins a large domain folded in alpha-helix conformation. Both proteins show a similar pattern of hydrophilicity which is more pronounced at the amino and carboxy-termini, leaving the middle of the protein neither hydrophilic, nor very hydrophobic.

The two programs Peptidestructure and Plotstructure (of the GCG packet, see methods) were used to generate the two-dimensional figures representing the secondary structure of the proteins (figure 3.18 and figure 3.19). Peptidestructure uses the original method of Chou and Fasman (1978) to predict helices, sheets and
Figure 3.18. Chou and Fasman secondary structure prediction of the *D. virilis* NONA protein.
Figure 3.19. Chou and Fasman prediction of the secondary structure of the *D. melanogaster* NONA protein.
turns. The minimum length of an alpha-helix is six residues and of a beta-sheet is four. Regions without adequate predictions are replaced by the conformational state of the next best probability. Glycosylation sites are also predicted for sites where the residues have the composition NXT or NXS. When X is D, W, or P, the site is taken to be a weak glycosylation site, otherwise it is a strong glycosylation site. The Plotstructure program plots the measures of secondary structure from the output file of Peptidestructure into a two dimensional representation. The two-dimensional pictures produced for the *D. virilis* and *D. melanogaster* NONAs (figure 3.18 and 3.19) nicely shows the striking difference between the two proteins from residues 150 and 250, where in *D. virilis* NONA the large region of alpha-helix conformation is represented by a sine wave, and the following uninterrupted region full of turns is indicated by 180 degree turns.

The Peptidestructure and Plotstructure programs were also used to obtain graphic representation of the secondary structure of the RNA-binding domain of the two NONA proteins, which are located approximately from amino acid 300 to 600 in each amino acid sequence (appendix 3.4 and 3.5). All RNA-binding structures determined to date reveal a structure composed by a large antiparallel beta-sheet which is sandwiched between two alpha-helices, and follow a general pattern of $\beta$-$\alpha$-$\beta$-$\beta$-$\alpha$-$\beta$ (Siomi and Dreyfuss, 1997). The functionally important structure seems to be the antiparallel beta-sheet which forms a non-sequence specific RNA-binding platform. The secondary structure of the RNA-binding domains in the *D. melanogaster* and *D. virilis* NONA proteins seems to fit with the general conformational rules for RNA-binding proteins just described: an antiparallel beta-sheet is found in both NONAs and it is sandwiched between two alpha-helices domains (appendix 3.4 and 3.5).

**Molecular evolution of the nonA gene**

We have seen that the first half of the NONA protein is quite diverged between *D. virilis* and *D. melanogaster*, while the remaining half maintains a high level of conservation. Although the high conservation of the RNA-binding domains and the
adjacent charged region is not surprising, given that the same domains are functionally important and show conservation in proteins found in such distantly related organisms such as Drosophila and humans (Rendahl et al., 1996), the divergence found in the first part of the protein could be explained in two opposite ways. Either the diverged regions of NONA are those regions of the protein which lack functional importance, and are thus free of selective constraints, or, alternatively, they serve species-specific functions and therefore their divergence reflects functional importance which could be the result of adaptive evolution.

Coding DNA sequences can undergo two types of substitutions: non-synonymus and, given the degeneracy of the genetic code, synonymous. Synonymous changes can be considered to be largely neutral, and therefore should reflect the neutral mutation rate, i.e. the ticking of the molecular clock (see Li, 1997). On the contrary, non-synonymous substitutions are likely to be mostly deleterious and therefore subjected to purifying selection. However, if a non-synonymous is advantageous, it will be fixed at a rate faster than the neutral mutation rate. When this happens, the number of non-synonymous substitutions can be greater than the number of the synonymous ones. In other words, if we calculate the number of non-synonymous substitutions (indicated as \( K_A \)) and the number of synonymous substitutions (\( K_S \)), a ratio \( K_A/K_S \) greater than one will indicate that positive selection is acting in mantaining the non-synonymous changes (Li and Graur, 1991).

The number of synonymous and non-synonymous substitutions per site between the D.virilis and the D.melanogaster nonA genes was calculated with the GCG NewDiverge program. The program is based on a variant of the method of Li et al. (1985), as modified by Li (1993) and Pamilo and Bianchi (1993). It uses a translation table to determine codon degeneracies, and applies Kimura's two-parameter method (1980) to correct for multiple hits and to account for the difference in substitution rates for transitions and transversions. The Assemble program of the GCG packet was used to remove DNA regions from each nonA sequence which could not be properly aligned, i.e. those regions which generated gaps in the proteins alignment.
The $K_A/K_S$ test was performed first on the whole coding sequence of nonA, and then in two separate domains of the sequence: the first including the most diverged part of the genes (from nucleotide 1 to 813 of each newly assembled DNA sequence), and the second constituted by the more conserved part (from nucleotide 814 to the end of each gene).

The $K_A/K_S$ test was performed also on the $D. virilis$ and $D. melanogaster$ per, tra and engrailed (en) homologous genes. Table 3.5 shows the results of the test for these genes, including the results obtained for the timeless (tim) gene by Ousley et al. (1998). It is clear that in all these genes the rate of non-synonymous substitutions is occurring at a much lower rate than the synonymous level. It is interesting that tra, known for the high divergence it displays across different species (O'Neil and Belote, 1992), shows the highest $K_A/K_S$ of all the genes here studied. The nonA gene displays a lower ratio than tra even when the most diverged part of the gene only is taken into consideration.

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<td>0.213</td>
<td>0.153</td>
<td>0.126</td>
<td>0.197</td>
<td>0.081</td>
<td>0.325</td>
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</table>

It has to be pointed out that the $K_S$ measure is likely to be an underestimate of the neutral mutation rate. It has been shown, in fact, that preferences in the usage of codons might exist, especially for highly expressed genes (Shiels et al., 1988, Sharp et al., 1988), these preferences being driven probably by the availability of different types of tRNAs (Kurland, 1987; Sharp et al., 1988). Biases in codon usage are inversely
proportional to the silent mutation rate, and have been shown to slow down the rate of silent substitutions by as much as one half in some genes (Sharp and Li, 1989).

Rapid divergence driven by positive selection has rarely been demonstrated at the molecular level. One of the few examples is presented in a recent study on the evolution of the lysozyme gene in primates (Messier and Stewart, 1997). Using the $K_a/K_s$ ratio as a tool to identify positive selection, Messier and Stewart have been able to show that adaptive changes occurred in the lysozyme gene. They identified two separate episodes of positive selection: one on the lineage leading to the common ancestor of the colobine mokeys, and a second, unexpectedly, on the lineage leading to the common ancestor of hominoids.

Genes encoding functional proteins are always expected to be subjected to some degree of purifying selection, eliminating most of the amino acids changes because of their deleterious effects. Whole-gene comparisons of $K_a/K_s$ ratios reflect both positive and negative selective forces, consequently the value of the ratio may often be less than one even if some adaptive substitutions have occurred. For example, when 363 homologous genes were compared between the mouse and rat, only one had a ratio higher than one (Wolfe and Sharp, 1993). It would be surprising however if only one of all those genes had undergone adaptive changes during the divergence of the two species (Sharp, 1997). One way to overcome this problem is to know where to look for adaptive evolution within a gene sequence. Knowing in advance which sites within a protein are good candidates for adaptive changes, would allow focusing only on the corresponding part of the sequence, without including in the analysis many other positions subjected to purifying selection. An example of this is the study of the class I major histocompatibility complex (MHC) genes (Hughes and Nei, 1988). Based on the knowledge of which residues of the protein are part of the antigen recognition site (ARS), Hughes and Nei (1988) were able to show that these sites specifically have a $K_a/K_s$ ratio greater than one, while in the remaining parts of these genes the ratio is always considerably lower. Similarly, analysis of the $K_a/K_s$ ratio in the homeodomain of the rodents homeobox gene, Pem, has shown that the divergence of this specific domain was driven by adaptive selection (Sutton and Wilkinson, 1997).
It is now recognized that most nuclear genes in *Drosophila* evolve at rates more than two times faster than those in rodents and other mammals (Moriyama, 1987, Moriyama and Gojobori, 1992, Sharp and Li, 1989). Moreover, the rate of synonymous substitutions can vary between different genes of *Drosophila* (Sharp and Li, 1989). The rate of nucleotide substitutions depends on the mutation rate (Kimura, 1983), the generation time (or the number of replications), and the degree of selective constraints on the nucleotide sequence when the changes are not selectively neutral (Moriyama and Gojobori, 1992). One explanation for the higher rate of synonymous substitutions in *Drosophila* compared to mammals could be the shorter generation time of *Drosophila*, which implies more DNA replication per year (Britten, 1986; Li and Tanimura, 1987; Kimura, 1983; 1991). Differences in the accuracy of DNA replication and efficiencies in DNA repair could also account for the different substitution rates between genes of *Drosophila* and mammals (Moriyama and Gojobori, 1992).

A consistent explanation for the high and variable synonymous substitution rates in *Drosophila* has not been provided so far. However several pieces of evidence suggest that the rates of synonymous substitutions are correlated with the base composition of the nuclear genes. It seems that the G+C and C content at the third codon position of nuclear genes is negatively correlated with synonymous substitution rates, while the A content is positively correlated (Moriyama and Gojobori, 1992). Moriyama and Gojobori propose two possible explanations for the observed relationships. The first is that spontaneous mutations directed from C to A or T may frequently occur in the *Drosophila* nuclear genome. As a result, A and T have accumulated and C has been reduced at those sites under low functional constraints. This hypothesis was supported by data obtained from pseudogenes which indicated an accumulation of A and T in third positions (Moriyama and Gojobori, 1992). The second possible explanation is that mutations that cause accumulation of A or T in the coding regions of *Drosophila* nuclear genes may be subjected to some kind of selection. Consequently, genes under higher selective constraints may have a more biased codon usage (presumably towards G+C), as described by Sharp and Li (1989).
Using an estimate of the divergence time between *D.melanogaster* and *D.virilis* of 40 million years (Moriyama, 1987), the rate of synonymous substitutions was calculated to be $15.52 \times 10^9$/site x year for *nonA*, $15.43 \times 10^9$/site x year for *per*, $18.94 \times 10^9$/site x year for *tra*, $15.8 \times 10^9$/site x year for *tim* and $12.94 \times 10^9$/site x year for *en*. Moriyama and Gojobori (1992) had already calculated the rate of synonymous substitutions in *per* to be about $16 \times 10^9$/site x year after comparing the sequences of *D.virilis* and *D.melanogaster*. The small difference with my calculations for *per* are likely to be due to different methods used to align the two sequences and to calculate the synonymous substitutions. Moriyama and Gojobori (1992) divided the *Drosophila* genes they studied into three groups according to their average synonymous substitution rate. These three groups have an average synonymous substitution rate of $11 \times 10^9$ (group I), $17.5 \times 10^9$ (group II) and $27.1 \times 10^9$ (group III) respectively. All the genes included in my study, except maybe *en*, can be included in group II, and seem to be evolving at a rate determined by a similar molecular clock.

**Codon usage**

If synonymous mutations are selectively neutral, we should expect all the synonymous codons for one amino acid to be used with more or less equal frequencies. However since the first studies of Grantham *et. al.* (1980, 1981), it has become apparent that the usage of codons is not random. Moreover related species show similar patterns of choice between synonymous codons, i.e. codon biases seem to be species specific (Grantham *et. al.* 1980). In most species, however, there is also a considerable difference in codon usage between different genes (Sharp *et. al.*, 1988).

Biases in codon usage could constrain the rate of silent substitutions, and in fact it has been shown that the rate of synonymous substitutions is negatively correlated with the degree of codon usage bias (Sharp and Li, 1989). Sharp and Li (1989) have concluded that the variation in silent substitution rates among different nuclear genes in *Drosophila* is mainly due to differences in the strength of selective constraints on synonymous codons. In addition to this, Moriyama and Gojobori
(1992) found that the rates of synonymous substitutions are also correlated negatively with the C content and positively with the A content at the third codon positions (see above).

Here, the codon usage patterns were studied in ten homologous genes of *D.melanogaster* and *D.virilis: nonA, en, hb, knirps, nanos (nos), oskar (osk), per, runt, sevenless (sev), and tra*. The Codonfrequency program of the GCG packet was used to count codons and write their frequencies into codon frequency tables. Table 3.6 shows, for each amino acid in each *D.virilis* gene, which codon was mostly used. The biased codons for each gene of *D.melanogaster* are shown between brackets only when different from those used by *D.virilis*. For amino acids encoded by only two different codons, a gap means that no preference was shown in the use of one or the other.

The Relative Synonymous Codon Usage (RSCU) has been defined by Sharp *et al.* (1988) as the observed number of codons divided by the number expected if all codons for that amino acid were used equally. In table 3.6, codons of the *D.virilis* genes studied with a RSCU greater than 1.5 are marked by one asterisk, those with a RSCU greater than 2 are indicated with two asterisks.

It is immediately clear that the codon usage for *nonA* differ from that of other genes. Firstly, with the only exception of *tra*, *nonA* is the gene which displays the larger difference in codon usage between *D.virilis* and *D.melanogaster*. In *nonA*, in 10 amino acids out of 18, a different codon is used in *D.virilis* compared to *D.melanogaster*. In all the other genes always fewer than 8 amino acids show a different codon usage in the two species, the only exception being *tra* in which 11 amino acids present different codon bias between *D.virilis* and *D.melanogaster*.

Another striking feature of the *nonA* sequence, is the high A/T content found in the third codon position. Table 3.7 shows the percentage, within the number of biased codons, of codons that end with A or T in all the 10 *D.virilis* and *D.melanogaster* genes analyzed. Overall in both *D.virilis* and *D.melanogaster*, it appears that A and T ending codons form a substantial proportion of biased codons only in *nonA* and *tra*. If
Table 3.6. Codon bias table for 10 *D. virilis* (in black) and *D. melanogaster* (in red and brackets) genes. The biased codons of *D. melanogaster* are shown only when different from those of *D. virilis*. The number below each amino acid indicates the corresponding number of degenerate codons. For amino acids coded by two different codons, a gap indicates no bias for one or the other. For *D. virilis* only (in black) biased codons with a RSCU >1.5 are marked (*), codons with a RSCU >2 are also marked (**).

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Table 3.7. For each *D. virilis* and *D. melanogaster* gene, the number and the percentage of biased codons ending in A or T is indicated.

<table>
<thead>
<tr>
<th>Gene</th>
<th>BIASED CODONS</th>
<th>A/T-ending</th>
<th>% A/T-ending</th>
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<td>10</td>
<td>58.8</td>
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<td></td>
<td><em>mel</em></td>
<td>16</td>
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</tr>
<tr>
<td><em>en</em></td>
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</tr>
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<td><em>hb</em></td>
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<td>3</td>
<td>16.7</td>
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<tr>
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<td><em>mel</em></td>
<td>18</td>
<td>5.6</td>
</tr>
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<td><em>knirps</em></td>
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<td>8</td>
<td>44.4</td>
</tr>
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<td></td>
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<td>5.6</td>
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<td>16.7</td>
</tr>
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<td></td>
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<td>18.8</td>
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<td><em>mel</em></td>
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<td><em>mel</em></td>
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<td>16.7</td>
</tr>
<tr>
<td><em>sev</em></td>
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<td>7</td>
<td>41.2</td>
</tr>
<tr>
<td></td>
<td><em>mel</em></td>
<td>16</td>
<td>43.8</td>
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Table 3.8. Number of codons with RSCU >1 and >2 for each *D. virilis* gene.

<table>
<thead>
<tr>
<th>Gene</th>
<th>&gt;1.5</th>
<th>&gt;2</th>
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<td><em>nonA</em></td>
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</tr>
<tr>
<td><em>en</em></td>
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<td>6</td>
</tr>
<tr>
<td><em>hb</em></td>
<td>6</td>
<td>4</td>
</tr>
<tr>
<td><em>knirps</em></td>
<td>5</td>
<td>2</td>
</tr>
<tr>
<td><em>nos</em></td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td><em>osk</em></td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td><em>per</em></td>
<td>7</td>
<td>8</td>
</tr>
<tr>
<td><em>runt</em></td>
<td>3</td>
<td>8</td>
</tr>
<tr>
<td><em>sev</em></td>
<td>7</td>
<td>6</td>
</tr>
<tr>
<td><em>tra</em></td>
<td>7</td>
<td>6</td>
</tr>
</tbody>
</table>
we focus on *D. virilis* only, we can add *nos* to this list. This unusual result could be interpreted to mean a lack of selective constraint on *nonA*, leading to the expression of the A/T mutation bias, as suggested by Moriyama and Gojobori (1992; see above). This should be reflected in a higher synonymous mutation rate for *nonA*. However, as described above, *nonA* does not seem to have elevated mutation rate compared to the other genes examined (*per, tra, tim* and *en*). Moreover, *tra* has a slightly lower number of A/T-ending biased codons compared to *nonA*, yet, instead of having a lower (or at least similar) synonymous mutation rate compared to *nonA*, it has a higher one. These observations therefore do not fit very well with those presented by Moriyama and Gojobori (1992).

Finally, to further complicate the picture, the *D. virilis* *nonA* pattern of codon usage seems to be very different from that of the other *D. virilis* and *D. melanogaster* genes (see table 3.6). In *D. virilis* *nonA* the codons GTT for Valine, CGT for Arginine, and ACA for Threonine have a RSCU greater than 2, meaning a very strong bias. Interestingly, *D. melanogaster* always prefers other codons for Arginine and Threonine in *nonA*. Similarly, in all the other genes studied, either in *D. virilis* or in *D. melanogaster*, the codons used for Val, Arg, and Thr are almost always different from those used in *nonA* of *D. virilis*. There could be an explanation for this if Val, Arg, and Thr were highly repeated amino acids, since repetition of a single amino acid could largely inflate the proportion of the codon encoding for it. However the *D. virilis* *nonA* amino acid sequence data does not support this hypothesis, since V, R, and T are not involved in large repeats. Similarly, the results from the cryptic symplicity analysis do not indicate the presence of any short repeat containing the motifs: GTT, CGT, or ACA. In contrast to these observations, those amino acids which in *D. virilis* *nonA* are actually involved in repeats, such as Gln, Ala, His and Gly, show a codon bias similar to that all the other *D. virilis* and *D. melanogaster* genes examined.

Biases in codon usage can result from a combination of several factors, for example mutation trends, translational selection among synonymous codons, or selection for particular structures in DNA. In general, highly expressed genes show
very strong biases towards a particular subset of codons, while lowly expressed genes show a more even codon usage (Sharp et al., 1988). In E.coli, the favoured codons in highly expressed genes are those recognized by the most abundant tRNA species (Gouy and Gautier, 1982; Ikemura, 1985). In contrast, in lowly expressed genes, the deviation from equal usage of synonymous codons is most likely to be a result of mutation trends (Sharp and Li, 1986; Bulmer, 1988). To have an idea of the strength of the codon bias in the 10 D.virilis genes studied here, the number of codons with a RSCU >1.5 and >2 was counted (see table 3.8). From table 3.8 it is clear that nos and osk have the lowest bias strength of all. Interestingly, nonA, with the only exception of nos, has the lowest number of codons with a RSCU >2, showing therefore a rather poor bias strength. If we assume that strength in codon bias reflects the level of expression of a gene, then, from the data presented here, the results (table 3.8) predict that nonA is not as highly expressed as most of the other genes included in the study. Any differences in nonA's codon usage would therefore simply be the result of mutational forces, and not selection based on tRNA availability. If this is the case, then these forces should give different codon usage patterns also to nos and osk (which also have weak bias - see table 3.8), compared to the other genes. However this is clearly not the case (see table 3.7), and nonA's unusual bias for codons GTT (Val), GCT (Arg), ACA (Thr), TTG (Leu) and AAA (Lys) (the first four having a RSCU >1.5), remains something of an enigma.

In conclusion, we have seen that the D.virilis nonA gene has diverged considerably from that of D.melanogaster, particularly in the N-terminal half. The next step is to find out whether this divergence has any repercussions for the biological functions of the nonA gene. In other words, will the D.virilis nonA gene rescue the behavioural defects of the D.melanogaster nonA mutant? O'Neil and Belote (1992) showed, for example, that the D.virilis tra gene, in spite of being highly diverged from its D.melanogaster orthologue, was able to restore "femaleness" to diplo-X D.melanogaster tra pseudomales. If the D.virilis nonA gene can restore normal song and visual behaviour to nonA mutants, then it will be possible to assess whether the
transformants show any evidence for *D. virilis*-like song behaviour. In the case of the *per* gene, which involved transforming the *D. simulans* *per* gene into arrhythmic *per*\(^0\) *melanogaster* hosts, robust interspecific rescue was obtained for the arrhythmic song phenotype, and the *simulans* characteristics were transferred to the host (Wheeler *et al.*, 1991). However, *D. simulans* diverged from *D. melanogaster* about two million years ago (Cariou, 1987), whereas the *D. virilis/D. melanogaster* relationship is far more remote (Moriyama, 1987; Beverly and Wilson, 1984). The following chapter describes how the *D. virilis nonA* gene was cloned into a vector and transformed into *D. melanogaster* hosts and includes the genetic and molecular analysis of the various lines that were generated.
CHAPTER 4

Transformation of $nonA^{vir}$
INTRODUCTION

In chapter 3 I described how the *D. virilis* genomic clone containing the putative *nonA*^vir^ gene was subcloned in two fragments into the pUC18 vector. In order to transform *D. melanogaster* flies with the *nonA*^vir^ gene it was necessary to rejoin the two DNA fragments by cloning them into a suitable transformation vector. The *P* element vector pW8 (Klemenz et al., 1987) was preferred among others (Ashburner, 1989) for its large polylinker containing 12 unique restriction sites. This vector, derived from the *P* element end containing construct Carnegie4 (Rubin and Spradling, 1983), carries, as a marker, the *white*^+^ gene fused to the *heat shock protein 70* (*hsp70*) gene promoter. Expression of the *w*^+^ gene completely or partially rescues the lack of red pigment in the eyes of the recipient transformant *w* flies.

METHODS

Figure 4.1 schematically illustrates the procedure used to join the two *nonA*^vir^ DNA fragments into the transformation vector pW8. As a first step, the ~6.5 Kb *nonA*^vir^ DNA fragment in pUC39 was shortened of ~0.5 Kb by cutting with *XbaI* which recognizes a site in the pUC polylinker and one at about 0.5 Kb from the 3' end of the *nonA*^vir^ DNA insert, and *SphI*. The released ~6 kb *nonA*^vir^ band was gel purified and subcloned into the *SphI*-*XbaI* sites of pUC18 (see chapter 2). This new construct was named pUC23. The 5' and 3' ends of the insert in pUC23 were sequenced using the pUC Universal and Reverse primers, in order to make sure that the 0.5 Kb fragment had been actually removed from the correct end of the insert. The next step was to clone the *SphI*-*XbaI* DNA fragment from pUC23 into the unique *SphI* and *XbaI* sites of the pW8 polylinker. To do so, the 6 Kb *SphI*-*XbaI* fragment was released from pUC23 by digesting with *SphI* and *XbaI*, gel purified, and ligated to pW8 digested with the same enzymes. Transformation of the construct in *E. coli* cells was
Figure 4.1.
Cloning steps performed to join two \textit{nonA}^{vir} fragments into the transformation vector pW8. The size of the plasmids is not in scale.
performed by electroporation as described in chapter 2. This plasmid was called pW824.

In the second step the ~6 Kb nonAvir DNA fragment contained in pUC7 was excised by digesting with Sphi and cloned into the Sphi site of construct pW824. Two insert-containing positive clones were obtained (pW824-12; pW824-25), and a restriction digestion confirmed that in both constructs the Sphi-Sphi fragment had inserted in the appropriate orientation reconstituting the intact nonAvir gene. To further check whether no frame-shift mutations had occurred at the two inserts' junction site, the DNA from both pW824-12 and pW824-25 constructs was manually sequenced (as in chapter 2) using primers annealing upstream and downstream of the Sphi junction. Large scale DNA preparation was made of construct pW824-12 (the procedure was described in chapter 2) and this DNA was used to transform flies as reported in chapter 2. From now on this construct will be referred to as the nonAvir construct.

**Transforming w; e Sb Δ2-3/TM6 flies**

When flies from the w; e Δ2-3 Sb/TM6 stock were used as the injectees of the nonAvir construct the source of transposase was provided by a stable P-element inserted on the third chromosome. This element, constructed in vitro by Laski et al. (1986) is missing the third (2-3) intron of the P element and although it is very stable, it causes mobilization of other elements at unusually high frequency (Robertson et al., 1988). The injection procedures are described in chapter 2. The nonAvir DNA was precipitated and resuspended in injection buffer (5 mM KCl and 0.1 mM NaH2PO4, pH 6.8) at a concentration of 600 ng/μl.

**Transforming w1118 flies**

The helper plasmid pUCHSΠΔ2-3 (a gift from J-M. Dura) was used as a transposase source when the injected flies were w1118. The DNA concentration of
nonAv construct: helper plasmid was 300:150 ng/μl in injection buffer. G₀ flies were backcrossed to the w¹¹¹⁸ stock and their progeny (G₁) screened for transformants (marked by the red-eye colour).

**Southern blotting of transformed lines**

The DNA from 50 flies of each transformed line was extracted using the method described in chapter 2. The DNA was digested with the appropriate restriction enzyme, run overnight on a 0.8% agarose gel and blotted according to the methods in chapter 2. The probe to be used in the standard hybridization step was the 5'P element end of pW8. This 600 bp DNA fragment was released from pW8 after digesting with HindIII-Xbal, and, after gel purification, 20ng of DNA were used for the hybridization as described in chapter 2. A probe annealing to the 3' P element end of pW8 was used in some cases. This probe was generated by PCR amplification of a 600 bp fragment from a D.melanogaster 2.9 Kb P element. However only ~180 bp of this probe actually annealed to the pW8 3' P element end making the hybridization signal rather poor.

**Mapping the inserts**

The crosses performed in order to simultaneously map and balance the transgenes are presented in figure 4.2. Flies of the double-balanced stock w; T2:3 apX⁺/SM5CyTM3Sb were crossed to flies from each transformant line. From the F₁, red-eyed individuals carrying the transgene and both balancers (marked with Cy and Sb) were crossed to double-balanced white-eyed flies. If the insert maps on the second chromosome, white-eyed, curly wing (w; Cy) F₂ flies are produced among the progeny. The proportion of the F₂ carrying these informative markers should be about 1/6 of the total progeny. If the insert is on the third chromosome, among the F2 progeny should be flies carrying both w and Sb.
Figure 4.2.
Crosses for mapping transgenic inserts (\(\nabla\)).
Mobilization of X-linked inserts

Transgenes that map to the first chromosome can be mobilized to other chromosome locations if a source of transposase is added in their genome. In order to do this, transformant females were crossed to males of the w; Sb e Δ2-3/TM6 stock (figure 4.3). Red-eyed males of the progeny marked with Sb carry the transposase-producing Δ2-3 element on their third chromosome. Transposase activity promotes mobilization of the insert in the germ cells of these males as well as in their somatic tissues. As a result some of these males displayed a red and white mosaic-eyed phenotype indicating that in some cells the insert had been lost. All the w⁺; Sb e Δ2-3 males were singularly crossed to w1 1 1 8 females and the male progeny screened for the presence of w⁺ which would indicate a jump to an autosomal location.

RESULTS

Lines 112 and 113

Transformant lines 112 and 113 were obtained from ~180 injected G₀ w; Sb e Δ2-3/TM6 flies. The insert in 112 was found to be homozygous viable on the X chromosome. Several extensive attempts to jump this insert into an autosomal location, using crosses to w; Sb e Δ2-3/TM6 flies as described above, failed. The position of insert 112 on the X chromosome was roughly estimated by crossing 112 flies to a yellow, white (y w) strain and backcrossing the F₁ females to y w males. The number of recombinant flies (333 y w⁺, 400 y⁺ w) over the total number of F₂ flies (1572) gave an approximate map position of 1-47, very close to the estimated position of the endogenous nonA locus (1-52, Kulkami et al., 1988). An in situ hybridization to polytene chromosomes of line 112 was performed by Federica Sandrelli at the University of Padova, using a ~3 Kb nonAvir specific probe generated by digesting the pUC7 construct (see figure 4.1 and chapter 3) with HindIII. This probe included the
Figure 4.3.
Crosses for mobilizing an X-linked transgene (\(\nabla\)).
Only the first and third chromosomes are indicated.
first exon of nonA\textsuperscript{vir}, some upstream sequence, and part of the first intron. From the results of the \textit{in situ} hybridization, the 112 insert was mapped to 18F-19-A, again very close to the endogenous nonA gene which lies at 14C1-2 by \textit{in situ} hybridization using a cDNA probe (Besser \textit{et al.}, 1990). Given the proximity of nonA\textsuperscript{vir} to the endogenous nonA in line 112, attempting to isolate a double cross-over of the 112 insert to an X chromosome carrying a deletion of the nonA locus (Stanewsky \textit{et al.}, 1993) or the nonA\textsuperscript{dis} mutation seemed impractical. Consequently any behavioural phenotype produced by the insert in line 112 could not be tested on a nonA mutant background (see next chapters).

Southern blots of line 112 hybridized with a probe for the 5' P element end always revealed the presence of two bands (figure 4.7, 4.10, 4.11) indicating that either the 5' P element end had been somehow duplicated during integration of the transgene, or that the X chromosome contained two inserts. However the former seemed to be the most plausible explanation since only one band was found when using hybridization probes annealing to either the 3' P element end (figure 4.4) or the 6 Kb 3' half of the nonA\textsuperscript{vir} gene (figure 4.5).

The transgene of line 113 integrated on the Sb $\Delta2$-3 chromosome. To avoid further mobilization of the insert, the $\Delta2$-3 element was recombined off the chromosome by crossing red-eyed females to $w$ males. To test for the loss of $\Delta2$-3 the DNA of ten transformant males not marked with Sb was individually amplified using primers for the $\Delta2$-3 element, and run on a gel alongside a positive control (amplified DNA of $w$; Sb $\Delta2$-3/TM6 flies). None of the ten samples displayed a band corresponding to the amplification product of the positive control, indicating the absence of $\Delta2$-3 (data not shown). Line 113 was initiated from the progeny of only one of these tested males after crossing to $w$ females. The insert in line 113 is homozygous lethal, which means that after balancing the transgene-carrying third chromosome with MKRS (marked with Sb), the only viable $w^+$ flies carry Sb.
Figure 4.4.

A: Southern blot with EcoRI digested DNA from lines 112, 113, 67-4, and 168-8. EcoRI digested DNA from w^1118 flies (lane w) was included as a P element-free negative control sample. HindIII digested λ DNA was used as a marker. The probe used annealed only partially to the 3' P element end of pW8 (see methods) making the hybridization signal rather poor. B: schematic illustration showing the position and orientation of the nonAv" construct in relation to the 5' and 3' P element ends of pW8 (named "5' P" and "3' P" in picture). The white gene in pW8 is also shown. The arrows above each gene indicate the 5' to 3' orientation. The annealing position of the 3' P element probe is also indicated. The construct is ~15 Kb long.
Figure 4.5.

A: Southern blot of Sphl digested DNA from transformant line 112. The hybridization probe used was the ~6 Kb Sphl-Xbal fragment of nonAvir (picture B). DNA from w^1118 flies served as a negative control in the hybridization. The presence of only one band in the 112 lane suggests that a single transgenic insert is present in this line. B: Schematic illustration showing the position and orientation of the nonAvir construct in relation to the 5' and 3' P element end of pW8, as in figure 4.4. The annealing position of the probe is indicated by the vertical arrow.
Lines 168-8 and 67-4

After the failure in mobilizing the X-linked 112 insert, an attempt was made to mobilize the third chromosome insert of line 113. In this case, even local jumps on the same chromosome, which are more likely to occur (Tower et al., 1993), would have been welcomed. The fact that the insert in line 113 is homozygous lethal was used as a means of screening for new insertions. If the transgene moves to another genomic location it should complement the lethality of insert 113. The cross designed to mobilize insert 113 is schematically represented in figure 4.6. About 200 males carrying insert 113 and the Sb e Δ2-3 chromosome were individually crossed to females of line 113 (i.e. carrying insert 113 balanced by MKRS). The progeny were scored for any red-eyed flies which were not carrying the Sb marker. In theory these flies should carry two copies of the transgene: one at the original position of 113, and one in a novel genomic location. Only one out of the 200 individual crosses (number 168) produced such flies among its F2. One male with red eyes and no Sb marker was crossed to w; TM2/MKRS females. All the flies generated from this cross had red eyes, indicating that the putative new insert was on the third chromosome. Half of these flies therefore carried the original 113 insert, while the other half carried the hypothetical jump. From these flies, 20 males with the insert (either the old 113 or the jump) balanced by MKRS were crossed again to w; TM2/MKRS females in order to generate 20 different balanced lines. In order to identify which of the 20 lines carried the newly mobilized insert, a red-eyed male carrying MKRS was taken from each of the lines and crossed back to females of the original 113/MKRS stock. Those whose progeny gave red-eyed Sb+ flies must originate from a line containing the insert in a novel position. Only one of these lines (number 168-8) was kept while all the others were discarded. The "new" insert in line 168-8 is homozygous lethal since the progeny of a male and a female carrying 168-8/MKRS consist only of w+ Sb flies.

Line 67-4 was also generated from the cross described in figure 4.6. Some flies among the progeny generated by one male (numbered 67), carrying the 113 insert and Δ2-3, crossed to 113/MKRS females, displayed a strikingly stronger eye colour than the parental one. However all these flies also carried Sb indicating that their insert was not complementing the lethality of insert 113. One male (number 4) from the flies
Figure 4.6.

Crosses for mobilizing the transgene in line 113 ($\nabla_{113}$) to the 2nd or 3rd chromosomes. Only the 1st and the 3rd chromosomes are indicated in the P and F1.
bearing the new eye colour was crossed to \( w \), \( \text{TM2/MKRS} \) females. Resulting males and females which had the insert balanced by \( \text{MKRS} \) were crossed to each other to generate line 67-4. Again, all the flies in this line are always \( w^+ \ Sb \) indicating that insert 67-4 is homozygous lethal. My first hypothesis on this line was that it could have originated by a jump that had left a copy of the insert in its original position. The fact that the eye-colour of 67-4 flies was much darker than 113 individuals also supported the idea that in line 67-4 there could be two copies of the insert. However, when Southern blots were performed on lines 113, 168-8 and 67-4, the results suggested a much more complicated scenario (see below).

When a Southern blot with \( \text{EcoRI} \) digested DNA from lines 113, 168-8 and 67-4 was hybridized with a probe for the 3' \( P \) element end, the band detected was of the same size for each line, suggesting that the insert was in the same position in lines 168-8 and 67-4 as in line 113 (figure 4.4). Since only one band was detected in the DNA of line 67-4, it appeared that this line carried only one copy of the insert. Moreover, when a probe for the 5' \( P \) element end was used, no signal was detected from the DNA of any of these lines suggesting that the insert in each one of them had lost its 5' \( P \) element end (figure 4.7). Consequently, since the original 113 insert had somehow lost the 5' \( P \) element end, it was very unlikely that it could have jumped to generate lines 168-8 or 67-4. \textit{In situ} hybridizations, performed by Federica Sandrelli at the University of Padova, also confirmed that the inserts in line 168-8 and 67-4 were localized exactly in the same position as in line 113 (80 A on the third chromosome).

A Southern blot was then performed to check whether the \( \text{XbaI} \) site adjacent to the 5' \( P \) element end had also been lost. This site marked the proximal extremity of the \( \text{nonA}^{\text{vir}} \) gene as it was cloned into pW8 (see figure 4.1), and its loss would indicate that also some of the 3' end DNA of the original \( \text{nonA}^{\text{vir}} \) had been lost. The Southerns in figure 4.8 and 4.9 show that this was the case for line 113 and consequently for line 168-8 and 67-4. However a \( \text{HindIII} \) site 2Kb from the 3' end of \( \text{nonA}^{\text{vir}} \) was retained in lines 113, 168-8 and 67-4 (figure 4.8) suggesting that the \( \text{nonA}^{\text{vir}} \) sequence lost in these lines could have been anything between 0 and 2 Kb. This \( \text{HindIII} \) site falls within the fourth \( \text{nonA}^{\text{vir}} \) intron (see chapter 3 and figure 4.8). No additional experiments
A: Southern blot of lines 112, 113, 67-4, and 168-8, using a probe for the 5' P element end of pW8. The DNA from flies of each line was either double-digested with EcoRI and BamHI or with XbaI and SstI as indicated. DNA from a P element-free w1118 (w) line was also included as a negative control.

B: Schematic illustration showing the position and orientation of the nonAvr construct in relation to the 5' and 3' P element ends of pW8 (named "5' P" and "3' P" in picture). The white gene in pW8 is also shown. The arrows above each gene indicate the 5' to 3' orientation. Some of the sites of pW8 multiple cloning site ("MCS") are indicated. The vertical arrow shows the annealing position of the 5' P element probe.
Figure 4.8.

A: Southern blot with HindIII and XbaI digested DNA from lines 112, 113, 67-4, and 168-8. DNA from w1118 was included as a negative control. As shown in picture B, an HindIII-XbaI cut releases two fragments of 2 and 3 Kb from the end of the nonA"vir construct. The probe, generated by PCR amplification of a -950 bp fragment spanning from the end of exon three to the beginning of exon 5 of nonA"vir, anneals mostly to the 3 Kb HindIII-HindIII fragment and in part also to the 2 Kb XbaI-HindIII fragment. From the hybridization results in A it is clear that all the tested lines retained the 3 Kb HindIII-HindIII fragment. However line 113, 67-4, and 168-8 appear to have lost the nonA"vir terminal XbaI site since they do not display the expected 2Kb XbaI-HindIII band (as in line 112). Instead they show a much larger band (marked with "?" in A) resulting by a XbaI genomic cut distal to the insert (the signal of line 168-8 is not evident). The size of this band seems to be the same for 67-4 and 113 indicating that the extent of the 3' terminal nonA"vir deletion is the same in both inserts.
Figure 4.9.

A: Southern blot with HindIII-XbaI digested DNA from lines 112, 113, 67-4, and 168-8. As shown in picture B the hybridization probe used was the 2 Kb XbaI-HindIII terminal fragment of nonA\(^{vir}\). For some reason, the use of this probe generated very poor hybridization results as seen by the quality of picture A. The results of this Southern confirm the data shown in figure 4.8. Again it is shown that only line 112 displays the expected 2 Kb XbaI-HindIII band while lines 113, 67-4, and 168-8 do not since in these lines the insert has lost the nonA\(^{vir}\) terminal XbaI site. The signal of line 113 here is not clear while the one of line 168-8 (which was not clear in the Southern of figure 4.8) is quite evident. The size of the band in 67-4 and 168-8 appears to be similar indicating that a similar deletion pattern occurred in the inserts of these two lines.
were performed in order to check whether the fifth nonA\textsuperscript{vir} exon was intact in the inserts of lines 113, 168-8 and 67-4. Further investigations aimed to evaluate the extent of 113, 168-8 and 67-4 nonA\textsuperscript{vir} sequence loss were unsuccessful due to the problems generated by the use of a probe for the 3' end of nonA\textsuperscript{vir} (see Southern in figure 4.9).

**Lines 72, 135, and 297**

\(w^{118}\) flies were injected as described in the methods above. From \(-300\) G\(_0\) injected lines only three generated transformant progeny. Transformant lines 72 and 297 both had a homozygous viable X-linked insert. The only autosomal insert, in line 135, was mapped and balanced as described in the methods (figure 4.2): it is homozygous viable on the third chromosome. A Southern blot was performed on these lines using the 5' \(P\) element end of pW8 (figure 4.10). Again, it was found that one of the three lines (line 72) had lost the 5' \(P\) element end. The presence of a single hybridization signal from the blotted DNA of 135 and 297 indicated that in these lines only one copy of the transgene had inserted into the genome.

**Lines 297-6, 297-64, 297-71**

The X-linked insert in line 297 was successfully mobilized with the method illustrated above. Three new independent autosomal lines were generated: 297-6, 297-64 and 297-71. The insert in 297-64 was mapped to the third chromosome. A Southern blot hybridized with the 5' \(P\) element end probe showed that this end had been lost in the insert of lines 297-64 and 297-71 (figure 4.11). The insert in line 297-6 appeared to be in single copy.

**Lines 75, 97, 191**

Three new independent transformant lines were generated from \(-200\) \(w^{118}\) injected flies. They all appeared to have autosomal inserts. A Southern blot checking
Figure 4.10.

A: Southern blot of *EcoRI* digested DNA from lines 112, 72, 135, and 297, using a probe for the 5' P element end of pW8. DNA from *w^{1118}* flies was included as a negative control. **B**: Same as figure 4.7B.
Figure 4.11.

A: Southern blot with EcoRI digested DNA from lines 112, 75, 97, 191, 297-6, 297-64, 297-71, and w^{118} (w), hybridized with a probe for the 5’ P element end of pW8. The signal in the lane corresponding to line 297-64 is an artifact due to a tear in the filter. No band was actually detected in 297-64 lane. B: Same as figure 4.7B.
the 5' P element end confirmed that all three independent inserts had retained it (figure 4.11). Only one copy of the transgene was present in each tested line. By crossing to w; T2,3 apX u /SM5CyTM3Sb flies (see methods) the inserts were mapped to the third chromosome in lines 97 and 191 and to the second in line 75.

DISCUSSION

In this chapter I reported how a construct containing the entire predicted sequence of the viridis nonA gene was created and transformed into the genome of D.melanogaster with the use of P element-mediated germline transformation. The main problem with P element-mediated transformation remains the fact that no control can be imposed on where in the genome the transformed gene will insert. As a consequence, every independently derived transgene is localized in a unique chromosomal context that may affect its expression. When studying the phenotypic effects of a transgene, several independent insertions of the same transgene must be isolated in order to rule out these possible position effects.

The first two lines obtained by transforming w; Sb e Δ2-3/TM6 flies with the nonAvir construct were line 112 and 113. Line 112 carried the insert on the X chromosome. The position of the insert was more precisely mapped and turned out to be quite close to the endogenous nonA locus, thereby making it difficult to test the effects of the 112 insert on a nonA mutant background without an extensive and complicated series of crosses. It was therefore necessary to obtain at least one other autosomal line in addition to line 113. The easiest way to obtain autosomal lines given an X-linked transgene is to mobilize it and select for new insertions on the autosomes. The transposase necessary for mobilization of the insert was provided by the Δ2-3 element. Several extensive attempts to jump the 112 insert onto an autosomal location all failed. In order to transpose, a P element must have intact 5' and 3' ends. Approximately 150 bp of DNA at each end is required for transposition (O'Hare and Rubin, 1983). Contained within these sequences are a perfect terminal inverted repeat
(IR) of 31 bp and an 11 bp sequence found in inverted orientation ~130 bp from each end (O'Hare and Rubin, 1983; Engels, 1989). Characterization of the transposase protein has shown that this DNA-binding protein interacts with an internal 10 bp consensus sequence adjacent to the 31 terminal inverted repeat of each 5' and 3' end (Kaufman et al., 1989). Another type of protein named IRBP (inverted repeat binding protein) interacts with the outer 16bp of both terminal inverted repeats (Staveley et al., 1995) and it has been proposed to mediate transposition by increasing the affinity of the transposase for the internal 10 bp target sequence (Kaufman et al., 1989). The presence of both 5' and 3' P element ends was checked in the 112 insert by Southern Blotting. However, no actual information was obtained about the integrity of the sequences required for transposition at each end. Therefore the stability displayed by insert 112 could be due to post-insertion damage to its terminal P element ends. Alternatively the genomic position of the 112 transgene could be responsible for its inability to transpose. Cases have been reported in which the genomic position of a certain P element was the only cause of abnormally low levels of transposition (Engels, 1989). A P element marked with the wild-type allele of the white gene (P(w\textsuperscript{+})) inserted at cytological position 17B was found to have very low levels of transposition and excision. However, when a rare transposition to a new position was finally obtained, the new site displayed normal transposition levels (Engels, 1989).

Whenever the DNA from line 112 was digested with an enzyme that did not cut within the 5' P element sequence, and probed with the 5' P end HindIII-XbaI fragment from pW8, two bands were always displayed in the autoradiographs (figure 4.7, 4.10 and 4.11). This would suggest either the presence of two copies of the 5' P element end in the 112 insert or the presence of two copies of the non\textsuperscript{A\textsuperscript{vir}} construct. Since only one band was found when probing for the 3' P element end (figure 4.4) or for the second half of the non\textsuperscript{A\textsuperscript{vir}} construct (figure 4.5), it seems more likely that there is a single insert in line 112. However its 5' P element end must have duplicated, possibly via some sort of slippage during the DNA replication required for the transgene's chromosomal insertion (Klecker, 1981; Shapiro, 1983).
An attempt was subsequently made to mobilize the autosomal insert 113. The advantage of jumping this third chromosome transgene was that any new local insertions within the same chromosome should be easier to obtain compared to interchromosomal hops (Tower et al. 1992). The 113 insert is homozygous lethal. P elements can be mutagenic for a variety of reasons, depending on the position of the insertion in relation to the affected gene. P elements seem to preferentially insert at or near genes' transcripton start sites (Engels, 1989) and consequently affect promoter function, usually reducing the level of wild type transcript. Alternatively, insertion into the S' untranslated regions of genes could affect the stability of the transcript or result in its premature termination. P elements can also insert in the coding region of some genes, disrupting the production of a functional protein. One such example is the white locus in which four insertions in the coding sequence were recovered, three of which mapped to the same nucleotide (Rubin et al., 1982; O'Hare and Rubin, 1983). Then there are more complicated cases of P element-gene interactions like the one described by Horowitz and Berg (1995). They showed that an in vitro generated P[lacZ, rosy+] element, caused a mutant phenotype by inducing aberrant splicing and termination of the pipsqueak gene transcript. It was found that the P element, which was inserted within an intron of pipsqueak, also carried the splice acceptor site and the transcription termination sequences of a gene called l(3)S12. Consequently an aberrant splicing event was joining the splice donor site upstream of the pipsqueak intron, in which the P element was inserted, to the acceptor site of the l(3)S12 intron. The result was a fusion pipsqueak-l(3)S12 transcript which yielded a protein with mutant effects.

Whatever causes the mutagenic effect of a P element insertion, it is possible to have reversion to wild-type simply by mobilizing the insert. However if the insert is located in the coding region of a gene, only a precise excision of the element can restore the proper reading frame. Bearing this in mind, the insert in line 113 was mobilized by providing a stable transposase source (the Δ2-3 element, see methods). It was believed that a new insertion could be detected because of its ability to complement the lethality of the original 113 insert. The third chromosome insert 168-8 displayed such a property in that flies heterozygous for 113 and 168-8 were fully viable. However when Southern analysis was performed, it was found that the insert in
168-8 was still at the same genomic position of 113. Further analysis revealed that both the original insert 113 and the new 168-8 had apparently lost their 5' P element end (figure 4.7). It was therefore unlikely that insert 113 could have transposed to a new genomic location in order to give the new insert 168-8.

However, even though insert 113 could not have transposed somewhere else, it is still possible that it might have been excised from the genomic DNA. The transposase could have mediated the excision after recognising the 10 bp target sequence internal to the terminal repeat of each P element end (Kaufman et al., 1989). In fact, even though the Southern performed indicated that most of the sequence of the 5' P element end was gone from the 113 transgene, it cannot be excluded that sequences important for transposase binding, or for other factors interacting with the transposase (such as the IRBP, Staveley et al., 1995; Kaufman et al., 1989) had been retained in 113. Alternatively the transposase could have recognized the 8bp repeated sequences found at each side of P elements after genomic insertion (Engels, 1989; Paques and Wegnez, 1993). The excision of 113 would have left a double stranded gap which could have been repaired using the sister chromatid as a template (Engels et al., 1990; Daniels and Chovnick, 1993; Staveley et al., 1995).

In order to explain the ability of 168-8 to complement 113 lethal phenotype, it must be assumed that during the process of repair an error might have occurred that changed either sequence within the nonA\textsuperscript{vir} construct or some flanking genomic DNA sequences. It is common that after excision of a P element the gap produced is expanded by exonuclease action. Thus an error could possibly occur even during the repair of the sequences that actually lay outside each end of the insert (Engel et al., 1990; Johnson-Schlitz and Engels, 1993). There is evidence that if the sequence within the P element end contains a fair amount of repetitive material, the repair process is especially error-prone due to frequent slippage-like events occurring during DNA synthesis (Paques and Wegnez, 1993; Paques et al., 1996; Golic, 1994; Staveley et al., 1995). How a sequence change occurring during gap repair, either within the nonA\textsuperscript{vir} construct or in some sequences outside it, could have reversed the effect of the lethal mutation caused by the 113 insert, remains a mystery. Moreover the "new" insert 168-
8 is also lethal when homozygous, suggesting a very complex mechanism underlying these events.

Line 67-4 was also isolated during the same screen that produced line 168-8. The insert in this case does not display complementation of the 113-caused lethality. However flies containing this insert have eyes which are of a much brighter and stronger shade of red than those of flies from line 113. Different eye colour displayed by transformant flies usually indicate different levels of expression of the w+ transgene suggesting different localization in the genome of recipient flies. The first hypothesis was that this line could have originated by a jump of 113 that had left a copy of the insert in the original position. This would have accounted for the lack of lethal complementation and the darker eye colour (since two copies of the white+ gene would have been present). However Southern blot analysis confirmed again that insert 113 had not moved in order to originate line 67-4 (figure 4.4). If we assume again that the 113 insert had undergone transposase-dependent excision, then 67-4 can be explained as the result of an event occurring during gap repair. Whatever happened to insert 67-4 must have affected the expression of the white+ gene encoded by the transgene. A possible duplication of the white+ gene cannot be excluded since it has not been tested.

As well as the 5' P element end, insert 113 and consequently inserts 168-8 and 67-4 have lost the teminal XbaI restriction site which marked the 3' end of the nonAvir construct (see methods and figure 4.1, 4.8 and 4.9). All the three lines may therefore be missing some other 3' terminal sequences of the original nonAvir construct. However little can be said about the exact extent of the deletion. A HindIII site found 2kb upstream of the 3' end of the nonAvir construct was however retained in all the three lines (figure 4.8) indicating that the sequence loss from the end of the original construct after insertion must be between 0 to a maximum of 2kb. Comparing the Southern results of line 113 with those of lines 168-8 and 67-4 (figure 4.8 and 4.9), it seems that the extent of the 3' terminal nonAvir deletion is the same in the three lines. Because of the way the Southern were designed (see legends in figure 4.8 and 4.9) an additional loss of 3' terminal nonAvir sequence in the inserts of line 168-8 and 67-4 would have resulted in a smaller band displayed after hybridization of their DNA.
Instead, the bands resulting from lines 168-8 and 67-4 seem to be of the same size as that of line 113. However it must be noted that small differences in DNA sizes cannot be revealed by Southern blot.

Transformant lines 72, 135, 297, 75, 97 and 191 were all generated by injecting the nonA\textsuperscript{vir} construct into embryos of the w\textsuperscript{1118} strain. This method proved to be more efficient than injecting w; e Δ2-3 Sb/TM6 flies mainly because of the much greater survival rate of the w\textsuperscript{1118} injected embryos. Lines 267-6, 267-64, and 267-71 were easily obtained by mobilizing the X-linked insert 297 to the autosomes. Southern blot analysis suggests that in lines 135, 297, 75, 97, 191 and 297-6 only one copy of the transgene has inserted into the genome (figure 4.10 and 4.11). When using a probe for the 5' P element end no hybridization signal was displayed by the DNA corresponding to lines 72, 297-64 and 297-71. Thus again the 5' P element end had been lost, possibly upon integration of the transgene into the genome of the recipient flies. Considering the fact that this end has been lost in a total of four independent integrations plus in another case it has been duplicated, it is seems that replication slippage events at this end of the construct occur quite frequently. This suggests that the sequence at the 3' terminal end of the nonA\textsuperscript{vir} construct might be highly repetitive.

In spite of the enigmatic nature of lines 168-8 and 67-4 and the fact that they are not independent lines from 113, they were still used in all the behavioural tests described in the following chapters. No direct proof was obtained that these lines retained an intact fifth exon in their nonA\textsuperscript{vir} transgenes. However indirect proof of the existence of this exon in the three lines can be obtained by the behavioural results since the fifth exon has been shown to be indispensable for all the nonA biological functions (Jones and Rubin, 1990; Rendhal et al., 1992). Among the other transformant lines, only lines 135, 97, 191 and 297-6 were used in the behavioural analyses.
CHAPTER 5

Viability tests
INTRODUCTION

This chapter reports the first tests performed on transformant flies carrying a nonA\textsuperscript{vir} gene. Stanewsky et al. (1993) have isolated a lethal reciprocal translocation between the first and the fourth chromosomes (T(1;4)9e2-10) which produces a small deletion uncovering the nonA gene and a partially overlapping, upstream locus (l(1)i19e) whose product is essential in D.melanogaster. By transforming flies carrying this deletion with transgenes encoding only for the essential gene l(1)i19e, Stanewsky et al.(1993) managed to create nonA\textsuperscript{(c)} flies (see figure 5.1). Hemizygous nonA\textsuperscript{(c)} males are subviable: only 10-30% of the expected number eclose from metamorphosis. The few nonA\textsuperscript{(c)} males which reach the imaginal state are deficient in locomotor activity and flight, and less than one-third of them survive to become fertile males (Stanewsky et al., 1993). Not surprisingly nonA\textsuperscript{(c)} flies are severely affected in all the nonA-related behaviours. Severe mutations affecting the first RRM domain (RRM1) of nonA have also proven to considerably reduce viability along with affecting both visual and song behaviours (Rendahl et al., 1996; Stanewsky et al., 1996). As a corollary, we can expect that mutations that affect viability will undoubtedly disrupt all the behavioural phenotypes controlled by nonA.

Since it was decided that the nonA\textsuperscript{vir} transgene should be tested on a nonA\textsuperscript{(c)} background, it seemed sensible to first assess the efficiency of the nonA\textsuperscript{vir} gene in rescuing the viability deficits caused by nonA\textsuperscript{(c)}. If the nonA\textsuperscript{vir} transgene does not ameliorate viability of nonA\textsuperscript{(c)} males, there is little possibility that it will show any rescue of their behavioural defects. Bearing in mind that the nonA\textsuperscript{vir} construct used for transformation contains more than 3 Kb of DNA upstream of the first nonA\textsuperscript{vir} codon (see chapter 3, figure 3.8), I suspected that all the sequence of the virilis homologue of the l(1)i19e gene would also be contained in the transforming fragment. Therefore I decided to place the nonA\textsuperscript{vir} construct not in a nonA\textsuperscript{(c)} background, but directly on a T(1;4)9e2-10 genetic background. I knew that if any transformant males carrying the nonA\textsuperscript{vir} construct and the l(1)i19e +nonA double deletion could be produced, it would
Figure 5.1. A: molecular and genetic map of the X chromosomal region 14B17-14C4. The physical position of the 9-21, l(1)19e, and nonA genes is indicated. The black bar on top of these genes shows the extent of the 9e2-10 deletion. The nonA primary transcript is also shown: black bars represent exons, thin lines stand for introns. Underneath, the two genomic DNA fragments used to generate nonA° flies are indicated: the 211s12 fragment encodes only the first two exons of nonA, while the 235R11-SacII has a frame shift mutation (indicated by the arrow) introduced at the SacII site in the first intron. Thus neither of these two DNA fragments can produce a full NONA protein.

B: chromosomal constitution of a nonA° male. Grey bars: X and Y chromosomes; black bars: second or third chromosome; red bars: fourth chromosome. The X chromosome carries the 9e2-10 deletion uncovering both l(1)19e and nonA. The reciprocal 1;4 translocation is schematically represented by the grey and red bars. On the autosomes (black bars) a transgene (V) encodes only the l(1)19e gene product, this could be either the 235R11-SacII or the 211s12 DNA fragment.
have meant that the non$A^{\text{vir}}$ transgene carries the virilis homologue of the $l(1)i19e$ gene and that this gene is functioning in a $D.melanogaster$ background.

However, placing the non$A^{\text{vir}}$ construct over the 9e2-10 deletion background does not distinguish between poor rescue of the $l(1)i19e$-induced lethality and poor rescue of the semilethality associated with non$A^{(c)}$. In fact, a reduction in the expected number of $T(1;4)9e2-10/Y$ hemizygous males carrying the non$A^{\text{vir}}$ construct can be attributed either to a non complete rescue of the $l(1)i19e$ deletion, or to an incomplete rescue of the semilethality of non$A^{(c)}$, or both. In this case the non$A^{\text{vir}}$ construct should be tested independently on a non$A^{(c)}$ background and on a $l(1)i19e$ lethal mutant background (i.e. the $l(1)i19e^2$ mutant allele, Rendahl et al., 1996; Stanewsky et al., 1966).

**METHODS**

**Placing the non$A^{\text{vir}}$ construct on a $T(1;4)9e2-10$ background.**

Males from each of the seven autosomal transformant lines described in chapter 4 ($113, 67-4, 168-8, 135, 97, 191$ and $297-6$) were crossed to females carrying the $T(1;4)9e2-10$ double deletion over the FM7 balancer (figure 5.2). Males with the $T(1;4)9e2-10$ deletion-carrying X chromosome and a non$A^{\text{vir}}$ autosomal insert should be distinguished by the presence of the morphological markers carried by the $T(1;4)9e2-10$ X chromosome ($\text{yellow}'$ (y'), $\text{crossveinless}$ (cv), $\text{vermillion}$ (v), $\text{forked}$ (f), and $\text{carnation}$ (car) (Stanewsky et al., 1993).

**Viability tests.**

Large numbers of males from each transformant line were crossed to $T(1;4)9e2-10/FM7$ females and all the males of the progeny were classified and
**Figure 5.2.** Cross for testing the viability of flies carrying the *T(l;4)9e2-10* translocation (marked with $y^c v^f c a r$) and the *nonA<sup>vir</sup>* transgene (represented by $V$, marked with $w^+$). The three types of males generated by the cross are indicated. Only the X and one of the autosomes (either the second or the third) are shown for males B and C. In type A males the fourth chromosome involved in the *T(l;4)9e2-10* translocation is also illustrated. Grey bar = X and Y chromosome. Red bar = fourth chromosome. Black bar = second or third.
counted. The ratio of \( y, cv, v, f, car; w^+ \) males \((T(1;4)9e2-10 + \text{nonA}^{vir}\) insert\) over the total number of males produced by the cross was calculated. The genotypes of the other two kinds of males obtained (figure 5.2) were \( y, w, B; w^+ \) and \( y, w, B; \) since the \( FM7 \) chromosome is marked with \( y, w, Bar (B), \) (Lindsley and Zimm, 1992), and the transgene is marked with \( w^+ \) (Klementz et al., 1987).

RESULTS

If males carrying the X-linked \( T(1;4)9e2-10 \) double deletion and a \( \text{nonA}^{vir} \) autosomal insert were 100\% viable, their number should equal 1/3 of the total number of males produced by the cross (i.e. 33.3 \%). From table 5.1, it is clear that males carrying the \( l(1)i19e + \text{nonA} \) deletion \((9e2-10) \) and a \( \text{nonA}^{vir} \) transgene are viable, confirming that the \( \text{nonA}^{vir} \) construct used in the transformation (chapter 3 and chapter 4) must encode the complete product of the \( virilis \) homologue of the \( l(1)i19e \) essential gene.

Most transformant lines show full rescue of the viability defects associated with \( l(1)i19e \) and \( \text{nonA}^{(c)} \). Unexpectedly, for most lines, far more than a third of the total males, produced by the cross, carried the \( \text{nonA}^{vir} \) transgene on a \( 9e2-10 \) deletion background. This effect is so marked that a statistical test is unnecessary. It is inevitable that the presence of the \( FM7 \) chromosome in the two other types of males produced by the cross (B and C in table 5.1.), given the chromosomal rearrangements within the balancer (Lindsley and Zimm, 1992), will affect their viability.

Strangely, only 1 male of type C (see table 5.1) was produced in the cross involving the 135 transgenic line. The reason for this bizarre result is unknown. Line 191 generated significantly less than 33\% \( T(1;4)9e2-10/Y; \text{V}/+ \) males \((\chi^2 = 14.7; df = 2; p < 0.001)\). A position effect is the most likely explanation for this result.
Table 5.1. Number of males of each genotyope (A, B, or C) resulting from a cross
T(1;4)9e2-10/FM7 females x transgenic nonA\textsuperscript{vir} males (see figure 5.2). A: males carrying the
X-linked deletion T(1;4)9e2-10 and the nonA\textsuperscript{vir} insert on an autosome (indicated with the \( \forall \) symbol). B: males with FM7 and an autosomal nonA\textsuperscript{vir} insert. C: males with FM7 only. In
the last column the percentage of the A type of males is calculated over the total amount of
males collected from the cross.

<table>
<thead>
<tr>
<th>TRANSGENIC LINE</th>
<th>A</th>
<th>B</th>
<th>C</th>
<th>% A/(A+B+C)</th>
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<td>64</td>
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<td>36.5</td>
</tr>
</tbody>
</table>

DISCUSSION

In this chapter I have demonstrated that the nonA\textsuperscript{vir} construct (see chapter 3 and 4) fully rescues the lethality associated with a translocation T(1;4)9e2-10 that
causes a simultaneous deletion of the essential l(l)H9e gene and the nonA gene
(Stanewsky et al., 1993). This suggests that the nonA\textsuperscript{vir} transgene must contain the
entire coding sequence for the D.virilis homologue of the D.melanogaster l(l)H9e
gene, and that this gene product can function properly in a D.melanogaster
background. Furthermore, the nonA\textsuperscript{vir} construct fully rescues the subviability
associated with the absence of a functional nonA gene product in nonA\textsuperscript{(+)} flies.
(Stanewsky et al., 1993). Therefore there is good indication that the nonA\textsuperscript{vir} transgene contains sequences encoding for the full product of the \textit{D.virilis} nonA gene, and that the \textit{virilis} NONA protein is working in a \textit{D.melanogaster} background at least at levels sufficient to perform basic functions.

A useful consequence of these results is that, in order to perform the subsequent visual and courtship behavioural tests, there will be no need to produce nonA\textsuperscript{-} flies (as illustrated in figure 5.1) to serve as a genetic background for testing the various nonA\textsuperscript{vir} inserts. These can in fact be studied directly on a \textit{T(1;4)9e2-10} genetic background, saving a considerable amount of time consuming crosses that involve very unhealthy flies (i.e nonA\textsuperscript{-} flies).
CHAPTER 6

Analysis of visual behaviour
The nonA locus is one of the many genes which affect the normal functioning of the visual system of D.melanogaster (reviewed by O’Tousa, 1990; Pak, 1991; Smith et al., 1991). nonA mutations were first isolated in screens for visual mutants which exhibited reduced phototaxis (Pak, 1970; Hotta and Benzer, 1970). Subsequently it was observed that the nonA visual phenotype involves multiple defects, as assayed electrophysiologically and behaviourally. nonA flies are deficient in optomotor responses (Heisenberg, 1971a) and exhibit an aberrant electroretinogram (ERG) (Hotta and Benzer, 1969; 1970; Heisenberg, 1971b; Pak et al., 1970).

The Drosophila visual apparatus, like that of Musca, consists of a peripheral retina and three postretinal structures, the lamina, the medulla and the lobula, which together form the optic lobe (figure 6.1). The retina is composed of approximately 700 structural subunits called ommatidia, each optically isolated from the others by a layer of screening pigment cells containing pterins and ommachromes. These pigments give rise to the red colouration of the eye, and have a function similar to that of the pupil in the vertebrate eye. As well as the imagine-forming dioptic apparatus, each ommatidium bears a photoreceptive system, the retinula, which is composed of eight retinula cells (photoreceptors). Each retinula cell carries a rhabdomere, a differentiated membrane structure which contains the visual pigment. From each ommatidium, the rhabdomeres of cells R1-6 transfer their excitation to the monopolar neurons of the lamina, while those of cells R7-8 synapse directly with the medulla.

Recording the electroretinogram (ERG) of a fly can give an idea about the physiological and structural integrity of the peripheral retina and lamina in the compound eye. However interpretation of the results can be difficult, (Pak, 1975). An ERG records the electrical potential change elicited by a light stimulus, between an electrode placed in the cornea and another electrode (the reference electrode) placed in some unresponsive portion of the animal (usually the abdomen or the dorsal surface of the head.) The electrical potential recorded arises from the summed electrical activities
Figure 6.1. Schematic drawing of the dipteran eye and optic ganglia. Cr: crystalline cone; La: Lamina; Le: Lenslet; Lo: Lobula; Me: Medulla; O: Ommatidium; Rc: Receptor cell. Insert figure: crossection of retinula cells in one ommatidium. From Heisenberg and Buchner (1977).
of the retinula cells as well as the neurons (and possibly the glial cells) of the optic lobe. A typical ERG displays a corneal-negative component which is maintained as long as the light is on. In addition, the ERG includes a rapid, corneal-positive transient (the "on-transient") when the light is first turned on, and a corneal-negative transient (the "off-transient") when the light is turned off. The corneal-negative, maintained component is believed to originate mainly in the retinula cells (Goldsmith, 1965), although minor contributions may come from cells in the optic lobe. A variety of evidence indicates that neither the on nor the off-transient originates from the retinula cells (Autrum and Hoffman, 1957, 1960; Burkhardt and Autrum, 1960; Wolbarsht et al., 1966; Eichenbaum and Goldsmith, 1968; Alawi and Pak, 1971). Major components of these transients seem to originate in the lamina (Pak, 1975; Coombe, 1986; Coombe and Heisenberg, 1986).

The characteristic feature of an ERG recorded from nonA mutants is the lack of both on and off-transient spikes (Pak et al., 1970; Hotta and Benzer, 1970; Heisenberg, 1971b). Thus the ERG of these mutants consists mainly of the corneal-negative, maintained component. An obvious interpretation of these ERGs is that in the mutants the receptor cells must be functioning normally, but either the transmission of the light signal to the neurons in the lamina is prevented, or the excitation of these laminar cells is somehow blocked (Pak et al., 1969; Heisenberg, 1971b). Also, the possibility that the mutants' visual defects are not confined to the lamina, but may extend to the medulla, is not ruled out by the electrophysiological evidence.

The hypothesis that nonA mutants have mainly a central defect in their visual system is supported by their poor optomotor response, which is the fly's tendency to follow movements of the visual environment (Götz, 1964). For an optomotor response to occur, the movement of the environment must be detected comparing the outputs from at least two photoreceptors. Since synaptic interconnections between different photoreceptor axons take place in the lamina, the first motion detectors are probably localized in this structure. Heisenberg (1972) characterized the flying and walking optomotor behaviour of the mutant nonA^{H2} which was further refined by Heisenberg and Buchner (1977). These investigators showed that, in optomotor tests,
nonA\textsuperscript{H2} seems to be defective in vision at high light levels in a system that specializes in the resolution of visual details (high acuity system). However nonA\textsuperscript{H2} defects did not seem to manifest themselves at the level of the retina, and were assumed to occur at a level proximal to it. Studies on the optomotor-blind\textsuperscript{H31} (omb\textsuperscript{H31}) mutant, in which the giant neurones of the lobula plate are missing or severely reduced, suggest that a set of three neurons (H-cells) at the anterior surface of the lobula plate may be part of the main pathway for the optomotor turning response (Heisenberg et al., 1978). Electrophysiological experiments on larger flies have shown that the H-cells are sensitive to horizontal movement around the fly (Dvorak et al., 1975: Hausen, 1976). Thus they are good candidates for mediating optomotor responses.

In this study the optomotor response of transformant flies carrying one, two or three copies of the \textit{D.melanogaster} (nonA\textsuperscript{+}) and \textit{D.virilis} non-A gene (nonA\textsuperscript{vir}) were analyzed and compared to that of mutants nonA\textsuperscript{dis} and wild-type flies. An initial attempt was made to record ERGs from these flies, using equipment designed to measure ERGs in crustaceans. No reliable ERG recordings could be produced due to technical reasons. Considering the fact that ERGs artefacts are common even using better equipment, a decision was taken to use only the optomotor test in studying the visual behaviour of the flies of interest. The optomotor test, giving more than a single positive or negative result (in contrast to ERG measurements in which the only results are either presence or absence of transient spikes), was viewed as a more powerful test for uncovering subtle differences in the visual behaviour of the transformants.

**MATERIALS & METHODS**

The walking optomotor test was performed as described in chapter 2. Two sets of stripe widths were used: in one set each black and white pair subtended a small angle (26.6°). In the “large” stripes each black and white pair subtended a larger angle of 72°. 10 males for each genotype were tested in both patterns, with every male
subjected to twenty trials, in which the direction of the rotating stripes was alternatively changed.

GENOTYPES

A: Controls

Males from the *D.melanogaster* strain *Canton S* were used as a wild type positive control. The *nonA<sup>disst</sup>* mutant male strain *disst/FM7a; ry<sup>506</sup>/MKRS* provided a negative control in the test. Unfortunately it proved almost impossible to generate *nonA<sup>c</sup>* flies to use as negative controls. As described in chapter 5, *nonA<sup>c</sup>* males can be produced by crossing females carrying the *T(1;4)9e2-10* translocation, which deletes both the *l(1)i19e* and the *nonA* genes, with transformant males carrying autosomal *235R11-SacII* or *211S12* inserts (which encode only the functional product of the *l(1)i19e* gene, see figure 5.1). For some reason, even after performing such a cross several times with large numbers of flies, only two *nonA<sup>c</sup>* males were ever produced. I realized therefore that it would have taken too much time to produce enough *nonA<sup>c</sup>* flies to serve as negative controls in any of behavioural tests, and decided to use *nonA<sup>disst</sup>* mutants instead.

Hemizygous *nonA<sup>c</sup>[nonA<sup>+</sup>]*

As the experimental flies to be tested were transformants carrying the *D.virilis nonA* gene (*nonA<sup>vir</sup>*), appropriate transformant controls carrying the *nonA<sup>+</sup>* gene from *D.melanogaster* were required. These were provided as follows. The translocation *T(1;4)9e2-10* created a simultaneous deletion of the essential gene *l(1)i19e* and the *nonA* locus (Stanewsky *et al.*, 1993, see chapter 5). Transformant flies carrying a 11kb DNA fragment encoding both the lethal gene *l(1)i19e* and the *nonA* gene of *D.melanogaster* (*235R11*, Jones and Rubin, 1990) were provided by the stock *P[(ry)235R11]D1* (see chapter 2) which carries the homozygous insert on the third
chromosome. Males from this stock were crossed to T(1;4)9e2-10 IFM7 females. Since the X chromosome in T(1;4)9e2-10 is marked with y, cv, v, f, and car, all the F1 red-eyed males carrying these alleles rescue simultaneously the loss of the lethal gene and nonA. These flies carried one copy of the nonA+ gene of D.melanogaster over a nonA(−) background and were named 235R11/d.

B: hemizygous nonA(−)[nonAvir]

Transformants carrying one copy of the nonA gene from D.virilis (nonAvir) over a nonA(−) background were generated by crossing T(1;4)9e2-10/IFM7 females to males for each of the seven transformed lines carrying autosomal nonAvir inserts discussed in the previous chapter. The seven lines were provided by the transgenic strains 113, 67-4, 168-8, 135, 97, 191, 297-6. From each cross, only F1 males carrying the T(1;4)9e2-10 chromosome morphological markers were selected for behavioural testing. These males carry one copy of the nonAvir gene over a nonA(−) background. The fact that such viable males are produced by these crosses indicates that the nonAvir construct inserted in each line must also carry the intact virilis homologue of the lethal gene l(l)19e (see chapter 5). The males tested were respectively named: 113/d, 67-4/d, 168-8/d, 135/d, 97/d, 191/d; 297-6/d, where d is used as a symbol for the double deletion.

C: heterozygous nonAvir/nonA+

Transformants carrying one copy of the nonAvir gene plus the endogenous copy of melanogaster nonA+ were generated as follows. The transformant males from line 112 (discussed in the previous chapter) were included in this analysis: these flies carry one copy of nonAvir inserted not far from their endogenous copy of nonA+ on the X chromosome (carrying w, see chapter 4). These transformants were named 112/m. In the transformant lines 113, 67-4, and 168-8 the insert on the third chromosome causes lethality if homozygous. As each strain carries the w mutation on the X chromosome,
every red eyed male in these lines has one copy of the nonAvir gene plus the endogenous nonA+. The males analyzed from line 67-4 and 168-8 were called 67-4/m, and 168-8/m respectively. Transformant line 113 flies are distinguished by a very weak red eye colour. Bearing in mind that white flies are optomotor blind due to the lack of pigments in their eyes (see Hall, 1994), it was necessary to test the effect of line 113 on a wild-type (red-eyed) background. In order to do this, males from line 113/MKRS (i.e. carrying the 113 transgene over the MKRS balancer) were crossed to females Canton S. F1 males carrying no balancer were then tested in the optomotor apparatus and are here referred to as 113/m. All the other nonAvir inserts gave a strong red-eye colour when placed with a w X chromosome, so further crosses were not considered necessary. In lines 135 and 97 flies homozygous for the insert are fully viable. Males from lines 135 and 97 were therefore crossed with females from a w1118 stock. Red eyed male progeny carry a single copy of the insert. These males are called 135/m and 97/m respectively.

D: 3 mel

Males from the P[(ry)235R11]D1 stock are homozygous for the melanogaster nonA+ transgene on the third chromosome and carry the endogenous copy on their X chromosome. These flies were named 3mel.

E: 2 mel

Males from the P[(ry)235R11]D1 stock were crossed to females of a Canton S strain. The resulting males (called 2mel) have one transformed copy of the melanogaster nonA+ transgene on the third chromosome plus the endogenous copy on the X.
**F: mel+2vir**

Females from line 112 (*nonA^vir* insert on the X) which had their X chromosome balanced with *FM7*, were crossed to males from line 113 with their third chromosome balanced by *MKRS*. Resulting F1 males carrying no balancers must have one copy of *nonA^vir* on the third chromosome and one on the first, plus the endogenous *melanogaster nonA^+* gene on the X. These flies were named *mel+2vir*.

**GENETIC BACKGROUND**

When comparing the behaviour of independent transformants lines it is best to try to reduce possible differences due to the different genetic backgrounds. For this reason I tried to keep the background of the transformed flies analyzed here as similar as possible.

Lines 113 and 112 were obtained by transforming flies from the stock *w;Δ2-3/TM6* and successively crossing the transformants to flies from the laboratory stock *w Tully*. Lines 67-4 and 168-8 share the same background being derivatives of line 113. Heterozygous *nonA^vir/mel nonA^+* lines 112/m, 67-4/m, and 168-8/m also have the same background, so does *mel+2vir*. 113/m is slightly different since males of line 113 were crossed to *Canton S* females.

Lines 135, 97, 191 and 297-6 were all obtained by injecting another strain of *white* flies (stock *w^118*) and crossing the transformants to flies of the same *w^118* stock. Heterozygous lines 135/m and 97/m have the same background.

In line 235R11 (produced in Jeff Hall's lab) the *melanogaster nonA^+* insert carries a *ry^+* marker and is on a *ry^506* background. This background is therefore shared with the two lines 2*mel* and 3*mel*. 
RESULTS

Table 6.1 shows, for each genotype, the number of males out of 10 which gave a significantly correct optomotor response when tested in the optomotor apparatus, meaning 15 or more correct turns in 20 trials. A correct turn is achieved every time the fly turns in the same direction as the rotating stripes. A fly that performs 15 correct turns generates a significant $\chi^2$ value of 5.0 (df=1; p<0.05) where the null hypothesis is random turning (i.e. 10 correct, 10 incorrect turns). A fly giving only 14 correct turns ($\chi^2=3.2; 0.1>p>0.05$), or less, is therefore considered as visually compromised.

As shown in table 6.1, for small and large angle stripes, the genotype which stands out is $nonA^{diss}$ with not a single male giving significantly correct responses. All the others generally produce predominantly correct responses.

Table 6.1. Number of males giving significantly correct optomotor responses. S and L stand for small and large stripes. $diss$ refers to $nonA^{diss}$.

<table>
<thead>
<tr>
<th>Stripe</th>
<th>S</th>
<th>L</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>0</td>
<td>10</td>
</tr>
<tr>
<td></td>
<td>0</td>
<td>10</td>
</tr>
</tbody>
</table>

A $\chi^2$ test of homogeneity, on the results obtained for small stripes, gave a significant value of 94.7 (df=19; p<0.001). The result was still significant even after excluding from the test the data of $nonA^{diss}$ males ($\chi^2=39.01; df=18; p<0.01$). A
similar $\chi^2$ performed on the data for all genotypes, with large stripes, gave a significant value of 68.1 (df=19; $p<0.001$). When the $\text{non}A^{\text{dis}}$ data were not included, the $\chi^2$ value was still significant ($\chi^2=32.6; \text{df}=18; p<0.02$). A 2x2 contingency test was performed on the results obtained with the two different stripe widths (excluding the data from $\text{non}A^{\text{dis}}$), the null hypothesis being that small and large stripes give the same results. The null hypothesis was rejected ($\chi^2=4.03; \text{df}=1; p=0.045$ with Yates' correction). From table 6.1 it is clear that genotypes 135/d, 67-4/m, 168-8/m, perform comparatively poorly with large stripes.

This preliminary analysis can be further refined by taking into account, for each of the ten flies tested per genotype, the number of the correct turns performed out of twenty trials. Table 6.2 shows the results, obtained with these data, of a two way ANOVA with genotypes (18 in total) and stripe width (small or large) as the main factors. The data obtained from the $D.\text{virilis}$ and the $\text{non}A^{\text{dis}}$ genotypes were not included since the $\text{non}A^{\text{dis}}$ data were clearly different, and the optomotor test of $D.\text{virilis}$ flies had to be performed using a different method (see chapter 2). Thus only differences between transformant lines carrying one, two or three copies of $\text{melanogaster} \text{non}A^+$ and $\text{non}A^\text{vir}$ genes are analyzed. The test scores of the positive control line $\text{Canton S}$ were also included. Table 6.2 reveals a significant effect of Genotypes ($p=0.0000$) and Stripe width ($p=0.0014$). The interaction is also marginally significant ($p=0.0389$).

**Table 6.2.** Two way ANOVA of walking optomotor test

<table>
<thead>
<tr>
<th>SOURCE</th>
<th>MS</th>
<th>df</th>
<th>F</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>GENOTYPE</td>
<td>21.22</td>
<td>17</td>
<td>5.96</td>
<td>0.0000*</td>
</tr>
<tr>
<td>STRIPE</td>
<td>36.74</td>
<td>1</td>
<td>10.33</td>
<td>0.0014*</td>
</tr>
<tr>
<td>G*S</td>
<td>6.10</td>
<td>17</td>
<td>1.71</td>
<td>0.0389*</td>
</tr>
<tr>
<td>ERROR</td>
<td>3.56</td>
<td>324</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Figure 6.2 illustrates the effect of stripe size collapsed over genotype. Clearly the larger set of stripes leads to significantly more turning errors than the narrower stripes. However, the presence of a marginally significant interaction indicates that this is not the case for all the genotypes analyzed. This is evident from figure 6.3 which illustrates the influence exerted by a change in stripe width on each genotype. In this picture the results obtained with nonA\textsuperscript{dis} and \textit{D. virilis} were also included. Although for most genotypes a change of stripes into larger ones creates more incorrect responses, some genotypes behaved in the opposite way and actually performed better in response to large stripes (e.g. 97/d, 191/d, 297-6/d). In contrast, the response of other genotypes did not seem to be significantly influenced by the change of stripe size. This is the case of \textit{Canton S}, 67-4/d, 135/m and 97/m.

**Hemizygous nonA\textsuperscript{(+)}[nonA\textsuperscript{+}]**

As expected, the visual behaviour of nonA\textsuperscript{(-)} males carrying one copy of the \textit{melanogaster nonA}\textsuperscript{+} gene on their third chromosome (line 235R11/d on figure 6.3) showed no statistical difference from the results obtained using wild-type \textit{Canton S} flies (see Newman-Keuls test results in appendix 6.1), indicating complete rescue of the visual defect of nonA\textsuperscript{(-)} flies.

**Hemizygous nonA\textsuperscript{(+)}[nonA\textsuperscript{vir}].**

Inspection of figure 6.3 reveals that the lines carrying one copy of nonA\textsuperscript{vir} on a nonA\textsuperscript{(-)} background (from line 113/d to line 297-6/d) can be divided into two groups:

- lines 113/d, 67-4/d, and 168-8/d: these lines gave on average more correct turns than the other four lines, indicating a better rescue of the mutant visual defect. Their performance was not statistically different from that displayed by wild type flies or line 235R11/d (see appendix 6.1), revealing that the nonA\textsuperscript{vir} gene can function as
Figure 6.2. Main effect of stripe width (collapsed over genotype) on correct turning in walking optomotor test. Note that, in general, flies responded better to small stripes.

Figure 6.3. Average number of correct turns (out of 20 trials) given by each genotype on small and large stripes.
well as the melanogaster nonA\(^+\) gene in rescuing the mutant visual deficiencies. At this point it is useful to remark that, as explained in chapter 4, the P-element insert in line 113/d has lost the 5’P-element end and may be missing part of the 3’ end of the original nonA\(^{vir}\) construct. Moreover it appears that lines 67-4 and 168-8 are not independent lines from 113, so even though the nature of any molecular alteration in lines 168-8 and 67-4 is enigmatic (see discussion in chapter 4), the insert is in the same chromosomal position in all the three lines.

-lines 135/d, 97/d, 191/d, and 297-6/d: those which achieved the poorest results were line 135/d and line 191/d. Line 135/d behaved much worse when tested using the larger set of stripes. Its performance using large stripes is marginally poorer (p=0.043, see Newman-Keuls test’s results in appendix 6.1) than that of Canton S, but not from 235R11/d flies. However the transgene in this line is still rescuing the visual defect rather well, since its optomotor scores show no significant difference when compared to those obtained from Canton S on small stripes or 235R11/d on both small and large stripes. Line 191/d surprisingly performed more poorly when tested on small stripes than on large ones (even though the difference between stripes is not significant, see appendix 6.1). When tested on small stripes, line 191/d has poorer scores than 235R11/d (p=0.038) and 113/d flies (p=0.040, see appendix 6.1). Therefore line 113 and its derivatives 67-4 and 168-8 rescue visual behaviour as well as the mel nonA\(^+\) transgene, and better than the intact nonA\(^{vir}\) transgenes 135, 97, 191, and 297-6. Lines 97/d and 297-6/d are not significantly different from Canton S, 235R11/d, 113/d, 67-4/d, or 168-8/d flies, for both sets of stripes.

**Heterozygous nonA\(^{vir}/nonA\(^+\)**

Inspection of figure 6.3 reveals that the six lines analyzed (from line 112/m to line 97/m) showed similar scores to their hemizygous counterparts. Line 168-8/m however showed relatively poor visual behaviour when tested on large stripes. Its performance was significantly poorer than the wild-type (p=0.005, Newman-Keuls test, appendix 6.1) and 235R11/d when tested on large stripes (p=0.006). Interestingly
line 168-8/m, when using large stripes, produced significantly more errors compared to the insert on the nonA\(^{(v)}\) background (line 168-8/d, \(p=0.007\)). The same happened in the case of 67-4/m. When tested on large stripes, this line gave significantly less correct turns when compared to line 67-4/d (\(p=0.048\)). Line 112/m, which carries one copy of nonA\(^{\text{vir}}\) inserted on the X chromosome was not significantly different when compared to Canton S or to flies carrying only one copy of the melanogaster nonA\(^+\) gene or one copy of nonA\(^{\text{vir}}\). In conclusion, for some inserts (113, 67-4, and 168-8) there is some evidence that adding a copy of the mel nonA\(^+\) gene increases the number of optomotor errors. However, for inserts 135, 97, 191, 297-6, there is little difference.

**2 mel**

Adding one extra copy of the gene nonA essentially had very little effect on the visual behaviour of the flies. As shown in figure 6.3, line 2mel showed a poorer performance when tested on large stripes, but in any case this performance was not significantly poorer than the one obtained with wild-type Canton S flies or with flies carrying only one copy of nonA\(^+\) (line 235R11/d).

**3 mel**

When tested on either stripes, flies carrying three copies of the mel nonA\(^+\) gene scored a little worse than flies carrying two copies of the gene but the difference is not statistically significant. The visual behaviour of flies carrying three doses of nonA\(^+\) was also not significantly different from the one displayed by Canton S flies or flies carrying only one copy of the mel nonA\(^+\) transgene.

**mel+2 vir**

The performance of flies carrying one dose of the mel nonA\(^+\) gene and two copies of the nonA\(^{\text{vir}}\) insert showed no significant difference to that of flies carrying only one copy of either gene.
DISCUSSION

In this chapter the visual behaviour of transformant flies carrying one, two or three copies of nonA+ and/or nonAvir has been analyzed using the walking optomotor test. It was confirmed that a transformant line carrying the putative D.melanogaster nonA+ gene (235R11; Jones and Rubin, 1990) completely rescues the visual defect of nonA(‘) flies (Rendahl et al., 1992).

In this study two different set of stripes were used in the optomotor test. The optomotor response is known to be dependent on the pattern wavelength of the visual stimulus (Götz, 1964; Heisenberg and Buchner, 1977; Blondeau and Heisenberg, 1982). The pattern wavelength \( \lambda \) is given by the angle subtended by a pair of black and white stripes. In the experiments reported in this chapter the pattern wavelength \( \lambda \) was 26.6° for the small stripes, and 72° in the case of the large stripes. An average response curve for walking wild-type flies, as a function of the pattern wavelength used, was first produced by Götz (1964), and is presented by Heisenberg and Buchner (1977). The optomotor response tends to be at an optimum when pattern wavelengths between 55° and 19° are used, while it decreases precipitously at larger, and rather more gently at smaller values. Based on these observations, it would be expected that the large stripes used in the experiments described in this chapter (\( \lambda=72° \)) represent a more challenging test when compared to the small ones. In contrast, the small ones should be eliciting the optimum optomotor response, under the same conditions. Indeed it was found that almost all genotypes performed better when tested on small stripes. However wild type flies were influenced very little by the change in wavelength pattern.

All the transformants carrying one copy of the putative nonAvir gene seem to rescue the visual defect of nonA(‘) rather well. I could not directly analyze nonA(‘)/Y males as they were almost impossible to obtain. Instead I used nonA\(_{dis} \) mutant males as the negative control. In my test, nonA\(_{dis} \) males cannot see the rotation of the stripes, as in Kulkarni et al. (1988) and Rendahl et al. (1992). Line 113/d and its
derivatives (line 67-4/d and 168-8/d) displayed the best optomotor responses indicating a complete rescue of the visual phenotype. Interestingly these lines carry a nonA
\textsuperscript{vir} insert that appears to have lost some non-coding sequence at its 3' end (see chapter 4). The fact that these lines completely rescue the poor nonA\textsuperscript{(c1)} optomotor response gives an indirect proof that their insert still carries the complete sequence for the fifth exon. This exon is in fact indispensable for the rescue of all the nonA-related biological functions (Jones and Rubin, 1990, Rendhal et al., 1992). Whatever sequence has been lost from the transgene in lines 113, 67-4, and 168-8, it has to carry no biological function related to visual behaviour. This is particularly interesting when compared to the situation found in the case of the complex optomotor-blind (omb) gene. At the core of this locus lies a lethal complementation group (lll/omb) whose mutations cause larval and pupal lethality (Plugfelder et al., 1990; Plugfelder et al., 1992). The recessive viable allele In(1)omb\textsuperscript{H31} which causes neuroanatomical and optomotor defects, is the result of an inversion affecting a proximal, non-transcribed part of the locus. This 40 Kb downstream regulatory region has been termed "OLR" for optic lobe regulatory region. Brunner et al. (1992) have shown that removing increasing amounts of DNA from this region results in a gradient of increasing severity of the neuroanatomical and visual defects.

On the basis of the results described above, it seems more likely that, in the case of nonA, all the neuroanatomical information related to vision is restricted to the coding region or to other sequences not very far from it. Recent studies involving in-vitro mutagenesis of nonA in order to study the behavioral effects of mutations affecting NONA's putative RNA-binding motives (Rendahl et al., 1996; Stanewsky et al., 1996), suggest that the N-terminal RNA-recognition motif (RRM1) is necessary for all the known biological functions of NONA. However, the C-terminal RRM2 domain appears to be primarily involved in vision. In agreement with this stands the fact that the chemically induced mutants with specific defects to the visual system (nonA\textsuperscript{H2}, nonA\textsuperscript{P14}) also map in or near the RRM2 domain (Rendahl et al., 1996). Lines 135 and 191 seem to be slightly less effective than the other lines in rescuing the nonA\textsuperscript{(c1)} defect, although their phenotype is far from being mutant. These discrepancies could simply be the result of position effects which imply that a particular genomic location
of the transduced DNA insert could lead to a lower than normal expression of the transformed gene (Hazelrigg et al., 1984; Baylies et al., 1987). Among all the in vivo and in vitro mutations isolated so far, not a single one has ever affected solely the singing behaviour. It seems as if, whenever a mutation in nonA occurs, the "first to go" is always the visual phenotype (Rendahl et al., 1992; Rendahl et al., 1996; Stanewsky et al., 1996). Bearing this in mind, one could be tempted to think that a transgene rescuing the visual defect should consequently fully rescue the song phenotype as well.

However there have been cases in which particular nonA+ transgenes seemed to rescue better the visual defect than song phenotype. Rendahl et al. (1992) found that two transformed lines, carrying the 11Kb EcoRI genomic fragment from Jones and Rubin (1990), restored visual function better than the song phenotype in nonA diss mutants. These investigators tried to explain this finding suggesting the possibility that the 11Kb fragment could be missing some particular "thoracic enhancers" causing poor restoration of the biological activity necessary for normal song. They argued that maybe larger genomic fragments might fully restore both nonA associated phenotypes. However this finding could be more parsimoniously explained as a result of a position effect, since other lines carrying the same fragment fully rescued the visual and song phenotypes (Rendahl et al., 1996). With this in mind it is critical to analyze many independent lines carrying a particular DNA insert. If at least one of these lines can rescue both phenotypes, then the transduced DNA fragment has to contain all the essential enhancers for the relevant behaviours. In this study genomic DNA containing the putative nonA vir gene seemed to rescue the optomotor deficiencies of nonA ( ) flies. Thus there is a strong possibility that the same fragment will also rescue the mutant song phenotype. Any deviation of the transformants' song pattern from wild-type could thus be due to species-specific song information encoded within nonA vir.

The changes in behavioural phenotype caused by one or more copies of a particular transgene have been studied with the clock gene per. Males and females carrying one versus two copies of the same transgene produce alterations in the circadian period (Cooper et al., 1994). In the case of per, the locomotor activity
rhythm of the fly is shortened by increasing the levels of *per* expression and lengthened by decreasing them (Cooper *et al.*, 1994; Baylies *et al.*, 1987). The possibility of a dosage effect of the *nonA*\(^+\) gene on visual behaviour was tested by Stanewsky *et al.* (1996). These investigators studied the optomotor response of flies carrying two or three doses of the *nonA*\(^+\) gene. They found that males of one line which carried a homozygous *nonA*\(^+\) insert on a *nonA*\(^+\) background (i.e. carrying three copies of *nonA*\(^+\)) seemed to have a mutant optomotor response. However this turned out to be due to the site of insertion of that particular transgene, since the same insert, when tested with a different *nonA*\(^+\) insert, plus the endogenous *nonA*\(^+\) gene, gave normal visual behaviour.

In this chapter the optomotor response of flies carrying two or three copies of *nonA*\(^+\) has been tested again, but using a different *nonA*\(^+\) insert than the one described by Stanewsky *et al.* (1996). The results with these transformants confirm the findings of Stanewsky *et al.* that increasing the doses of *nonA*\(^+\) has no influence on the visual behaviour. Here we also tested the effect of adding one or two doses of *nonA*\(^{vir}\) to flies with a *nonA*\(^+\) background. In two cases (168-8/m and 67-4/m), when compared to hemizygous males carrying *nonA*\(^+\) or *nonA*\(^{vir}\), a significant increase of the number of errors made in the optomotor test was observed. Furthermore a similar result is obtained with line 113. However, 168-8 and 67-4 were derived from 113, and these effects are not observed with the other lines where the insert appears to be intact. When males bearing one dose of *nonA*\(^+\) and two copies of *nonA*\(^{vir}\) were tested on the optomotor (*mel+2vir*), they performed similarly to hemizygous males carrying only one copy of either gene. Thus increasing the dose of the *nonA*\(^{vir}\) gene seems to have no effect on optomotor behaviour.

Given the successful rescue of optomotor behaviour, song analysis in these transformants should provide a reasonably unambiguous interpretation of whether the *nonA*\(^{vir}\) gene rescues the mutant phenotype. These song studies are reported in the next chapter.
CHAPTER 7

Song analysis
INTRODUCTION

This chapter presents a detailed analysis of the courtship song produced by the same transformant lines whose visual behaviour has been described in the previous chapter. From the results of chapter 6 it is clear that the nonAvir transgene is able to rescue the visual defect of nonA\(^{(c)}\) flies. Here the lovesongs produced by nonA\(^{(c)}\) males carrying the nonAvir transgene will be compared to the songs generated by nonA\(^{dis}\) mutants, nonA\(^{(c)}\)[nonA\(^{+}\)] transformants, and D.virilis males. The aim is to investigate whether the nonAvir gene rescues the nonA\(^{(c)}\) song phenotype, and whether it carries any species-specific song characteristic. The songs of heterozygous nonA\(^{+}\)/nonAvir, and 2mel, 3mel and mel+2vir (see chapter 6) males will also be analysed in order to test the possibilities of nonA dosage effects, and if the nonAvir gene carries any semidominant virilis-like song characteristics.

The D.virilis song differs from that of D.melanogaster in several quantitative aspects. Firstly, the sine song component, which, from my own observations, in D.melanogaster can often account for as much as half of the total song produced, is completely absent from the songs produced by D.virilis (Hoikkala, 1985). Figure 7.1 shows clearly the differences between the pulse song of D.melanogaster and D.virilis. D.virilis pulses are more polycyclic than those of D.melanogaster (range of 4.5-6.5 cycles compared to the 1-3 of D.melanogaster). Furthermore, the average D.virilis intrapulse frequency (PF) is higher, usually > 400 Hz in D.virilis compared to < 350 Hz in D.melanogaster (see results). In addition, pulse trains are shorter in D.virilis (rarely containing more than 10-11 pulses), and many of the pulses tend to fuse together so that there are short gaps between the end of one pulse and the beginning of the next. Consequently, D.virilis IPIs are shorter than those of D.melanogaster (Hoikkala, 1982).

The nonA\(^{dis}\) mutant lovesong has already been described in chapter 1. Its salient feature is the tendency of pulses towards the end of long trains (containing > 6 pulses) to become high both in cycles number (CPP) and amplitude (Kulkarni et al.,
Figure 7.1. Pulse song of *D. melanogaster*, *nonA\textsuperscript{dis}*, and *D. virilis*. 
Figure 7.1. Pulse song of *D.melanogaster*, nonA<sup>diss</sup> and *D.virilis*. 
1989, see figure 7.1). Wheeler et al. (1989) also extended the study of these mutant songs and found previously undetected qualitative aberrations in the pulse and sine waveform shapes. In this chapter only quantitative characteristics of the songs will be taken into consideration. The parameters studied for each song were described fully in chapter 2 and are:

- **CPP**: cycles per pulse.
- **bCPP**: value of the CPP regression slope.
- **SSP**: sine song proportion.
- **PF**: pulse frequency.
- **bPF**: value of the PF regression slope.
- **SSF**: sine song frequency.
- **IPI**: mean interpulse interval.

**MATERIALS & METHODS**

**GENOTYPES**

The genotypes analyzed were the same as those tested for optomotor behaviour (see chapter 6). *nonA* mutant songs were recorded from males of the strain *diss/Y; ry^{506}/MKRS*. Other *diss* strains tested failed to produce an easily recognizable mutant song. These stocks had been kept in J.C. Hall's lab for several years and had probably accumulated modifiers. No songs were recorded from the *nonA ( ) genotype (see chapter 5) since no flies of this genotype could ever be generated (see chapter 6). Line 235R11/d (see chapter 6) provided wild-type control songs. The same line had already been used as the positive control (under a different name, J. Hall, personal communication) in the song analysis performed by Rendahl et al. (1996). Therefore I expected to obtain reliable *melanogaster*-like wild-type songs from it.
SONG ANALYSIS

10 males from each genotype were recorded, as described in chapter 2. For most genotypes 10 min of song from each fly were processed and analyzed using the Spike2 program and a Microsoft Excel 5.0 macro written by M. Couchman, as described in chapter 2. Only 6 songs were analysed for the genotype 67-4/m. 5 min of song only were analyzed from flies of the genotypes diss, 97/d, 191/d, 296-6/d, and 135/m and vir.

RESULTS

It became clear during song recording that some flies sang more than others. In some cases the difference could be over an order of magnitude. For example, within ten minutes of recording, one fly of the 3mel genotype produced a total of 171 seconds of song, while one fly of the 113/m genotype sang for a total of only 15 seconds. If any of the song characters, such as CPP or bCPP, changed systematically in relation to the amount of song, then this would have to be covaried out of the song measure. For example, imagine that one fly sang twice as much as another, and CPP increased linearly with the amount of song, perhaps due to fatigue. An adjustment would have to be made between the two flies' CPP values, via regression, so that they could be compared in the absence of any differences in global song output.

As an index of song output I decided to use the sum of the total amount of pulse song plus the total amount of sine song produced by each fly (PS+SS). PS and SS were calculated as described in chapter 2. For each genotype, Pearson product-moment correlations were then performed between the PS+SS parameter and each of the song characters of interest. The results are presented in table 7.1.
Table 7.1. Correlation matrix between song output parameters (PS+SS) and other song characters in each genotype. CPP= cycles per pulse. bCPP= cycles per pulse gradient. SSP= proportion of sine song over the total amount of song (pulse song + sine song). PF= pulse frequency. bPF= pulse frequency gradient. SSF= sine song frequency. IPI= mean interpulse interval. * indicates significant r values at p <0.05. **indicates significant r at p<0.01.

<table>
<thead>
<tr>
<th>PS+SS</th>
<th>CPP</th>
<th>bCPP</th>
<th>SSP</th>
<th>PF</th>
<th>bPF</th>
<th>SSF</th>
<th>IPI</th>
</tr>
</thead>
<tbody>
<tr>
<td>diss</td>
<td>-0.56</td>
<td>0.54</td>
<td>0.82**</td>
<td>-0.19</td>
<td>0.31</td>
<td>0.68*</td>
<td>0.03</td>
</tr>
<tr>
<td>235R11/d</td>
<td>0.81**</td>
<td>-0.50</td>
<td>0.41</td>
<td>-0.40</td>
<td>0.46</td>
<td>0.18</td>
<td>-0.15</td>
</tr>
<tr>
<td>113/d</td>
<td>-0.30</td>
<td>-0.12</td>
<td>0.71*</td>
<td>-0.29</td>
<td>-0.04</td>
<td>0.25</td>
<td>0.18</td>
</tr>
<tr>
<td>67-4/d</td>
<td>-0.03</td>
<td>-0.18</td>
<td>-0.01</td>
<td>-0.23</td>
<td>0.42</td>
<td>0.52</td>
<td>0.04</td>
</tr>
<tr>
<td>168-8/d</td>
<td>0.20</td>
<td>0.34</td>
<td>0.72*</td>
<td>-0.60</td>
<td>0.44</td>
<td>0.40</td>
<td>-0.06</td>
</tr>
<tr>
<td>135/d</td>
<td>0.65*</td>
<td>0.05</td>
<td>-0.03</td>
<td>0.22</td>
<td>0.15</td>
<td>0.48</td>
<td>-0.13</td>
</tr>
<tr>
<td>97/d</td>
<td>0.27</td>
<td>-0.04</td>
<td>0.46</td>
<td>-0.43</td>
<td>0.27</td>
<td>0.51</td>
<td>-0.54</td>
</tr>
<tr>
<td>191/d</td>
<td>0.53</td>
<td>0.11</td>
<td>0.32</td>
<td>-0.41</td>
<td>0.24</td>
<td>-0.04</td>
<td>-0.11</td>
</tr>
<tr>
<td>297-6/d</td>
<td>0.16</td>
<td>0.44</td>
<td>0.73*</td>
<td>-0.37</td>
<td>0.22</td>
<td>0.05</td>
<td>0.24</td>
</tr>
<tr>
<td>112/m</td>
<td>0.60</td>
<td>-0.16</td>
<td>0.45</td>
<td>-0.42</td>
<td>-0.06</td>
<td>-0.71*</td>
<td>0.70*</td>
</tr>
<tr>
<td>113/m</td>
<td>0.59</td>
<td>-0.24</td>
<td>0.56</td>
<td>-0.62</td>
<td>0.58</td>
<td>0.00</td>
<td>-0.03</td>
</tr>
<tr>
<td>67-4/m</td>
<td>-0.63</td>
<td>-0.34</td>
<td>0.92**</td>
<td>-0.64</td>
<td>0.75</td>
<td>-0.18</td>
<td>-0.21</td>
</tr>
<tr>
<td>168-8/m</td>
<td>-0.59</td>
<td>0.26</td>
<td>0.74*</td>
<td>-0.15</td>
<td>-0.31</td>
<td>0.11</td>
<td>0.17</td>
</tr>
<tr>
<td>135/m</td>
<td>0.59</td>
<td>-0.33</td>
<td>0.87**</td>
<td>-0.61</td>
<td>0.57</td>
<td>-0.07</td>
<td>0.29</td>
</tr>
<tr>
<td>97/m</td>
<td>-0.32</td>
<td>-0.04</td>
<td>0.22</td>
<td>-0.60</td>
<td>0.58</td>
<td>-0.17</td>
<td>0.49</td>
</tr>
<tr>
<td>2mel</td>
<td>0.20</td>
<td>0.03</td>
<td>0.34</td>
<td>-0.53</td>
<td>0.60</td>
<td>0.28</td>
<td>0.17</td>
</tr>
<tr>
<td>3mel</td>
<td>0.54</td>
<td>-0.65*</td>
<td>0.28</td>
<td>-0.25</td>
<td>0.63</td>
<td>-0.32</td>
<td>0.05</td>
</tr>
<tr>
<td>mel+2vir</td>
<td>0.13</td>
<td>-0.49</td>
<td>0.33</td>
<td>-0.69*</td>
<td>0.81**</td>
<td>0.60</td>
<td>-0.20</td>
</tr>
<tr>
<td>vir</td>
<td>0.76*</td>
<td>-0.64*</td>
<td>--</td>
<td>-0.51</td>
<td>-0.24</td>
<td>--</td>
<td>0.41</td>
</tr>
</tbody>
</table>

From table 7.1 it is clear that no consistent relationship between PS+SS and characters such as CPP, bCPP, bPF, SSF or IPI can be found. In some genotypes the correlation between song output index and each one of these parameters is positive, while in some others it is negative. Furthermore in almost all the cases these correlations are not significant. A different result is found when correlating PS+SS with SSP. In this case almost all the genotypes display a positive correlation between these two characters, and in several genotypes the relationship is highly significant. PF also seems to be related to the song output parameter: consistent negative (but largely non-significant) correlations between PS+SS and PF can be found for all the genotypes except one.
In conclusion, there is no compelling evidence that characters such as CPP, bCPP, or bPF, which could potentially be used to distinguish between species-specific, *virilis*-like song rescue, or incomplete, *diss-*mutant like rescue, are related to quantity of song output. However, SSP and PF do seem to genuinely related to the amount of song generated, so caution must be exercised in using these characters, and any effect of differential song production between genotypes must be removed before any comparison.

**bCPP.**

The salient feature of a mutant *nonA*mutant song is that the number of cycles per pulse (CPP) is not constant during each single pulse train. In long trains (containing more than 5-7 pulses) the pulses toward the end of the train become increasingly more polycyclic, sometimes bearing up to 30 cycles per pulse. Wild type songs never show such a feature and in general their pulse trains contain a constant number of cycles per pulse (usually between 1 and 3) (Kulkarni et al., 1987).

The first song feature to be taken into consideration in the analysis of the songs recorded was therefore the CPP regression gradient (bCPP). A CPP least squares regression was performed for each song as described in chapter 2. The bCPP gives an indication, for each song, of how the pulses change during pulse trains: a significant positive bCPP indicates a tendency of the number of CPPs to increase as the train of pulses progresses. A non significant, very small bCPP will suggest a tendency of the pulses to remain constant during pulse trains.

For each genotype, the number of songs out of ten which had a significant CPP regression gradient (bCPP) was calculated (table 7.2). For genotype 67-4/m this number was calculated out of only 6 songs analyzed. The significant gradients were divided into positive or negative. A negative gradient would suggest that the pulses at the beginning of the train were actually more polycyclic than those towards the end. The average value of bCPP was also calculated. Notice that the genotypes which had
very few songs with significant regressions also had a very small average bCPP value (see 113/m or 3mel). In these genotypes it seems that the CPP remains constant during the progression of a pulse train.

Table 7.2. bCPP (cycle per pulse regression gradient) for different genotypes. N= number of significant bCPPs out of 10 songs (* out of 6 songs). + = significant positive bCPPs. − = significant negative bCPPs. The mean bCPP value per genotype is also indicated.

<table>
<thead>
<tr>
<th>GENOTYPE</th>
<th>N</th>
<th>+</th>
<th>−</th>
<th>Mean bCPP</th>
</tr>
</thead>
<tbody>
<tr>
<td>diss</td>
<td>10</td>
<td>10</td>
<td>0</td>
<td>0.257</td>
</tr>
<tr>
<td>235R11/d</td>
<td>4</td>
<td>1</td>
<td>3</td>
<td>-0.002</td>
</tr>
<tr>
<td>113/d</td>
<td>7</td>
<td>7</td>
<td>0</td>
<td>0.004</td>
</tr>
<tr>
<td>67-4/d</td>
<td>8</td>
<td>8</td>
<td>0</td>
<td>0.019</td>
</tr>
<tr>
<td>168-8/d</td>
<td>8</td>
<td>8</td>
<td>0</td>
<td>0.016</td>
</tr>
<tr>
<td>135/d</td>
<td>8</td>
<td>8</td>
<td>0</td>
<td>0.022</td>
</tr>
<tr>
<td>97/d</td>
<td>7</td>
<td>6</td>
<td>1</td>
<td>0.012</td>
</tr>
<tr>
<td>191/d</td>
<td>7</td>
<td>6</td>
<td>1</td>
<td>0.024</td>
</tr>
<tr>
<td>297-6/d</td>
<td>7</td>
<td>6</td>
<td>1</td>
<td>0.014</td>
</tr>
<tr>
<td>112/m</td>
<td>5</td>
<td>3</td>
<td>2</td>
<td>0.001</td>
</tr>
<tr>
<td>113/m</td>
<td>2</td>
<td>0</td>
<td>2</td>
<td>-0.003</td>
</tr>
<tr>
<td>67-4/m*</td>
<td>4</td>
<td>4</td>
<td>0</td>
<td>0.014</td>
</tr>
<tr>
<td>168-8/m</td>
<td>5</td>
<td>5</td>
<td>0</td>
<td>0.010</td>
</tr>
<tr>
<td>135/m</td>
<td>5</td>
<td>4</td>
<td>1</td>
<td>0.005</td>
</tr>
<tr>
<td>97/m</td>
<td>3</td>
<td>3</td>
<td>0</td>
<td>0.004</td>
</tr>
<tr>
<td>2mel</td>
<td>8</td>
<td>7</td>
<td>1</td>
<td>0.009</td>
</tr>
<tr>
<td>3mel</td>
<td>2</td>
<td>0</td>
<td>2</td>
<td>0.000</td>
</tr>
<tr>
<td>mel+2vir</td>
<td>3</td>
<td>2</td>
<td>1</td>
<td>0.001</td>
</tr>
<tr>
<td>vir</td>
<td>10</td>
<td>10</td>
<td>0</td>
<td>0.186</td>
</tr>
</tbody>
</table>

As expected, all the diss song recorded had a strongly significant positive gradient and the average value of the positive slope was high. Unexpectedly, the results obtained with vir were strikingly similar to those given by diss: all the songs recorded had a significantly positive regression slope and the average value of bCPP was also very similar to that found in diss. Only four songs from the 235R11/d genotype had a significant CPP regression and in three this regression had a negative
slope. The average bCPP for this genotype had a slightly negative value. Therefore in line 235R11/d the amount of CPP did not increase with the progressing of the pulse train as opposed to what happens in *diss*, thus confirming that this line, which carries one copy of *D.melanogaster nonA* over a *nonA(·)* background, gives effective wild-type rescue of this song character (Rendahl *et al.*, 1992; 1996).

All the transformant lines carrying one copy of *nonA* over *nonA(·)* background (from genotype 113/d to 297-6/d in table 7.2) produced songs with mostly significant positive CPP regressions. Also the average value of bCPP was positive in all these lines even though it was generally at least 10 times smaller than the average slope value for *diss* regressions (see also figure 7.2a). Heterozygous lines carrying one copy of *nonA* plus the endogenous *nonA* gene (genotypes 112/m to 97/m in table 7.2) had a much lower number of significant regression slopes when compared to their hemizygous counterparts. Also some of the significant slopes were negative (in 113/m 2 out of 2 significant slopes were negative). The average bCPP had also much lower values in these genotypes if compared with those of the *nonA* hemizygous lines. In general these values resembled closely those found for line 235R11/d. The only two exceptions were perhaps lines 67-4/m and 168-8/m for which all the significant regression gradients were positive and the average bCPP values were about ten times greater than those from the other heterozygous lines. A t-test comparing the proportions of positive significant slopes in hemizygous *nonA(·)[nonA]* lines with those in heterozygous *nonA*/*nonA* revealed a significant difference between these two groups (*t*=3.701; *p*=0.003).

Line 2*mel* which carries two copies of the *nonA* gene had twice as many significant CPP regressions than the line carrying one copy of *nonA* (235R11). Also almost all the gradients were positive. Consequently the average bCPP value was also slightly positive. Line 3*mel* carrying three copies of *nonA* had only two significant regressions both of them with a negative slope. The average value of bCPP for this genotype was 0 indicating that the song pulses maintain the same number of cycles throughout the pulse trains. The results obtained from the line carrying two copies of *nonA* and one of *nonA* (*mel+2vir*) were very similar to those of line 3*mel*. Both
lines behave very similarly to line 235R11/d, indicating that no consistent dosage effect was exerted on this song character, in spite of differences between line 2mel and line 235R11/d.

One way ANOVA was performed using the bCPP values of the songs analyzed for each genotype. When the data for all the genotypes was included, the result was highly significant (F= 11.155; df = 18, 167; p << 0.001). When the bCPP values of diss and vir were excluded from the analysis (note in figure 7.2a the huge variance in bCPP values for these two genotypes), one-way ANOVA still gave a significant main effect (F= 4.263; df= 16, 149; p << 0.001). Figure 7.2a shows the mean, the standard error and standard deviation of the bCPP values in each genotype, and figure 7.2b illustrates the same values for the transformant lines only. Newman-Keuls a posteriori test revealed that all the transformant lines were significantly different from both diss and vir.

From figure 7.2b it is evident that all the hemizygous nonA(\nonAvir) lines have, in general, higher bCPP values than the other transformant lines. A planned comparison between the control line 235R11/d and all the hemizygous nonA(\nonAvir) lines (from line 113/d to line 297-6/d, in figure 7.2) revealed that these two groups differ significantly (F= 17.321, df= 1, 149; p << 0.001). In particular, a Newman-Keuls test, performed excluding the data from vir and diss (see appendix 7.1), revealed that lines 67-4/d, 135/d and 191/d differ significantly from the positive control line 235R11/d. Heterozygous nonAvir/nonA+ lines showed, in general, similar or slightly higher bCPP values than those of the control 235R11/d. A planned comparison between all the nonAvir/nonA+ lines (from line 112/m to line 97/m in figure 7.2) and line 235R11/d showed that these two groups do not differ significantly (F= 2.824; df= 1, 149; P= 0.095) from each other.

From figure 7.2b it is clear that all the hemizygous nonA(\nonAvir) lines have higher bCPP values than the heterozygous nonAvir/nonA+ group. A planned comparison between these two groups showed in fact that they significantly differ (F= 20.921; df= 1, 149; p << 0.001) from each other. When comparing each hemizygous nonA(\nonAvir) line with its heterozygous nonAvir/nonA+ counterpart (for example
Figure 7.2.

A: bCPP values of all genotypes.
B: bCPP values of transformant lines only.
113/d with 113/m, in figure 7.2b) it is evident that the hemizygous lines have always higher bCPPs than their heterozygous counterparts. (although not significantly, see Newman-Keuls results in appendix 7.1)

bCPP values of line 2mel are not significantly different from the positive control 235R11/d, neither are those of lines 3mel and mel+2vir (see Newman-Keuls in appendix 7.1), indicating that there are no dosage effects on this song character.

In conclusion, results suggest that the bCPP values of hemizygous nonA(−) [nonA vir] lines are in general higher than those of the control line 235R11/d and the other transformant lines. This implies that in these genotypes there is a slight tendency of the cycles per pulse to increase towards the end of pulse trains. However the bCPP values of these lines are still closer to those of the control line (note that only lines 67-4/d, 135/d and 191/d differed significantly from 235R11/d) than to those of either diss or vir. These results can be interpreted in two ways. We could assume that the rescue of this song character in nonA(−)/nonAv, r lines is not complete, so the CPPs still tend to increase slightly toward the end of each song. However vir songs also show a tendency of CPP to increase during pulse trains (the bCPP average values of vir are in fact very similar to those of diss, see table 7.1) so we could also argue that the higher bCPP values might be the result of a species-specific effect exerted by the nonAvir gene. Other songs characters have therefore to be analyzed in order to try to distinguish between these alternatives.

CPP.

D. virilis pulse songs are in general more polycyclic than those of D. melanogaster, the average CPP value of virilis falling between 4 and 6 (Hoikkala et al., 1982), as opposed to 1-2 in melanogaster. In the analysis of song produced by transformant lines it was essential to check whether transformants carrying the nonAvir gene produced more polycyclic songs than transformants carrying the nonA+ gene.
For each song, CPP values can be simply extrapolated from the CPP regression curve $y=bx+c$ at different values of $x$, where $x$ represents pulse position in a train. An alternative analysis could consist of using the raw CPP values found in each song. Both approaches were used. In the first kind of analysis CPP values were calculated using the CPP regression of each song when $x=1$ (CPP1) and $x=10$ (CPP10). The second analysis consisted in calculating in each song how many pulses could be found that had a CPP value of 1, 1.5, 2, 2.5, 3, 3.5, 4, and so on. The results are presented below.

**CPP1.**

The CPP values based on the regression function (CPP=$bx+c$) when $x=1$ were calculated for each song. An one way ANOVA was employed to analyze these data. When the CPP1 values from all genotypes were used, the test gave a highly significant $F$ ratio ($F= 22.696; \text{df} = 18, 167; p << 0.001$). From figure 7.3a it is clear that in vir the first pulse in a train contains a relatively high number of cycles, and that this number tends to be quite variable. Large variation is found in the value of CPP1 of diss as well, even though in this genotype the average CPP1 value is lower than in vir, and more similar to that of the transformant lines. Newman-Keuls comparisons confirm that no significant difference exist between the average CPP1 value of diss and that of the transformant lines.

The ANOVA was repeated including only CPP1 values from the transformant lines. The $F$ ratio was highly significant ($F= 15.501; \text{df} = 16, 149; p << 0.001$). Figure 7.3b shows the mean, standard error and standard deviations of CPP1 values of the transformant lines only. A planned comparison between the control line 235R11/d and the group composed of of hemizygous non$A^{c[nonA^{vir}]}$ lines showed that they differ significantly ($F= 16.413; \text{df} = 1, 149; p << 0.001$). In particular, Newman-Keuls a posteriori comparisons (appendix 7.2) revealed that CPP1 values in 67-4/d, 168-8/d, 135/d and 297-6/d are significantly higher than in the control line 235R11/d.
Figure 7.3.

A: CPP values at x=1 for all genotypes.
B: same as in A excluding diss and vir.
C: CPP values at x=10 for all genotypes.
A planned comparison showed that even the group composed of all the heterozygous nonA\textsuperscript{vir}/nonA\textsuperscript{+} lines differs significantly from the control line (F= 36.084; df= 1, 149; p<< 0.001). Heterozygous nonA\textsuperscript{vir}/nonA\textsuperscript{+} lines also differ from the hemizygous nonA\textsuperscript{c}[nonA\textsuperscript{vir}] ones (F=14.870; df= 1, 149; p<< 0.001). From figure 7.3b it is clear that lines 67-4/m and 168-8/m have much higher CPP1 values than any other transformant line. These two lines significantly differ from all the others (see Newman-Keuls results in appendix 7.2). In general, with the exception of line 135, heterozygous nonA\textsuperscript{vir}/nonA\textsuperscript{+} lines bear higher CPP1 values than their hemizygous nonA\textsuperscript{c} [nonA\textsuperscript{vir}] counterparts, however these differences are not always statistically significant (Newman-Keuls, appendix 7.2).

Interestingly, line 2mel (carrying 2 copies of nonA\textsuperscript{+}) also significantly differs from line 235R11/d (p= 0.0436, see appendix 7.2) suggesting a possible dosage effect. However having 3 copies of nonA\textsuperscript{+} (3mel) or one copy of nonA\textsuperscript{+} and two of nonA\textsuperscript{vir} (mel+2vir) seems not to have much effect on CPPs since these lines do not differ significantly from 235R11/d (see appendix 7.2).

**CPP10.**

The CPP value when x=10 was calculated using the regression function of each song. The data was tested with one-way ANOVA which gave a highly significant F ratio (F= 52.501; df= 18, 167; p<< 0.001). As expected, figure 7.3c shows how the average CPP values at pulse 10 have more than doubled in diss since the first pulse (compare with figure 7.3a). The average CPP values at pulse 10 are also higher than in pulse 1 in vir songs. Slight increases of the CPP number are observed also in the transformant lines.

One-way ANOVA, excluding the data from diss and vir, resulted in a highly significant F ratio (F= 19.555; df= 16, 149; p<< 0.001). A posteriori Newman-Keuls comparisons (see appendix 7.3) showed that at pulse 10 all the transformant hemizygous nonA\textsuperscript{c}[nonA\textsuperscript{vir}] lines, except 113/d and 97/d, are significantly more
polycyclic than the control line 235R11/d. Also all the heterozygous nonA\textsuperscript{vir}/non-A\textsuperscript{+} lines, except 112/m and 113/m, differ significantly from line 235R11/d. 67-4/m and 168-8/m are again significantly more polycyclic than any other line. All heterozygous nonA\textsuperscript{vir}/nonA\textsuperscript{+} lines are still in general more polycyclic than their hemizygous nonA\textsuperscript{(c)} [nonA\textsuperscript{vir}] counterparts, with the only exception of 135/m. At pulse 10 the CPP values of 3mel and mel+2vir are still not significantly different from the control line 235R11/d, while 2mel is significantly more polycyclic (see appendix 7.3).

To summarize, hemizygous nonA\textsuperscript{(c)}[nonA\textsuperscript{vir}] lines are in general more polycyclic than the control line 235R11/d both at pulse 1 and 10. However at pulse 10 they also significantly differ from either diss or vir (which are much more polycyclic). Heterozygous nonA\textsuperscript{vir}/nonA\textsuperscript{+} lines are also more polycyclic than the control line at both pulse 1 or 10. In general, they are also more polycyclic than their hemizygous counterparts, even though pairwise comparisons are not always significantly different. Again, these results could be interpreted in two ways. We could assume that the increased polycyclicity seen in nonA\textsuperscript{(c)}[nonA\textsuperscript{vir}] lines when compared to the control nonA\textsuperscript{(c)}[nonA\textsuperscript{+}] is actually due to a species-specific effect connected to the presence of the nonA\textsuperscript{vir} gene. The fact that heterozygous nonA\textsuperscript{vir}/nonA\textsuperscript{+} lines are actually more polycyclic than their hemizygous nonA\textsuperscript{(c)}[nonA\textsuperscript{vir}] counterparts could be similarly explained invoking some sort of dominant effect linked to the nonA\textsuperscript{vir} gene. However, the mel+2vir genotype is normal, contradicting the notion of virilis dominance.

Alternatively, the polycyclicity of nonA\textsuperscript{(c)}[nonA\textsuperscript{vir}] lines could be explained by assuming that the nonA\textsuperscript{vir} gene is not giving complete rescue, and the results obtained with nonA\textsuperscript{vir}/nonA\textsuperscript{+} lines could be due to a dosage effect. The higher polycyclicity of 2mel could also be interpreted by invoking a dosage effect. However no dosage effect can be seen in the case of 3mel and mel+2vir.

CPP distributions.

For each song recorded the proportion of pulses containing 1, 1.5, 2, 2.5, 3,
3.5, etc.. cycles was estimated. The average proportions per genotype were then calculated. Frequency histograms showing the average proportion of pulses with 1, 1.5, 2, 2.5, and so on, cycles in each genotype are presented in figure 7.4. On the x-axis the maximum number of cycles considered was 10, however it has to be noted that a small proportion of pulses containing more than 10 cycles could be found in both diss and vir (in some diss songs pulses with more than 30 cycles were found). To see how these CPP distributions differed between genotypes, three parameters were taken into consideration: the mode of the distribution, the skewness and the kurtosis.

The mode of each song was computed and a Kruskal-Wallis non parametric ANOVA was performed on these data. In order to reduce the number of independent (grouping) variables (which in this case are the different genotypes), all the hemizygous nonA\(^{+}\) [nonAvir] genotypes were grouped together as vir/d, and all the heterozygous nonAvir/nonA\(^{+}\) as vir/m. The result of the test was highly significant (\(\chi^2 = 88.504; \text{df}= 7; p<< 0.001\)). However, when diss and vir were excluded from the analysis, the test was non-significant (\(\chi^2 = 8.663; \text{df}= 5; p= 0.123\)). A Kolmogorov-Smirnov two sample test performed on each pair of genotypes, confirmed that the only line that differs from all the others is vir. From figure 7.4 it is clear, in fact, that the mode of most genotypes, including diss, is 2. The fact that diss has a mode of 2 should not be surprising considering that in this mutant most of the pulses (all the pulses of shorter trains and all the pulses at the beginning of long trains) are still wild-type.

The skew values of the CPP distributions in each song were also analyzed using a Kruskal-Wallis ANOVA. Again, the number of independent variables was reduced by grouping all the hemizygous nonA\(^{+}\) [nonAvir] as vir/d and all the heterozygous nonAvir/nonA\(^{+}\) as vir/m. The test gave a significant result only when data from diss and vir were included in the analysis (\(\chi^2 = 22.914; \text{df}= 7; p= 0.0018\)). A Kolmogorov-Smirnov test conducted on pairs of genotypes, showed that diss was the genotype that differed from all the others. No significant difference was found between vir and any other transformant line.

The same kind of analysis was performed on the kurtosis values of each song CPP distribution. This time, the Kruskal-Wallis ANOVA was significant when the
Figure 7.4. CPP distributions in each genotype.
values of \( \text{diss} \) and \( \text{vir} \) were included (\( \chi^2 = 33.157; \text{df}=7; p=0.0000 \)), and when they were excluded (\( \chi^2 = 18.529; \text{df}=5; p=0.0024 \)). In addition significance was just missed when comparing transformants carrying one, two or three copies of \( \text{nonA}^+ \) \( (235R11/d, 2\text{mel} \) and \( 3\text{mel}; \chi^2 = 5.6; \text{df}=2; p=0.0608) \). When comparisons were made using the Kolmogorov-Smirnov test (in appendix 7.4), it was found that the \( \text{vir/d} \) genotype group (hemizygous \( \text{nonA}^+ \) [\( \text{nonAv ir} \) lines] differed significantly from the control line \( 235R11/d \) \( (p<0.05) \). However when tests were performed on each individual \( \text{nonA}^+ \) [\( \text{nonAv ir} \) line] against the control \( 235R11/d \) line, only \( 297-6/d \) gave a significant difference \( (p<0.05) \). The \( \text{vir/d} \) group was significantly different from the mutant \( \text{diss} \) \( (p<0.001) \), but did not differ from \( \text{vir} \), and this was confirmed in all individual comparisons with the Kolmogorov-Smirnov test (see appendix 7.4).

On the contrary, the \( \text{vir/m} \) genotype group (heterozygous \( \text{nonA}^{\text{vir}} \) [\( \text{nonA}^+ \) lines] did not differ from the control line \( 235R11/d \) (in individual comparisons only the line \( 135/m \) differed significantly from the control, \( p<0.05 \)) but did differ from either \( \text{diss} \), \( p<0.001 \), with all individual comparisons significant) and \( \text{vir} \), \( p<0.05 \), with two individual comparisons significantly different, \( 112/m \) and \( 113/m \). The same was found for \( 3\text{mel} \) and \( \text{mel}+2\text{vir} \) which differed from \( \text{diss} \) \( (p<0.001) \) and \( \text{vir} \), \( p<0.001 \) but not from the control. Interestingly \( 2\text{mel} \) differed from \( \text{diss} \), \( p<0.001 \) but not from either the control \( 235R11/d \) or the \( \text{vir} \) genotype (see appendix 7.4).

In conclusion, the mode and the skew of the CPP distributions in different genotypes turned out not to be very informative. However when kurtosis was considered the results revealed that hemizygous \( \text{nonA}^{\text{vir}} \) [\( \text{nonAv ir} \) lines] had CPP distributions more similar to those found in \( \text{vir} \) than those of the control line. Moreover this is unlikely to be due to incomplete rescue since hemizygous \( \text{nonA}^{\text{vir}} \) [\( \text{nonAv ir} \) lines] also differed from the mutant \( \text{diss} \). In addition, \( \text{vir/d} \) differed from the control \( 235R11/d \) as a group, whereas \( \text{vir/m} \) did not. This is probably the first evidence indicating that the \( \text{nonA}^{\text{vir}} \) gene could be carrying species-specific information. This information however does not appear to be dominant since heterozygous \( \text{nonA}^{\text{vir}} \) [\( \text{nonA}^+ \) (as a group) and \( \text{mel}+2\text{vir} \) lines differ from \( \text{vir} \), but do not differ from the \( 235R11/d \) control. Finally \( 2\text{mel} \) shows results which fall between the \( 235R11/d \) control and \( \text{vir} \). This could
possibly be due to some dosage effect linked to the *mel nonA* gene. Note however that lines carrying three copies of *mel nonA* (3*mel*) do not show this effect.

bPF.

For each song analyzed, a PF (pulse frequency) least square regression was performed as described in chapter 2 (see figure 2.2). Table 7.3 shows the number of songs, out of ten per genotype (out of six for 674/m), which had a significant PF regression gradient (bPF).

**Table 7.3.** bPF (pulse frequency gradient) for different genotypes. N= number of significant bPFs out of 10 songs (* out of 6 songs). + = significant positive bPFs. — = significant negative bPFs. The mean bPF value per genotype is also presented.

<table>
<thead>
<tr>
<th>GENOTYPE</th>
<th>N</th>
<th>+</th>
<th>−</th>
<th>Mean bPF</th>
</tr>
</thead>
<tbody>
<tr>
<td>diss</td>
<td>5</td>
<td>2</td>
<td>3</td>
<td>0.155</td>
</tr>
<tr>
<td>235R11/d</td>
<td>9</td>
<td>0</td>
<td>9</td>
<td>-4.107</td>
</tr>
<tr>
<td>113/d</td>
<td>10</td>
<td>0</td>
<td>10</td>
<td>-3.613</td>
</tr>
<tr>
<td>67-4/d</td>
<td>10</td>
<td>0</td>
<td>10</td>
<td>-4.104</td>
</tr>
<tr>
<td>168-8/d</td>
<td>8</td>
<td>0</td>
<td>8</td>
<td>-4.083</td>
</tr>
<tr>
<td>135/d</td>
<td>10</td>
<td>0</td>
<td>10</td>
<td>-5.041</td>
</tr>
<tr>
<td>97/d</td>
<td>8</td>
<td>0</td>
<td>8</td>
<td>-5.851</td>
</tr>
<tr>
<td>191/d</td>
<td>9</td>
<td>0</td>
<td>9</td>
<td>-6.841</td>
</tr>
<tr>
<td>297-6/d</td>
<td>10</td>
<td>0</td>
<td>10</td>
<td>-6.572</td>
</tr>
<tr>
<td>112/m</td>
<td>9</td>
<td>0</td>
<td>9</td>
<td>-2.781</td>
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<td>10</td>
<td>0</td>
<td>10</td>
<td>-3.464</td>
</tr>
<tr>
<td>67-4/m*</td>
<td>6</td>
<td>0</td>
<td>6</td>
<td>-1.530</td>
</tr>
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<td>0</td>
<td>9</td>
<td>-2.293</td>
</tr>
<tr>
<td>135/m</td>
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<td>0</td>
<td>10</td>
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<td>8</td>
<td>-2.988</td>
</tr>
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<td>9</td>
<td>-3.336</td>
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<tr>
<td>mel+2vir</td>
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<td>0</td>
<td>10</td>
<td>-4.725</td>
</tr>
<tr>
<td>vir</td>
<td>1</td>
<td>0</td>
<td>1</td>
<td>-0.630</td>
</tr>
</tbody>
</table>

From table 7.3 it is clear that in all the transformant lines (from line 235R11/d to line *mel+vir*) almost all the songs analyzed had a significant negative bPF. In other
words, in all the transformant lines analyzed there was a trend for the pulses at the beginning of each train to have a higher frequency than those towards the end. *diss* and *vir*, on the contrary, showed a very different PF regression pattern: only 5 *diss* songs had a significant bPF, 2 of which were positive. Only one *vir* song had a significant negative PF gradient. This means that in these two genotypes the frequencies of pulses at the beginning of a pulse train are not very different from the frequencies of pulses at the end. The average bPF values per each genotype were also calculated and are shown in table 7.3. These values plus standard error and standard deviation are given in figure 7.5.

**Figure 7.5.** bPF values.

![Graph showing bPF values for different genotypes](image)

One-way ANOVA was employed on the bPF values of each song analyzed. The result was highly significant (F= 8.066; df = 18, 167; p<< 0.001). When bPF values of *diss* and *vir* were removed from the analysis, the ANOVA still gave a highly significant F ratio (F= 4.773; df = 16, 149; p<< 0.001). From figure 7.5 it appears that *diss* and *vir* have the highest bPF values of all the genotypes analyzed. *A posteriori* Newman-Keuls test (see appendix 7.5) confirms that these two lines significantly differ from the control line 235RII/d and from most of the other transformant lines, while they do not differ from each other.
The mean bPF values in hemizygous $nonA^{c\{nonAvir\}}$ lines were found to be either similar or more negative than those of the control line $235R11/d$ (see figure 7.5). A planned comparison showed that the group including all the $nonA^{c\{nonAvir\}}$ lines does not significantly differ from the control line ($F= 2.465; \text{df} = 1, 167; p= 0.118$). Newman-Keuls $a \text{ posteriori}$ comparisons revealed that only lines $191/d$ and $297-6/d$ have significantly lower bPF values than line $235R11/d$.

Heterozygous $nonA^{vir}/nonA^+$ lines have bPF values close, or slightly higher than those of the control $235R11/d$ (figure 7.5). A planned comparison between these lines and line $235R11/d$ gave a non-significant $F$ ratio ($F= 3.35; \text{df} = 1, 167; p= 0.07$). According to the Newman-Keuls test (see appendix 7.5), no heterozygous $nonA^{vir}/nonA^+$ line differs from the control. However most of these lines do not differ from $\text{vir}$ either (except $113/m$ and $135/m$). Line $67-4/m$ is also not significantly different from $\text{dis}$.

A planned comparison between hemizygous $nonA^{c\{nonAvir\}}$ and heterozygous $nonA^{vir}/nonA^+$ showed that these two groups are significantly different ($F= 41.083; \text{df} = 1, 167; p< 0.001$). From figure 7.5 it appears that each hemizygous $nonA^{c\{nonAvir\}}$ line displays lower bPF values than its heterozygous $nonA^{vir}/nonA^+$ counterpart. Finally lines $2mel, 3mel$ and $mel+2vir$ do not differ from the control $235R11/d$.

PF.

As discussed above, pulse frequency is one of the two song characters that seem to generally correlate (in this case negatively) with the total amount of song produced by a fly (see table 7.1). Therefore for analysis, it is necessary to take into consideration the amount of real song recorded for each fly. Consequently, for each song, the (PS+SS) value was used as the covariate for PF, which itself was calculated at $x= 1$ from the regression function $y=bx+c$ (PF1). Analysis of covariance revealed a highly significant $F$ ratio ($F= 18.793; \text{df} = 18, 166; p= 0.00$). Corrected PF1 averages were produced for each genotype and their values are graphically plotted in figure 7.6.
Figure 7.6. Adjusted PF1 average values per genotype.

Again, it is obvious that the two genotypes that stand out are *diss* with much lower PF1 values than the rest, and *vir* which has much higher PF1s than the other lines. All the hemizygous *nonA*([*nonA*]<sup>vir</sup>) lines, except 113/d, show lower PF1 values than the control line 235R11/d. However Newman-Keuls comparisons reveal that only lines 67-4/d and 168-8/d significantly differ from 235R11/d. These two lines are also significantly different from *diss*, as are the rest of the *nonA*([*nonA*]<sup>vir</sup>) lines.

In figure 7.6 it is clear how the heterozygous *nonA*<sup>vir</sup>/non-*A*<sup>+</sup> lines have in general even lower PF1 values than their hemizygous counterparts, although the only significant difference is between 97/d and 97/m. According to Newman-Keuls tests, four out of the six heterozygous *nonA*<sup>vir</sup>/non-*A*<sup>+</sup> lines (67-4/m, 168-8/m 135/m and 97/m) significantly differ (p<0.05) from the control 235R11/d, even though they also differ from *diss*. Lines 2mel 3mel and mel+2vir had average PF1 values similar to the control line 235R11/d and in fact they do not significantly differ from it.

To conclude, the *nonA*<sup>vir</sup> gene, either on a *nonA*<sup>−</sup> or in a *non-A*<sup>+</sup> background, does not seem to have any species-specific effect on the pulse frequency of transformant flies. Furthermore the mutant *diss* pulse frequency appear to be rescued in *nonA*<sup>−</sup>[*nonA*<sup>vir</sup>] lines. In fact, while *vir* PF values are on average quite high, all the
transformants carrying one or two copies of the nonAvir gene display either similar or lower PFs than the control line transformed with non-A\(^+\) (235R11). Two hemizygous nonAvir(nonAvir) lines have PF values that seem to be intermediate between the control and the mutant diss. This could be explained as a result of non-perfect rescue of this song character. However it is more difficult to explain why four of the heterozygous nonAvir/nonA\(^+\) also display low PF values.

SSP.

As previously described, the courtship song of D.melanogaster is not exclusively composed of trains of pulses. A second song component consisting of ~160 HZ hums (sine or hum song) is generally present interspersed within the pulse song. D.virilis males, on the contrary, produce courtship songs containing only trains of polycyclic pulses. It seemed therefore interesting to study the amount of sine song produced by transformant males carrying one or two copies of the nonAvir gene in order to establish whether this gene had any possible effect in repressing sine song production.

The proportion of sine song produced during coutship by each male was calculated as the amount of sine song over the total amount of song (PS+SS) (see chapter 2). All the males of each genotype, including males of the diss line, produced at least one or two bursts of sine song. The only exception were two males of the mel+2vir line whose song was completely devoid of sine song.

SSP, as discussed above, seems to be almost always positively correlated with the total amount of song recorded for each fly (PS+SS). This was the case in all the genotypes analyzed except in line 67-4/d and 135/d for which the correlation was negative although not significant (see table 7.1). It seemed opportune therefore to take into account the PS+SS value of each song when analyzing SSP values. Analysis of covariance (where SSP was the dependent variable and PS+SS the covariate) gave a highly significant F ratio (F= 12.258; df genotype= 17, 157; p<< 0.001). The adjusted
SSP means per each genotype were calculated and are graphically represented in figure 7.7.

**Figure 7.7.** Adjusted SSP average values per genotype.

From figure 7.7, large variation of SSP averages can be noticed within each of the two groups constituted by hemizygous \( nonA^c[nonA^v] \) and heterozygous \( nonA^v/nonA^c \) lines. In general \( nonA^c[nonA^v] \) lines seemed to have higher SSPs than \( nonA^v/nonA^c \). Only line \( 135/d \) had significantly lower SSP values than the control \( 235R11/d \), according to a Newman-Keuls test, while line \( 97/d \) had actually significantly higher SSPs. All the other \( nonA^c[nonA^v] \) lines did not differ from the control.

Lines \( 112/m \) and \( 113/m \) had the lowest SSP values among all the genotypes analyzed. However the four other \( nonA^v/nonA^c \) lines had much higher values which did not significantly differ from the control (\( 235R11/d \)). When comparing each hemizygous \( nonA^c[nonA^v] \) line with its heterozygous \( nonA^v/nonA^c \) counterpart, no coherent pattern can be noticed. Finally, \( 2mel \) and \( 3mel \) had similar SSP values to those of the control \( 235R11/d \), while \( diss \) and \( mel+2vir \) had significantly lower SSPs.
In conclusion, there is no argument to support the idea that the nonA\textsuperscript{vir} gene causes suppression of sine song. Neither \textit{nonA\textsuperscript{vir}}
\textit{[nonA\textsuperscript{vir}] or nonA\textsuperscript{vir}/nonA\textsuperscript{+}} lines show a consistent pattern of SSP values, and adding multiple doses of the \textit{mel} or \textit{vir nonA} gene does not help to clarify the picture.

SSF.

Wild type \textit{D.melanogaster} sine song frequencies usually range between 100 and 200 Hz, with median values varying between 120 and 160 Hz (Wheeler \textit{et al.}, 1989). SSF values were calculated for each song as the average frequency of ten sine song bouts, as described in chapter 2. The average SSF values found in each genotype are shown in figure 7.8.

**Figure 7.8.** Average SSF values for each genotype.

One-way ANOVA performed on the SSF values of each song analyzed gave a highly significant F ratio (F=10.882; df = 17, 157; p<< 0.001). From figure 7.8 it is clear how \textit{diss} seems to be normal in respect to its SSF values when compared to the
control 235R11/d. The Newman-Keuls *a posteriori* test confirmed that no significant difference exists between the SSFs of *diss* and those of 235R11/d (see appendix 7.6).

Hemizygous *nonA*[^]*[nonA*vir*]* lines had SSF values either similar or higher than those of the control 235R11/d. A planned comparison between the group including all the *nonA*[^][nonA*vir*] lines and the control gave a significant F ratio (F= 6.320; df= 1, 157; p= 0.013). In particular Newman-Keuls tests showed that 168-8/d and 135/d display significantly higher SSFs than 235R11/d (see appendix 7.6).

Heterozygous *nonA*[^][nonA*vir*] lines show a complicated pattern since four of them have higher SSF values than the control (67-4/m and 97/m have significantly higher SSFs than 235R11/d, see Newman-Keuls in appendix 7.6) while other two (112/m and 113/m) have significantly lower values (see appendix 7.6). Not surprisingly a planned comparison between line 235R11/d and the *nonA*[^][nonA*vir*]+ lines was non-significant (F= 0.745; df= 1, 157; p= 0.389).

A planned comparison between hemizygous *nonA*[^][nonA*vir*] lines and heterozygous *nonA*[^][nonA*vir*]/nonA was significant (F= 9.316; df= 1, 157; p= 0.003). However, comparing each hemizygous *nonA*[^][nonA*vir*] line with its heterozygous *nonA*[^][nonA*vir*]/nonA+ counterpart (for example 113/d with 113/m) does not reveal any coherent pattern: in some cases the hemizygous line has higher value than the heterozygous (this is the case of lines 113, 168-8 and 135), in others the situation is reversed (compare 67-4/d with 67-4/m and 97/d with 97/m in figure 7.8). Finally, line 2mel has higher SSFs than the control, even though the difference is not significant, while 3mel and mel+2vir have values similar to 235R11/d.

In summary, it seems that the *nonA*[^][nonA*vir*] gene plays no role in affecting the sine song frequency. This is not altogether surprising considered that it has been shown above that the *nonA*[^][nonA*vir*] transgene does not affect the frequencies of single pulses either. *D. virilis* flies produce very high frequency sounds but this characteristic is obviously not determined by the *nonA* gene.
IPI.

The mean IPI values were calculated for each song as described in chapter 2. One way ANOVA performed on these data gave a highly significant F ratio ($F=64.492; df=18, 167; p<0.001$). The result of the ANOVA was still significant even after excluding the data from $vir$ and $diss$ ($F=11.295; df=16, 149; p<0.001$). Figure 7.9 graphically illustrates the mean, standard error and standard deviation of the IPI values of each genotype.

**Figure 7.9.** Mean IPI values of each genotype.

From figure 7.9 it is clear that $diss$ displays the highest IPI values of all the genotypes analyzed. These values differ significantly from the IPI values shown by the control line $235R11/d$, as confirmed by the *a posteriori* test (see appendix 7.7). As expected from previous analysis of *virilis* songs (Hoikkala *et al.*, 1982), the mean IPIs of this species are very short.

The mean IPI values of hemizygous $nonA^{vir}$*$[nonA^{vir}]$ lines are either similar or lower than those of the control $235R11/d$. A planned comparison between line $235R11/d$ and all the hemizygous lines $nonA^{vir}$*$[nonA^{vir}]$ (from line $113/d$ to line $297-$
6/d, see figure 7.9) showed that these two groups are indeed significantly different (F= 5.298; df= 1, 167; p= 0.023). In particular, a Newman-Keuls test revealed that the IPI values of lines 168-8/d and 135/d are significantly lower than those of the control 235R11/d (see appendix 7.7).

From figure 7.9 the mean IPIs of heterozygous nonA\( ^{\text{vir}}/\text{nonA}^+ \) lines appear to be either similar or higher than those of line 235R11/d. A planned comparison showed that this group of genotypes significantly differs from the control line 235R11/d (F= 6.763, df= 1, 167; p= 0.010). A Newman-Keuls test pointed out that the mean IPI values of lines 113/m and 112/m are significantly higher than those of the control 235R11/d. These two genotypes have IPI values that do not differ from those of diss. Also lines 135/m and 97/m do not significantly differ from diss, but they also do not significantly differ from the control 235R11/d (see appendix 7.7).

Another planned comparison between hemizygous nonA\( ^{\text{c}^1}\)[nonA\( ^{\text{vir}} \)] and heterozygous nonA\( ^{\text{vir}}/\text{nonA}^+ \) lines revealed that these two groups are significantly different from each other (F= 83.368; df= 1, 167; p<< 0.001). Note from figure 7.9 that each hemizygous nonA\( ^{\text{c}^1}\)[nonA\( ^{\text{vir}} \)] line always has a lower IPI value than its heterozygous counterpart. This was confirmed by Newman-Keuls tests (see appendix 7.7).

Finally, lines 2mel, 3mel, and mel+2vir have IPIs values higher than the control 235R11/d. All three lines do not significantly differ from diss, however, only line 3mel has significantly higher IPI values than the control line (see Newman-Keuls in appendix 7.7).

Considering that vir songs have very short mean IPIs, the fact the songs of flies carrying the nonA\( ^{\text{vir}} \) gene on a nonA\( ^{\text{c}^1} \) background tend to have lower IPI values than those of the control line is interesting and unexpected. However this effect on the IPIs is quite marginal, since, as seen above, only two nonA\( ^{\text{c}^1}\)[nonA\( ^{\text{vir}} \)] lines differ significantly from the control. Dosage effects could explain why some nonA\( ^{\text{vir}}/\text{nonA}^+ \) heterozygous lines, and lines with more than one copy of nonA\( ^+ \) (for example 3mel) have higher IPIs than the control. In fact adding copies of nonA\( ^+ \) (either virilis or mel)
increases IPIs, yet heterozygotes (\textit{nonA}^{vir}/\textit{nonA}^+, or \textit{mel}+\textit{2vir}) have in general lower IPIs than their hemizygous 2-dose or 3-dose \textit{mel} counterparts.

The fact that \textit{diss} has such long IPIs could however be the result of an artifact due to the high CPP and low frequencies of the pulses in this line. IPIs were here measured as the average distance between the maximum peak of the intrapulse oscillation of one pulse to the maximum peak of the next pulse. Thus low frequency, polycyclic pulses will have longer IPIs because the distance from the maximum peak of one such pulses to the peak of the next is actually greater than the same distance measured between high frequency, monocyclic pulses. In order to overcome this possible problem, I decided to correct each line’s IPI values by taking into account their respective CPP (at x=0), bCPP, PF (at x=0) and bPF values. An analysis of covariance was performed in which CPP, bCPP, PF and bPF were the covariates of the IPI values. The resulting F ratio was highly significant (F= 22.083; df= 18, 163; p<< 0.001). Figure 7.10 shows the adjusted IPI values for each genotype.

\textbf{Figure 7.10.} Adjusted IPI values.

![Adjusted IPI values](image)

Note that after correction the IPI values of \textit{diss} become very similar to those of the control line \textit{235R11/d}. Hemizygous \textit{nonA}^{vir}/[\textit{nonA}^{vir}] lines still have similar or
lower values than those of the control. Newman-Keuls *a posteriori* comparisons showed that the values of 67-4/d, 168-8/d, and 135/d are significantly lower than those of line 235R11/d. After correction the IPIs of lines 112/m and 113/m are still significantly higher than those of the control, while those of lines 67-4/m and 168-8/m are actually significantly lower. The adjusted IPI values of 2mel, 3mel and mel+2vir are not significantly different from those of the control line 235R11/d.

In summary, the most interesting result obtained after correcting the IPIs values for their respective CPP, bCPP, PF and bPF values, is that *diss* becomes indistinguishable from the control line. On the contrary, hemizygous nonA[^vir][nonA[^+]] lines retain generally lower values than line 235R11/d. After correction, the situation of the heterozygous nonA[^vir]/nonA[^+] lines changes slightly, since two of these lines now show significantly lower values than the control (before correction all nonA[^vir]/nonA[^+] lines had similar or higher values than 235R11/d, see figure 7.9). Also the results obtained with 2mel, 3mel and mel+2vir, were changed after correction, since these lines no longer differ from the control strain.

**MULTIVARIATE ANALYSES**

So far, the analyses have examined individual characteristics and some evidence for a modest transfer of *virilis*-like song phenotypes to *melanogaster* transformants has been presented. No compelling evidence was produced that the transgenes 113, 67-4, and 168-8 were any different from the other lines. Thus loss of 3' material did not generate a clearly identifiable song phenotype. When several variables are investigated as in this case, a more comprehensive picture can often be obtained if all the phenotypes are studied simultaneously. A number of multivariate tests are available, and discriminant analysis (Anderson, 1963) was chosen as a means of detecting differences between the experimental groups. SYSTAT 7.0 software was used and I am grateful to Yuri Dubrova for running the analysis on his computer and explaining the method and results to me.
All the six song variables, CPP, bCPP, PF, bPF, SSF, and IPI, were used. SSF of course could not be used in tests which included D. virilis songs. One diss fly, which showed a negative intercept on its CPP and a correspondingly high bCPP was removed to generate a normal distribution for these characters. In practice, this makes little difference to the analysis, as diss mutants form a clearly separate group (see below).

Figure 7.11a and b show the results of the discriminant analysis on the whole sample. This first analysis placed all individuals into 5 groups: vir (D. virilis), diss, vir/d (all the nonA⁻/nonA⁺ transformants), the group named MEL which includes the control line 235RII/d (or 1mel), 2mel and 3mel, and the "hybrids" group (HYB) composed by vir/m (all the nonA⁺/nonA⁻ transformants) and mel+2vir. In figure 7.11a individual scores are plotted plus the ellipses defining the 95% confidence limits, thereby giving an idea of the overlap between the 5 groups, and in figure 7.11b the mean for each group is shown for clarity. The vir species and diss mutants form two outlying, nonoverlapping groups. Table 7.4 presents the summary of the analysis and clearly indicates that each behavioural characteristic contributes equally highly to the differentiation of these 5 groups. An extremely low probability of misclassification of groups (the F-statistic estimates the probability of Wilk’s lambda value occurring by chance) is observed. This analysis reveals that about 87% of the total variance of all traits corresponds to the differences between vir and the rest of the groups (factor 1), whereas approximately 9% of the total variance appears due to the differences between diss and the others. Within this initial analysis, the transformant lines form one, largely overlapping group.

Table 7.4. Results of the discriminant analysis for the whole sample.

<table>
<thead>
<tr>
<th>Traits</th>
<th>F (4, 180)</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>IPI</td>
<td>217.04</td>
<td>&lt;&lt;0.0001</td>
</tr>
<tr>
<td>PF</td>
<td>27.15</td>
<td>&lt;&lt;0.0001</td>
</tr>
<tr>
<td>CPP</td>
<td>68.41</td>
<td>&lt;&lt;0.0001</td>
</tr>
<tr>
<td>bCPP</td>
<td>122.84</td>
<td>&lt;&lt;0.0001</td>
</tr>
<tr>
<td>bPF</td>
<td>20.74</td>
<td>&lt;&lt;0.0001</td>
</tr>
<tr>
<td>SSF</td>
<td>--</td>
<td>--</td>
</tr>
</tbody>
</table>

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Figure 7.11. Discriminant analysis of all genotypes grouped as follows: MEL = 235R11/d, 2mel, 3mel, vir/d = 113/d, 67-4/d, 168-8/d, 135/d, 97/d, 191/d, 297-6/d, HYB = 112/m, 113/m, 67-4/m, 168-8/m, 135/m, 97/m, mel+2vir. diss = nonA\textsuperscript{ass}, vir = D.virilis. A: individual scores plus the ellipses defining the 95% confidence limits. B: mean score for each group. Factor 1 accounts for 87.6% of the total variability, while factor 2 accounts for 9.5%.
A second series of discriminant analyses were performed within each of the three transformant groups (MEL, vir/d, and HYB), where individuals were placed within their transformant line. A highly significant differentiation was observed between melanogaster transformants carrying one (235R11/d or 1mel), two (2mel) and three (3mel) doses of D. melanogaster nonA+ (see figure 7.12a and b). There is little overlap between the three groups (figure 7.12a). Factor 1 accounted for about 67% of the variance and was due to the difference between having one (1mel) and two copies of nonA+ (2mel) whereas factor 2 accounts for about one-third of the variance and is caused by differences between three dose nonA+ flies (3mel) and the others. There is no simple dosage effect within the MEL group, as 3mel and 2mel have similar scores on factor 1, as do 1mel (235R11/d) and 2mel on factor 2. Appendix 7.8 shows the summary statistics for this analysis, with bPF failing to contribute significantly to the discrimination.

The vir/d transformants (all the nonA[nonA vir] lines) gave a more complex picture (figure 7.13a and b). Fig 7.13a shows significant overlap between most groups, but 113/d and its derivatives 168-8/d and 67-4/d do not overlap, and form the outliers. This is seen more clearly in figure 7.13b where the means for each line are presented. The different transformant lines cluster in two main sub-groups. 168-8/d and 67-4/d lie together in negative space for both factors, whereas all the other transformants except 113/d, (from which the former two were derived by ‘jumps’, see chapter 4) lie in positive space. Thus all the nonA vir insertions which are known to be intact, appear to cluster away from those which may be missing some regulatory material (see chapter 4). The bCCP phenotype does not contribute significantly to this discrimination (see Appendix 7.9). This pattern is repeated with the vir/m hybrids, again the 67-4/m and 168-8/m transformants clustering away from the other groups (see figure 7.14a and b and appendix 7.10).
Figure 7.12. Discriminant analysis of the MEL group (1mel, 2mel, 3mel)  
A: the individual scores for each genotype are indicated plus the ellipses defining the 95% confidence limits.  
B: mean score for each genotype. Factor 1 accounts for 67.5% of the total variability, while factor 2 accounts for 32.5%.
Figure 7.13. Discriminant analysis of the vir/d group (all the nonA\[^{+}\] [nonA\[^{+}\]] transformants). A: scores of the individuals of each genotype plus the ellipses defining the 95% confidence limits. B: mean for each transformant line. Factor 1 accounts for 69.5% of the total variability, while factor 2 accounts for 15.4%.
Figure 7.14 Discriminant analysis of the hybrids (HYB) group (including all the nonA⁺/nonA⁺ transformants and mel+2vir). A: scores of the individuals of each genotype plus the ellipses defining the 95% confidence limits. B: mean score for each line. Factor 1 accounts for 86.9% of the total variability, while factor 2 accounts for 10.6%. 
The final discriminant analysis puts all the individual lines into their respective groups, and maintains the different groups in terms of gene dosage. Thus \(1\text{mel}, 2\text{mel}\) and \(3\text{mel}\) are treated separately. Line \(\text{mel}+2\text{vir}\), and all the lines in which the insert is not intact (113, 67-4 and 168-8, see chapter 4) were excluded from the analysis. 5 groups were generated, \(1\text{mel}, 2\text{mel}\) and \(3\text{mel}\), the \(\text{vir/d}\) group which included the 4 intact transgenes (135, 97, 191, and 297-6), and the hybrid group \(\text{vir/m}\), again containing only intact \(\text{virilis}\) transgenes (135, 97 and 112). Figure 7.15a and b reveal a clear pattern, with the \(\text{vir/d}\) transformants clustering away from the others on factor 1, and accounting for about 78% of the variance. Differences between \(1\text{mel}\) and \(2\text{mel}\) account for 14% of the variance on factor 2. The summary statistics are presented in appendix 7.11. CPP just fails to reach significance in contributing to the discrimination, but all other factors, including SSF, contribute significantly. Thus a separation on factor 1 between \(\text{vir/d}\) transformants and each of the three \(\text{melanogaster}\) dosage groups (\(1\text{mel}, 2\text{mel}\) and \(3\text{mel}\)) is observed. The hybrids (\(\text{vir/m}\)) cluster with the \(\text{melanogaster}\) groups on factor 1, while on factor 2 they display differences from groups \(1\text{mel}\) and \(2\text{mel}\), but not from \(3\text{mel}\), or \(\text{vir/d}\), thereby revealing a recessive phenotype with respect to \(\text{nonA}^\text{vir}\) on factor 1, but a dominant one for factor 2.

In conclusion, the multivariate analyses have shown reasonably clearly that the \(\text{vir/d}\) transformants can be discriminated from the other groups by the simultaneous comparison of all the song phenotypes.

**DISCUSSION**

Table 7.5 shows a synopsis of the results presented. For each transformant line the trend shown by each song character analyzed (only informative characters are included in the table) has been classified as dissonance-like (D), \(\text{virilis}\)-like (V), or \(\text{melanogaster}\)-like (M). The classification was based on the very conservative \textit{a posteriori} tests described in the results section. Any character which showed a trend
Figure 7.15. Discriminant analysis of all the genotypes, excluding the transformant lines in which the insert is not intact (113/d, 67-4/d, 168-8/d, 113/m, 67-4/m, 168-8/m and mel+2vir; see text and chapter 4). 

A: scores of the individuals belonging to each group plus the ellipses defining the 95% confidence limit. 
B: mean of each group. Factor 1 accounts for 77.7% of the total variability, while factor 2 accounts for 14.1%.
similar, but still significantly different from, let us say, diss, was marked as D followed by an asterisk (D*). Therefore, a line which is marked as D (and no asterisk) for one character, not only shows a trend similar to diss, but also does not significantly differ from it in that character. Asterisks were also used in the same way for M and V. Note that in the last column (IPI adjusted), as diss and mel values are indistinguishable (see figure 7.10), only M or V classifications are used.

**Table 7.5.**

Trends in song characters displayed by each transformant line. M = melanogaster-like; D = dissonance (nonA_diss)-like; V = virilis-like. sign. = significant slope. distrib. = distribution.

<table>
<thead>
<tr>
<th>BCPP sign.</th>
<th>BCPP distrib.</th>
<th>CPP10 sign.</th>
<th>CPP10 distrib.</th>
<th>IPI sign.</th>
<th>IPI adjusted</th>
</tr>
</thead>
<tbody>
<tr>
<td>113/d</td>
<td>D,V*</td>
<td>M</td>
<td>M</td>
<td>M</td>
<td>M</td>
</tr>
<tr>
<td>67/d</td>
<td>D,V*</td>
<td>M</td>
<td>M,Y</td>
<td>M</td>
<td>M,V</td>
</tr>
<tr>
<td>168-8/d</td>
<td>D,V*</td>
<td>M</td>
<td>M,V</td>
<td>M</td>
<td>M,D</td>
</tr>
<tr>
<td>135/d</td>
<td>D,V*</td>
<td>M</td>
<td>M</td>
<td>M</td>
<td>M</td>
</tr>
<tr>
<td>97/d</td>
<td>D,V*</td>
<td>M</td>
<td>M</td>
<td>M</td>
<td>M</td>
</tr>
<tr>
<td>191/d</td>
<td>D,V*</td>
<td>M</td>
<td>M</td>
<td>M</td>
<td>M</td>
</tr>
<tr>
<td>297-6/d</td>
<td>D,V*</td>
<td>M</td>
<td>V</td>
<td>M</td>
<td>M</td>
</tr>
<tr>
<td>112/m</td>
<td>M</td>
<td>M</td>
<td>M</td>
<td>M</td>
<td>M,D</td>
</tr>
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<td>M</td>
<td>M,D</td>
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</tr>
<tr>
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<td>3mel</td>
<td>M</td>
<td>M</td>
<td>M</td>
<td>M</td>
<td>M,D</td>
</tr>
<tr>
<td>mel+2vir</td>
<td>M</td>
<td>M</td>
<td>M</td>
<td>M</td>
<td>M,D</td>
</tr>
</tbody>
</table>

**Poor rescue or species-specific effect?**

The most striking feature of table 7.5 is the observation that in most of the cases it was impossible to distinguish between a diss-like phenotype (D) or a vir-like (V) one. In fact the unexpected positive regression found for CPP in D.virilis complicated interpretation of the results, since the mutant, nonA_diss carries a similar
feature. Thus a positive gradient could be interpreted as either virilis-like or mutant diss behaviour.

It was clear that all the hemizygous nonA<sup>c</sup>[nonA<sup>vir</sup>] transformants had significantly more positive regressions than the control 235R11/d and the heterozygous nonA<sup>vir</sup>/nonA<sup>+</sup> lines (table 7.2 and figure 7.2b). If this was simply a reflection of poorer rescue in nonA<sup>c</sup>[nonA<sup>vir</sup>] lines, then we should expect CPP measures to show a similar pattern, i.e. higher CPP in nonA<sup>c</sup>[nonA<sup>vir</sup>]> nonA<sup>vir</sup>/nonA<sup>+</sup> > 235R11/d. Yet this was not the case: CPP values (at both x=1 and x=10) for nonA<sup>vir</sup>/nonA<sup>+</sup> lines were constantly higher than those of nonA<sup>c</sup>[nonA<sup>vir</sup>], which in turn were higher than 235R11/d. Since the CPP gradients of lines nonA<sup>vir</sup>/nonA<sup>+</sup> were definitely wild-type-like, it is difficult to explain this result as a mutant-like phenotype (possibly caused by a dosage effect). Perhaps this result reflects species-specific information conveyed by the nonA<sup>vir</sup> gene? However, if this was true, then why does line mel+2vir have completely normal bCPP and CPP values?

The CPP distributions were informative when the fourth moment (kurtosis) was examined. As a group, the nonA<sup>c</sup>[nonA<sup>vir</sup>] transformants had distributions more similar to those of virilis, and differed significantly from the mel control (235R11/d) and from the mutant diss. The group composed by nonA<sup>vir</sup>/nonA<sup>+</sup> lines instead did not differ from the control, but significantly differed from either diss or vir. These results were also partially supported when individual comparisons were made between lines. Most importantly, each transformant line differed significantly from diss, suggesting that robust rescue was achieved for this song character. Furthermore, most of the individual transformant lines carrying the nonA<sup>vir</sup> gene had distributions falling in between mel and vir (see table 7.5). Consequently, there is again some evidence for species-specific effects being transferred, and the results are difficult to place within an "incomplete rescue" scenario.

When the PF regressions were examined, all the nonA<sup>c</sup>[nonA<sup>vir</sup>] transformants behaved in a manner indistinguishable from the control line, showing good rescue for this character. When the bPF values were analyzed, some heterozygous nonA<sup>vir</sup>/nonA<sup>+</sup> lines gave values that fell between mel and vir, and in one case (67-4/m) it was not
possible to distinguish from a diss-like phenotype. Therefore, again, this could reflect some evidence for a transfer of species-specific information, but, the fact that diss behaved similarly to vir also in this song character, again created problems (at least for line 67-4/m) in distinguishing between a mutant or a virilis-like phenotype.

When PF was analyzed, after correction for song output (figure 7.6), both the nonA( )[nonA^vir] and nonA^vir/nonA^+ group lines gave values generally lower than the control 235R11/d and vir. In particular four out of the six heterozygous nonA^vir/nonA^+ lines analyzed, had PFs rather similar to (even though still statistically different from) diss. If this result was to be considered as evidence of poor rescue or mutant-like phenotype, then it is again difficult to understand why heterozygous nonA^vir/nonA^+ lines that show perfectly normal CPP gradients have actually lower PF values than the hemizygous nonA^+/[nonA^vir] lines whose CPP gradient might be considered as reflecting incomplete rescue. An alternative hypothesis to the results obtained with PF is that perhaps, in D.melanogaster, simply having more CPP predispose to lower PFs. This would explain why nonA^vir/nonA^+ lines, which have in general more CPP than nonA( )[nonA^vir] lines, also have lower PF.

SSP and SSF were not informative. My idea that nonA^vir might act as a factor repressing sine song in D.virilis was not supported. The only gene known so far to suppress SS in D.melanogaster is doublesex (dxs) (Villella and Hall, 1996).

Finally, the IPI results were quite interesting in that the hemizygous nonA( )[nonA^vir] flies had IPIs in the D.virilis direction, and generally lower than those of heterozygous nonA^vir/nonA^+ lines. Some lines of the heterozygous nonA^vir/nonA^+ group had IPIs falling in between the control and diss, and two lines had IPI values actually more similar to diss than to the control. However, the higher diss IPI may be an artifact due to the increased CPP and lower PF of these mutants. Because of the way IPIs were calculated (i.e. as the distance from the highest peak of the intrapulse oscillation of one pulse to the highest peak of the next pulse), it may be inevitable that pulses with higher CPPs and low PFs show longer IPIs. In fact, after performing a covariance analysis in which the IPI values were corrected by taking into account the
corresponding CPP, bCPP, PF and bPF values, the IPIs of diss were found to be very similar to those of the control 235R11/d.

However, even with diss it is difficult to know at what level IPI control occurs: in other words, does the actual space between two pulses stay the same even between polycyclic pulses, so that the IPIs measured as the peak to peak distance between such pulses is really longer than for monocyclic pulses? Or is it that when pulses are polycyclic the actual space between them is reduced so that the distance between the maximum peak of one pulse and the maximum peak of the next pulse remains unaltered if compared to monocyclic pulses? If the first case is true, then making a compensation for CPP and PF in the measure of IPIs is the correct thing to do. If the second case is correct, then the covariance analysis may not be meaningful.

To summarize, because of the positive bCPP values of hemizygous nonA(\textsuperscript{-}) [nonA\textsuperscript{vir}] lines, it proves very difficult to conclude whether they are completely rescued in their song phenotype, or whether they actually show any virilis song features. It must be noted however that in the PF characters they behave decisively like the control, and in a few characters resemble the vir phenotype (see their CPP distributions, and IPIs).

The existence of some species-specific information being conveyed by the nonA\textsuperscript{vir} gene can however be reasonably supported when taking into consideration the results obtained with the heterozygous nonA\textsuperscript{vir}/nonA\textsuperscript{+} lines. The fact that these lines have unusually high CPP in their songs, cannot be simply explained as a mutant phenotype caused by the males carrying one copy of nonA\textsuperscript{vir} and one endogenous copy of nonA\textsuperscript{+}. In most of the song characters analyzed, these lines are indistinguishable from the control, and even for those characters which present some inconsistencies (i.e. PF1 and IPI), alternative explanations, other than mutant effects, can be argued.

In addition, the multivariate analyses also suggest that, in the vir/d transformants, species-specific transfer of some virilis-like information may have passed into the melanogaster hosts. The multivariate analyses clearly show a differentiation between the vir/d males and the others transformants. In these analyses
all the song characters contribute significantly to the differentiation. This includes measures of CPP (CPP or bCPP) PF (PF or bPF) IPI and SSF. Again, the differences seen between the diss mutants and D.virilis are so large (Fig 7.11a and b) that it is not possible to say whether the subtle differences observed between transformant groups reflect mutant or virilis-like behaviour. However, if we imagine that the nonAvir transgene is robust enough to rescue visual defects, but that its "dosage" is not enough to ensure proper song rescue, (perhaps via regulatory region divergence), then we might expect the partial hybrids vir/m, to have a dosage that reflects more than one copy of the melanogaster gene but less than two copies. Thus the vir/m hybrids should fall in multivariate space between one and two dose melanogaster flies (1mel and 2mel). Yet, on the main factor of the discriminant analysis the hybrids lie at the same point as 1mel and 2mel (figure 7.15b). On factor 2, which accounts for only 14% of the variance, they lie between 1mel and 2mel but are also at the same level as 3mel. It is therefore difficult to argue that the vir/d phenotype is mutant due to less than optimal dosage, particularly as no clear picture on dosage emerges even from comparing 1mel, 2mel, and 3mel transformants (see figure 712b).

Of additional interest was that the discriminant analysis also differentiated between the lines containing insert 113 (113/d and 113/m), and those containing its two 'jumped' derivatives 67-4 and 168-8 (67-4/d and 67-4/m, 168-8/d and 168-8/m, see figures 7.13a,b and 7.14a,b). While the 113 insert may be missing some of the flanking DNA for nonAvir, the multivariate analysis argues that perhaps further regulatory material has 'gone' in inserts 67-4 and 168-8, as the transformants carrying them are clearly outliers.

Line mel+2vir is something of an enigma, since it is consistently indistinguishable from the control line 235R11/d in all the individual song characters analyzed. When the multivariate analysis is performed, this line clusters with other heterozygous lines carrying only one dose of nonAvir and mel nonA+ (figure 7.14b), as if the extra copy of the nonAvir gene made no difference.
Dosage effect?

As discussed above, having one dose of $\text{nonA}^{vir}$ and one of $\text{mel nonA}^{+}$ does not seem to cause a song phenotype that could be reasonably classified as mutant. However, the results obtained from flies carrying two copies of $\text{nonA}^{+}$ ($2\text{mel}$) are somehow puzzling. Even though their songs’ bCPP values are not statistically different from those of the control $235R11/d$, most of the $2\text{mel}$ songs analyzed had significantly positive CPP regressions. Moreover $2\text{mel}$ had significantly higher CPP values than the control, and the CPP distribution in this line was intermediate between that of the control line and $D.virilis$. $2\text{mel}$ IPI values were also quite high (between those of $235R11/d$ and those of $\text{diss}$), however after correction with covariance they actually become very similar to those of the control. These results could only be explained by assuming that $2\text{mel}$ flies are slightly mutant in their singing behaviour. Maybe having two doses of the $\text{nonA}$ gene in males somehow affects its proper functioning. However if this is true, then why is it that flies carrying one copy of $\text{nonA}^{vir}$ and one of $\text{mel nonA}^{+}$, 3 copies of $\text{nonA}^{+}$ ($3\text{mel}$), or one copy of $\text{mel nonA}^{+}$ and two of $\text{nonA}^{vir}$ do not seem to be similarly affected? As already mentioned, the multivariate analyses also showed that it is not possible to classify the results obtained with $1\text{mel}$, $2\text{mel}$, $3\text{mel}$, and the hybrids with a simple dosage effect explanation.

To conclude, this exhaustive analysis of the song data does bring a final message: clearly, no striking species-specific effect is carried over with the $\text{nonA}^{vir}$ gene, as was the case with the $\text{per}$ gene of $D.simulans$ (Wheeler et al., 1991) and $D.pseudoobscura$ (Petersen et al., 1988). However it may be that $\text{nonA}$ does convey some relevant species information, but only a small proportion of the interspecific variation is transferred.
CHAPTER 8

DISCUSSION
The aim of this thesis was to investigate whether the nonA gene carries species-specific information concerning the male's lovesong. The success of similar studies using D.simulans and D.pseudoobscura per transgenes to examine lovesong and circadian locomotor patterns was a major motivation for my experiments (Petersen et al., 1988; Wheeler et al., 1991). The courtship song of Drosophila is undoubtedly important for recognition of conspecific mates (Kyriacou and Hall, 1980; 1982; Cowling and Burnet, 1981), and therefore is a character which could be directly involved in speciation (Coyne, 1992). In addition, locomotor activity patterns could also be viewed as an important factor in sexual isolation: if one species is active only in the morning while the other is active only at night, individuals from the two different species are temporally isolated and may actually never come into contact with each other even if they cohabit the same niche. This would ensure that no interspecific matings occur, thereby reducing the risk of producing wasteful inviable or sterile hybrid offspring. By controlling directly species-specific characteristics of courtship song and locomotor activity in flies, the per gene can contribute directly to the speciation process, and has therefore been named a "speciation gene" by Coyne (1992).

Superficially, nonA resembles the per gene, in that it also controls characteristic elements of the lovesong. Consequently, could nonA be in any way responsible for the differences seen in the lovesong of D.melanogaster and that of another species? To address this question it was essential to find a species that differed from D.melanogaster in song characters that might be controlled by nonA. The best candidate was D.virilis, whose song is very different from that of D.melanogaster, and whose polycyclic pulses actually resembled those found in the mutant nonA^diss (see figure 7.1). Other Drosophila species songs that have been well characterized seemed too similar to the D.melanogaster song to be considered useful for this study (Cowling and Burnet, 1981). Furthermore, in the case of D.melanogaster and D.simulans it had been shown that characters such as the number of cycles per pulse or the interpulse interval are actually controlled by autosomal genes (Wheeler et al., 1988). Studies conducted on the species-specific differences between D.virilis and other species from the montana phylad instead suggested that lovesong characters such as IPI, CPP and
pulse train length might be controlled by X-linked genes (Hoikkala and Lumme, 1987). In the case of the differences between the song of *D. virilis* and *D. littoralis*, the involvement of sex-linked genes was actually demonstrated (Hoikkala and Lumme, 1987). Could *nonA* (which in *D. virilis* maps on the X as in *D. melanogaster*, Hoikkala, personal communication) be one of these genes? The fact that apparently no gene on the X chromosome controls lovesong characters such as CPP or IPIs between the *D. melanogaster* and *D. simulans* species is not contradictory, as it does not mean that *nonA* cannot be controlling these song differences between *D. virilis* and *D. melanogaster*. It is possible that in the lineage leading to *D. virilis*, changes which determine lovesong traits accumulated in the *nonA* gene. In the 1-2 million years since the divergence of *D. melanogaster* from *D. simulans*, clearly no X-chromosome gene (including *nonA*) which could affect these song characters, have had time to diverge. In fact, initial work I performed on the *D. simulans nonA* gene showed that its sequence is almost completely identical to that of *D. melanogaster*.

The fact that the *nonA* gene is sex-linked actually has implications for the evolution of new song patterns. As Ewing (1969) discussed, sex-linkage could provide the conditions for rapid fixation of a new song characteristic in a population, since a gene controlling song traits, if located on the X chromosome, would have immediate expression in the males. The process of fixation of a new song pattern could be further accelerated if the same gene controlling song production in the males, also controlled song receptivity in heterozygous females. If heterozygous females were able to respond to the new song pattern (implying some form of codominance), they would produce progeny in which the males produce the new pattern and the females are capable of responding to it. Under certain conditions the selective advantage for the new song characteristics could be high, and by virtue of being sex-linked, it could rapidly become fixed in the population. Such behavioural and genetic coupling between song production and song reception has sparked a controversial debate as to how this might evolve (Alexander, 1962; Kyriacou and Hall, 1986; Butlin and Ritchie, 1989).
The major problem in choosing to transfer the *D.virilis nonA* gene into *D.melanogaster* mutant hosts became evident only at the end of my work during the song analyses (chapter 7). As described in that chapter, it was found that the songs of *D.virilis* have positive CPP gradients, a result that was completely unexpected, since none of the many publications from Hoikkala's group had identified this characteristic (Hoikkala et al., 1982; Hoikkala, 1985; Hoikkala and Lumme, 1987). This complicated the interpretations of my results, since one of the principle characters studied in order to determine whether a song is mutant or not, is the CPP gradient (bCPP). Wild-type *D.melanogaster* flies produce songs with very small or negative gradient, while mutant songs (i.e. *nonA*\textsuperscript{dis}) have very pronounced positive gradients (Kulkarni et al., 1988; Rendahl et al., 1992; 1996). When the transformant *nonA*\textsuperscript{-} [nonA\textsuperscript{str}] flies exhibited a mild positive CPP gradient in their lovesong, it became difficult to determine whether this was due to an incomplete rescue of the song phenotype, or a transfer of the *D.virilis* characteristic.

The work described in this thesis consisted in cloning, and characterizing the *D.virilis nonA* gene and studying its functional properties by *P* element-mediated transformation into *D.melanogaster* hosts. This kind of interspecific analysis of genes, including their molecular and functional characterization is not new. Several *D.melanogaster* and *D.virilis* orthologues have been studied (see for example Colot et al., 1988; Treier et al., 1989; Michael et al., 1990; Heberlein and Rubin, 1990; Kassis et al., 1986; O'Neil and Belote, 1992; Hart et al., 1993, Wallrath and Friedman, 1991; Yao and White, 1991; Ousley et al., 1998) and several *D.virilis* genes have been tested, via *P* element-mediated transformation, for their ability to function properly in a *D.melanogaster* background (O'Neil and Belote, 1992; Wallrath and Friedman, 1991; Hart et al., 1993; Heberlein and Rubin, 1990). In all the cases cited, it was found that the *D.virilis* gene was able to partially or completely rescue the corresponding mutant phenotype in *D.melanogaster*. Particularly interesting was the case of the *D.virilis tra* gene which, in spite of displaying a very high level of divergence from its *D.melanogaster* orthologue (the identity of the two species' TRA proteins is only 50%, see chapter 3), was able to rescue, at least partially, the mutant diplo-X *tra* pseudomale phenotype (O'Neil and Belote, 1992). These data were encouraging when initially
planning my project, as the chances of robust rescue of nonA\(^{+}\) phenotype was considered high.

The overall identity of the *D. virilis* and *D. melanogaster* NONA proteins was calculated to be 75%, a figure that did not significantly differ from the levels of identity found between several other *D. virilis* and *D. melanogaster* proteins (see chapter 3, table 3.4). The NONA protein could be divided into two distinct parts: the N-terminal which showed high level of divergence between the two species, and the C-terminal which was almost identical in the two proteins. Not surprisingly the C-terminal contains the two RNA-binding domains and the acidic domain of nonA, which are conserved between NONA and other mammalian and *Drosophila* putative RNA-binding proteins (Rendahl *et al*., 1996). If the major role of NONA is indeed splicing of transcripts that are involved in the determination of visual and lovesong phenotypes, then we could expect that the *D. virilis nonA*, given that its RNA-binding domain is almost identical to that of *D. melanogaster*, might indeed provide these basic functions in *D. melanogaster* flies.

This was the case in the study of the *elav* gene, which also encodes a RNA-binding protein, and whose vital function appears to be that of splicing nervous-system-specific transcripts (Robinow and White, 1988; Robinow and White, 1991). Comparison of the *D. melanogaster* and *D. virilis* ELAV proteins revealed, as in NONA, a highly diverged N-terminal which was also rich in homopolymeric repeats (in this case alanine and glutamine). Three RNA-binding domains encoded within the C-terminal showed an astonishing 100% identity between the two species. Transformation experiments proved that the *D. virilis elav* ORF functioned in a manner indistinguishable from that of the *D. melanogaster*. Moreover, in trying to address the question of whether the diverged region of the protein carried functional information, or whether its presence was simply the result of lowered functional constraints in that part of the protein, Yao and White (1991) deleted the alanine/glutamine-rich domain by 40 amino acids. This deletion had no influence, but the consequences of a complete removal of the alanine/glutamine repeat were not tested.
One of the first decisions which had to be taken in the designing of the experiments, was "how much" of the \( \text{nonA}^{\text{vir}} \) clone was to be used to transform \( D.melanogaster \) flies. Most interspecific transformation experiments resort to using only the coding region of the donor species gene and placing it under the control of the corresponding \( D.melanogaster \) promoter. This ensures that the gene is properly expressed temporally and spatially in the \( D.melanogaster \) hosts (see for example Petersen et al., 1988; Yao and White, 1991). Therefore, transforming a \( \text{nonA}^{\text{vir}} \) DNA fragment, which contained approximately 3 Kb upstream of the beginning of translation and probably more than 1 Kb after the stop codon (although this cannot be precisely determined, see chapter 3, figure 3.8), was slightly risky. There was the possibility that the 3 Kb upstream the ORF did not contain the intact promoter for \( D.virilis \) \( \text{nonA} \). Similarly, I had no proof that the 3' untranslated material was enough to ensure proper termination and adenylation of the \( \text{nonA} \) transcript. Also, there was the possibility that, given the differences in sequence between the 5' regulatory region of \( D.virilis \) and \( D.melanogaster \) (see chapter 3), the promoter of \( \text{nonA}^{\text{vir}} \) would not function properly in the \( D.melanogaster \) background.

The transformation experiments, as described in chapter 4, did not go smoothly. First of all, the construct had the tendency to insert into the X chromosome, which made it difficult to cross into a \( \text{nonA}^{(\cdot)} \) background. The mobilization of insert 113 caused major problems since I was initially confident in the results of the genetic crosses, which indicated that the insert had jumped to a new chromosomal position, at least in line 168-8, because it complemented the lethal 113 homozygous phenotype. As discussed in chapter 4, I do not know what really happened in lines 168-8 and 67-4 and I did not have the time to investigate further. I am curious to find out why the 5' \( P \) element end of the \( \text{nonA}^{\text{vir}} \) construct was lost, possibly at the moment of insertion in the genome, on so many occasions (inserts 113, 72, 297-64 and 297-71) and duplicated in one case (112). My suspicion is that the 3' end of the \( \text{nonA}^{\text{vir}} \) construct (which, after cloning into the \( pW8 \) vector, is immediately adjacent to the 5' \( P \) element end, see chapter 4) might contain numerous repeated sequences. The presence of such repeats could create mispairing of DNA strands, and slippage events during the processes of chromosomal insertion of the construct, which might be likely to result in
the loss or the duplication of the *P* element end. It would therefore be of some interest to sequence the 3' untranslated region of the *nonA*<sup>vir</sup> construct. I also would have liked to find out more about lines 168-8 and 67-4. I showed that the insert in both lines still contained a restriction site which indicated that the sequence lost at the 3' end of each insert could be from a minimum of few bp to a maximum of 2 Kb. Digesting the DNA of 168-8 and 67-4 flies with an enzyme that cuts frequently, would cut this 2 Kb terminal region into several small pieces. After Southern blotting, a probe binding to this 2 kb sequence could be used (as in chapter 4, figure 4.9). This would hybridize to a series of small fragments from the 2 Kb terminal region. If a control in which the terminal 2 Kb region is intact was included in the analysis, it would be possible to determine, by comparison, which of the small fragments are missing in the blots of 168-8 and 67-4, thus discovering more precisely the extent of the deletion (if any) affecting these two inserts.

Chapter 5 presented evidence that the *nonA*<sup>vir</sup> gene completely rescues the semilethality caused by a loss of the *nonA* locus (*nonA*<sup>c</sup>). Furthermore, it was shown that the *nonA*<sup>vir</sup> construct included the sequence of the complete *l(l)H9e* essential gene since it also rescued the deletion affecting both *nonA* and *l(l)il9e* (*T(l;4)9e2-10*, see chapter 5, figure 5.1 and 5.2). These results indicated that *nonA*<sup>vir</sup> was functioning in *D.melanogaster* at least at a basic level.

The visual behaviour experiments (chapter 6) revealed that the *nonA*<sup>vir</sup> gene rescues the mutant (*nonA*<sup>c</sup>) phenotype as well as the *mel nonA*<sup>+</sup> transgene (control line 235R11/d). Only one transformed line (*191/d*) performed significantly worse than the control when tested on small stripes. This result is very likely to be due to position effect.

Two main results need to be underlined from the song analysis of the transformants described in chapter 7. First is the finding on CPP values. The fact that *nonA*<sup>vir</sup>/*nonA*<sup>+</sup> transformants had CPP values in general higher that the corresponding *nonA*<sup>c</sup>/*nonA*<sup>vir</sup> transformants, but wild-type CPP gradients, could only be explained as a transfer of species-specific information (for higher CPP number) from the *D.virilis* gene. This result was reminiscent of the *D.pseudoobscura per* gene which manifested
its effects on the locomotor activity patterns when placed on a *D.melanogaster* wild-type background (Petersen *et al.*, 1988). When placed on the mutant *per*<sup>0</sup> background, its effects were masked by incomplete rescue. Similarly, the effects of the *nonA<sup>vir</sup>* gene on the mutant background (*nonA<sup>c</sup>*) are difficult to interpret because in these transformants the presence of a (moderately) positive CPP gradient does not allow us to rule out the hypothesis of incomplete rescue versus species-specific transfer. Since the *nonA<sup>vir</sup>/nonA<sup>+</sup>* transformants do not show a positive CPP gradient, they cannot be classified as "mutants" and therefore their increased CPP values are likely to be the result of the presence of *nonA<sup>vir</sup>*. The second encouraging result was obtained when studying the fourth moment (kurtosis) of the distribution of CPP values in the *nonA<sup>c</sup> [nonA<sup>vir</sup>]* transformants. The results indicated that these lines were more similar in this aspect to *D.virilis*.

The other interesting result was obtained after applying multivariate analysis to the song data. As seen from figure 7.15, when all the song characters are considered simultaneously, the *nonA<sup>c</sup>/nonA<sup>vir</sup>* (*vir/d*) transformant lines are clearly different from the other transformants carrying the *mel nonA<sup>+</sup>* or the combination of *nonA<sup>vir</sup>* and *nonA<sup>+</sup>* (hybrids *nonA<sup>vir</sup>/nonA<sup>+</sup>*). These results also suggest that subtle effects from the *nonA<sup>vir</sup>* transgene percolate through to bCPP, PF, bPF, IPI and SSF, as all these variables contributed to the significant discrimination between groups. Thus *nonA<sup>vir</sup>* has a small but detectable effect on several song features. While the results are not as striking or as clear-cut as those obtained with the *per* gene (Wheeler *et al.*, 1991), it must be remembered that *per* only transferred the IPI rhythm from *D.simulans* to *D.melanogaster*, but no other song character was affected (Wheeler *et al.*, 1991).

It would be worthwhile to assess whether the song phenotypes associated with the *nonA<sup>vir</sup>* transgene can be attributed to the coding sequence, its regulatory regions or to both. The only definitive way to answer this question would be to construct and transform chimaeric genes composed of reciprocal combinations of the *D.virilis* and *D.melanogaster* coding and regulatory regions. Examination of Northern, Western blots, or immunohistochemistry (assuming the *D.melanogaster* NONA antibody recognizes the *D.virilis* epitopes) would reveal whether the *nonA<sup>vir</sup>* transgene is
expressed normally in the transformant lines. Is it expressed in the right regions, for example in the thoracic ganglia responsible for song production, (Schilcher and Hall, 1979; Hall, 1994)? Are there any indications that the differences seen between the songs of nonA(\textsuperscript{c}[^{nonA^{+}}]) and nonA(\textsuperscript{c}[^{nonA^{vir}}]) transformants could be due to incomplete rescue, perhaps by a quantitative or spatial difference in nonA expression?

If the promoter of nonA\textsuperscript{vir} is not functioning optimally in a D. melanogaster background, we might expect to see differences in the amount or even localization of the nonA product compared to that of the control line. This approach would also reveal whether some of the discrepancies in the results found for different transformant lines (see for example the optomotor results in chapter 6) could be ascribed to changes in expression. However, if the song characteristics carried by the nonA\textsuperscript{vir} transgene were to be attributed exclusively to its coding region, this would further support the assumption that the nonA(\textsuperscript{c}[^{nonA^{vir}}]) transformants are not mutant in their song behaviour, because these flies are normal in their visual behaviour. No coding region mutation of nonA has been isolated that affects solely the song behaviour without affecting vision, so the visual phenotype appears to be the most sensitive to mutation (Rendahl et al., 1992; 1996, Stanewsky et al., 1996). Logically, then, if the nonA(\textsuperscript{c}) [nonA\textsuperscript{vir}] transformants are rescued in their visual behaviour, they should be rescued in their song, indicating species-specific transfer of song traits.

As seen in chapter 3, the N-terminal of the NONA protein has diverged considerably between D. virilis and D. melanogaster. This makes it the most likely candidate for conveying any species-specific characteristics. Interestingly, the most striking difference in the N-terminal of nonA\textsuperscript{vir} is the presence of a very long uninterrupted poly-Glycine repeat. It would be valuable to see whether deleting (or shortening) this repeat has any effect on the visual or song behaviour. It was mentioned in chapter 3 that poly-Glycine regions in RNA-binding proteins could serve as protein-protein interaction domains (Bandziulis et al., 1989). In the case of the per gene, the repetitive Threonine-Glycine tract, which lies in one of the most diverged regions of the protein (Peixoto et al., 1993), is responsible for the transfer of species-

Many more experiments are needed to fully understand the functions of the *nonA* gene. For instance, the *nonA* gene appears to be ubiquitously expressed throughout all the stages of the *Drosophila* life cycle (Rendahl *et al.*, 1992). The limited and specific arrays of phenotypes displayed by *nonA* mutants is therefore rather surprising. One of the questions that could be addressed is where (i.e. in which organs), and when, is the product of *nonA* needed for the fly to display normal behaviour? The question "when" is the *nonA* product necessary has already been answered by Rendahl and Hall (1996), (as mentioned in chapter 1) by placing *nonA* under the control of a heat shock promoter and switching its expression at determined developmental stages. Federica Sandrelli, a doctoral student from Padova, Italy, and I have started to investigate "where" the product of *nonA* must be expressed to generate normal behaviour. We have performed a functional dissection of the *D.melanogaster nonA* promoter, by making a series of nested deletions. Promoter sequences containing deletions ranging from a minimum of ~200 bp to a maximum of ~1700 bp were placed 5' of the yeast transcriptional activator gene GAL4 (Brand and Perrimon, 1993), while the *D.melanogaster nonA* cDNA (Besser *et al.*, 1990) was cloned downstream of the GAL4-responsive UAS (Brand and Perrimon, 1993). A transformant line carrying the fusion of UAS with the reporter gene *lacZ* was also used in order to determine the expression pattern of the different *nonA*-GAL4 constructs. At the same time, crosses between the *nonA*-GAL4 lines with the UAS-*nonA* line permitted us to study the behavioural effects (on a *nonA*<sup>dis</sup> mutant background) of the promoter deletions. The ultimate goal of this project is to correlate changes in the temporal and spatial expression of UAS-*nonA* with the behavioural phenotypes of the transformants, and with the relevant promoter sequences. At the moment we are waiting for the behavioural results of this project.

The recent cloning of the Dmca1A-encoding locus (Smith *et al.*, 1996) whose *cac* mutation produces song and visual aberrations strikingly similar to those observed for *nonA*<sup>dis</sup> (Kulkarni and Hall, 1987) permits a test of whether the Dmca1A transcript
could possibly be the target of NONA. The transcript of the DmcalA-encoding locus displays a complicated pattern of alternative splicing which could be regulated by NONA, as already discussed in chapter 1. nonA mutants (nonA^{cl} or nonA^{dist}) flies could be transformed with any of the DmcalA-encoding cDNAs under the control of a heat shock promoter. If such a transgene rescued the visual and song phenotypes of nonA mutants it would be an indication that the DmcalA-encoding gene is in fact regulated by the nonA gene, presumably by splicing. Under this scenario, nonA^{vir} might splice a different isoform of DmcalA in the D.melanogaster transformants, suggesting a different profile of DmcalA cDNAs in D.virilis.

To conclude, if my interpretation is correct, nonA^{vir} may carry some species-specific information for the song characteristics that I have studied, because subtle but detectable changes were observed in the transformants. However, the larger differences in the song patterns between D.melanogaster and D.virilis are undoubtedly encoded by many genes. nonA may contribute a small component of this variance, thereby conforming to the traditional view that species-specific characteristics accumulate by small changes at many loci (Mayr, 1977; Coyne, 1992).
APPENDICES

FOR CHAPTER 3
Appendix 3.1.

Sequence and position (relative to the D. virilis nonA gene sequence, figure 3..) of the primers used in the sequencing of the D. virilis nonA gene. * = the position of this primer is relative to the D. melanogaster nonA gene sequence by Jones and Rubin (1990).

<table>
<thead>
<tr>
<th>PRIMER</th>
<th>SEQUENCE</th>
<th>POSITION</th>
</tr>
</thead>
<tbody>
<tr>
<td>5′E1</td>
<td>5′-ATGGAAAAATTCTGTAAA-3′</td>
<td>1-19</td>
</tr>
<tr>
<td>3′E1</td>
<td>5′-AAATCCACCTTTGTCG-3′</td>
<td>276-261</td>
</tr>
<tr>
<td>3′7A</td>
<td>5′-ATTTCCTCCGTCTGCTTT-3′</td>
<td>393-376</td>
</tr>
<tr>
<td>5′7B</td>
<td>5′-CACAACCTCAAGGCAGGC-3′</td>
<td>464-483</td>
</tr>
<tr>
<td>3′POL</td>
<td>5′-CTAATGCTGCGTAACCGTTG-3′</td>
<td>791-772</td>
</tr>
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Appendix 3.2. Translation in the three reading frames (a, b, c) of the beginning of the *D. virilis* nonA intron four.

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   a  VCKRTGHS X VSCYI * HMAYG
   b  YVNQDIVL * AAIYSIWHMV
   c  M * T YRT * YCEL LLY I AY GIWS

61 CGACTCCCATATGAAACCAGATGCGCCGCGCCTGTGCCCTTTATATATAACATAAAAGT 120
   a  RLPEYETRCRPRPVPLLI * HK S
   b  DSHMKPDAAGICLPY * YNIKV
   c  TPI * NQMPACALINIT * KL

121 TACCATTGTTGCGCGTCATTTATCGTTATATAATTCAATGACTGTCTCTCATCAT 180
   a  YHCRRSFFIGYISMTVLT H
   b  TIVAVALHLSVI * FQLFSLM
   c  PLLPSLlYRLYNFNDSCSHSC

181 GTAGATAGCAATCAA 197
   a  VDSDKS
   b  * IANQ
   c  R * QIK
Appendix 3.3.  Alignment of the *D. virilis* and *D. melanogaster nonA* DNA sequence. The sequence homologous to the DBHS domain (see Introduction) is highlighted in yellow. The two RNA-binding domains (RRM1 and RRM2) are underlined. The RNP1 motifs are shown in blue and the RNP2 in red.
vir 1438 GGCTCCAGGTGGAAACAGTTGCAGATGCATATTCATAAAGCAAGCAGATGC    1487
mel 1489 GTTTCGCGCTGGAAGCGCTGACAATTTGTCTAACAAAGGAGGAGGACGC    1538
vir 1488 TCTCAAGCGTGAAGCTCAAAAATGGAAGAAGAAAACTTGTGATCGCATATGGG    1537
mel 1539 ACTCAAACCGGAGCTGAAATGGAAGAGGACAAGTGTGGCTGACAGTGCA    1588
vir 1538 AGTATGCTCGCTATGAGCAAGAAGACAGCTTTTGCGTCAAGAATAATG    1587
mel 1589 AGTATGCGACCTAGCTAGGAGAGAAACTTGAATCTGTTGCGTACAGAGTCG   1638
vir 1588 AAACGCGAATCTGATATGAGCAAGAAGACAGCTTTTGCGTCAAGAATAATG    1637
mel 1639 AAACGCGAATCTGATATGAGCAAGAAGACAGCTTTTGCGTCAAGAATAATG    1688
vir 1638 GAARACAGCCCGAGGAAATGCGCAACACCGGAGAGGAGGACCATCGCTGCTC    1687
mel 1689 GAARACAGCCCGAGGAAATGCGCAACACCGGAGAGGAGGACCATCGCTGCTC    1738
vir 1688 ATCAAAACCAGAAATGCAAGGCGCTATGCGGCAAGAGGAGGATAATGCGT    1737
mel 1739 ATCAAGACCGAGATGCAAGGCGCCACATAGAATCGCCAGAGGAGGACATGCTC    1788
vir 1738 CGTCGTACAGCAGGAAACACATATTATTCAGAGCAGAGCAGCTAAACTC    1877
mel 1789 CGTCGCGACCGAGGAG...ACACTGTTCATGAAAGCAGACAGCTCATTTC    1835
vir 1788 GCTGCTGGATGTACACAGGAAAGGCTTCTGCGGCGGCAACGCTGTGGCGGCG    1837
mel 1836 GTTGTGCGATGAGCAAGGAGGATTT......................... 1859
vir 1838 GCCGGCGTGAGGCCGGCGGCCGGCGGTGGCAATACATTTTGATAACTTC    1887
mel 1860 .......TGGTGCGCGCGCGCGCTAACAATCTCCACCTTGTGACAGACTTC    1902
vir 1888 GGCGGCAACACGAACTCCTCCTGTTGAAGTGTGCAGGAG...........TAATAA    1931
mel 1903 GCTGGCAATAGCAATTCGCCATTCGAGGTTGTTTAGAGGCAATACAATAA    1952
vir 1932 CAACCTGTCAGCTTGCAAAACATGGCCGCGGCTGCGCAAAATACGAC    1981
mel 1953 CAATTCCCAAAATGATTGGAACAAATGCTGGCCGGCCACACA........... 1992
vir 1982 AGCAGGACTCTCGTCTGCTATTTGAAATTTGGGTTAACAATATGACCAAA    2031
mel 1993 ..CAGGACTCTTCC...GCTTTGTGAATTTGGGTTAACAATATGACCAA    2037
vir 2032 GCCGGCAATCAAGCTGGAAATAATGGA...GAAAATATGTTTCCATGGG    2078
mel 2038 GCCGGCAATCAAGCTGGAAATAATGGA...GAAAATATGTTTCCATGGG    2087
vir 2079 ACGACGTGCTTTTAGAC 2096
mel 2088 ACGACGTGCTTTTAGAC 2103
Appendix 3.4. Chou-Fasman secondary structure prediction of the RNA-binding domain of the *D. virilis* NONA protein.
Appendix 3.5. Chou-Fasman prediction of the secondary structure of the RNA-binding domain of the *D.melanogaster* NONA protein.
APPENDICES

FOR CHAPTER 6
Appendix 6.1. Newman-Keuls probability matrix based on the two way ANOVA for the optomotor test results. Any significant probability (p< 0.05) is shaded in grey. Bold numbers at the head of each column indicate each genotype's average correct turns.
APPENDICES

FOR CHAPTER 7
Appendix 7.1.

Newman-Keuls probability matrix based on the ANOVA for bCPP. Only transformant lines are included. Any significant probability (p< 0.05) is shaded in grey.

Bold numbers at the head of each column indicate each genotype's average bCPP value.

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Appendix 7.2.
Newman-Keuls probability matrix based on the ANOVA for CPP1. Only transformant lines are included. Any significant probability (p< 0.05) is shaded in grey.
Bold numbers at the head of each column indicate each genotype's average CPP1 value.

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{2}

6 7 -4 /d

{3}

1 6 8 -8 /d
1 3 5 /d
9 7 /d
1 9 1 /d

{4}
{5}
{6}
{7}

{3 }

{4}

{5 }

{6}

{7}

{8}

{9}

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2 .0 0 8 7 0 0

1 .9 5 5 0 7 0

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1 .7 2 2 7 6 0

1 .9 6 6 7 1 0

1 .7 1 2 7 0 0

1 .7 5 2 7 8 0

0 .9 3 1 1 5 1

2 3 5 R 1 1 /d {1 }
1 1 3 /d

{2 }

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{1 3 }

{1 4 }

{1 5 }

{1 6 }

{1 7 }

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1 .8 0 3 2 2 0

1 .8 4 0 9 2 0

1 .8 5 3 1 0 0

1 .7 4 2 1 6 0

1 .5 6 5 8 5 0

0 .0 0 0 0 4 1

0 .0 0 0 4 1 4

0 .6 9 2 7 2 4

0 .4 4 0 0 1 7

0 .0 0 0 2 4 4

0 .3 5 7 6 3 8

0 .4 1 6 9 4 4

0 .0 0 0 0 2 6

0 .0 0 0 0 2 3

0 .1 5 3 6 9 5

0 .0 5 5 8 5 7

0 .0 4 3 6 2 5

0 .4 0 8 3 4 4

0 .8 7 5 0 2 6

0 .0 0 0 0 3 6

0 .0 0 0 0 3 8

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0 .0 0 0 1 5 7

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0 .5 0 8 8 5 8

0 .0 1 0 2 0 6

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0 .5 1 3 6 8 1

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0 .8 5 5 6 1 0 | 0 .8 8 2 7 7 0

0 .0 0 1 1 6 8

0 .0 4 1 8 5 7

1 1 2 /m

{9}

0 .3 5 7 6 3 8

0 .4 6 5 1 5 6

0 .0 0 7 8 4 8

0 .0 0 8 1 8 1

0 .0 4 4 4 4 5

0 .3 3 0 4 8 1

0 .8 9 8 5 9 0

1 1 3 /m

{1 0 }

{1 1 }
1 6 8 -8 /m {1 2 }

{1 2 }

0 .0 0 0 0 4 0

2 9 7 -6 /d

6 7 - 4 /m

{1 1 }
2 .3 6 3 9 8 3

0 .0 3 5 0 4 0

0 .4 1 6 9 4 4

0 .4 3 8 8 1 1

0 .0 2 2 9 5 1

0 .0 2 6 1 3 1

0 .0 7 7 3 6 0

0 .5 7 4 9 2 8

0 .9 2 3 3 6 7 | 0 . 0 7 3 1 5 8 I 0 .9 5 7 2 6 9

0 .0 0 0 0 2 6

0 .0 0 0 0 2 9

0 .0 0 0 0 4 0

0 .0 0 0 0 4 0

0 .0 0 0 0 2 3

0 .0 0 0 0 2 3

0 .0 0 0 0 1 7

0 .0 0 0 0 2 1

0 .0 0 0 0 2 0

0 ,0 0 0 0 1 2

0 .0 0 0 0 2 3

0 .0 0 0 0 2 6

0 .0 0 0 5 0 4

0 .0 0 0 2 0 1

0 . 0 0 0 1 1 1 0 .0 0 0 0 2 0

0 .0 0 0 0 1 5

0 .0 0 0 1 2 7

0 .0 0 0 0 1 7

0 .0 0 0 0 1 0

0 .4 3 8 1 7 4

0 .6 3 2 9 0 7

0 .1 5 3 6 9 5

0 .1 5 5 1 6 2

0 .1 0 5 9 0 0

0 .1 2 4 8 3 0

0 .2 1 7 9 3 6

0 .2 7 6 7 7 9

0 .7 3 8 0 7 9

0 .2 3 2 5 5 1

0 .7 8 1 4 8 4

0 .5 2 2 7 9 3

0 .0 0 0 0 1 0

0 .0 0 0 0 3 2

9 7 /m

{14}

0 .0 5 5 8 5 7

0 .0 5 3 4 5 7

0 .2 2 5 3 1 0

0 .2 7 3 8 6 5

0 .3 1 7 2 2 1

0 .1 2 6 5 6 1

0 .5 6 4 3 9 8

0 .3 8 2 1 7 3

0 .5 8 2 2 0 5

0 .5 0 3 7 2 6

0 .0 0 0 0 3 2

0 .0 0 0 0 2 6

0 .6 3 2 9 0 7

2m el

{1 5 }

0 .0 4 3 6 2 5

0 .0 4 0 3 7 3

0 .2 1 4 0 2 3

0 .2 8 0 0 4 7

0 .1 9 6 3 9 1

0 .1 0 7 4 4 2

0 .5 6 4 2 4 8

0 .3 2 0 6 6 3

0 .5 6 2 4 7 6

0 .5 8 1 4 6 7

0 .0 0 0 0 2 6

0 .0 0 0 0 2 0

0 .8 0 2 5 2 2

0 .8 7 7 3 7 3

3m el

{1 6 }

0 .4 0 8 3 4 4

0 .4 4 7 5 7 1

0 .0 1 9 0 7 0

0 .0 2 0 9 7 5

0 .0 7 5 6 9 1

0 .5 3 3 2 5 8

0 .8 0 5 8 6 3

0 .0 6 6 8 7 4

0 .9 2 6 0 9 1

0 .8 9 2 9 7 8

0 .0 0 0 0 1 5

0 .0 0 0 0 1 2

0 .7 1 9 2 6 2

0 .5 9 4 1 2 2

0 .6 2 4 0 6 5

0 .8 7 5 0 2 6

0 .6 8 4 8 8 7

0 .0 0 0 0 2 5

0 .0 0 0 0 2 8

0 .0 0 0 0 6 6

0 .8 1 1 4 3 9

0 .3 4 9 1 4 1

0 .0 0 0 0 4 6

0 .3 3 8 8 8 8

0 .2 5 7 0 2 2

0 .0 0 0 0 3 3

0 .0 0 0 0 2 9

0 .0 6 5 8 1 1

0 .0 1 7 7 1 6

0 .0 1 2 2 9 7

1 3 5 /m

{1 3 }

m e l+ 2 v ir { 1 7 }

175

0 .8 7 7 3 7 3

0 .5 9 4 1 2 2

0 .0 1 7 7 1 6

0 .6 2 4 0 6 5

0 .0 1 2 2 9 7
0 .2 7 7 2 7 0

0 .2 7 7 2 7 0


Appendix 7.3.

Newman-Keuls probability matrix based on the ANOVA for CPP10. Only transformant lines are included. Any significant probability (p< 0.05) is shaded in grey. Bold numbers indicate each genotype's average CPP10 value.

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<td>0.000026</td>
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<td>0.818781</td>
<td>0.01552</td>
<td>0.425991</td>
<td>0.933398</td>
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<td>0.000017</td>
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<td>0.000026</td>
<td>0.093715</td>
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<td>0.01552</td>
<td>0.425991</td>
<td>0.933398</td>
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<td>0.425991</td>
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<tr>
<td>135/d (5)</td>
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<td>0.000026</td>
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<td>0.608517</td>
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<td>0.047555</td>
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<td>0.156081</td>
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<td>mel+2vir (17)</td>
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<td>0.000026</td>
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<td>0.000028</td>
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<td>0.101987</td>
<td>0.081398</td>
<td>0.450081</td>
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Appendix 7.4.

Kolmogorov-Smirnov tests. A: hemizygous nonA\textsuperscript{v}!(nonA\textsuperscript{v}) and heterozygous nonA\textsuperscript{v}!nonA\textsuperscript{+} lines are respectively grouped as vir/d and vir/m. B: each transformant line is compared to 235R11/d, diss and vir. Maximum differences between pair of genotypes are indicated. * = significant differences at p< 0.05. ** = significant differences at p< 0.01.

### A:

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<th>vir/d</th>
<th>vir/m</th>
<th>vir</th>
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<td>-0.900**</td>
<td>-0.843**</td>
<td>-0.893**</td>
<td>-0.800**</td>
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<tr>
<td>235R11/d</td>
<td>-0.486*</td>
<td>-0.429</td>
<td>-0.700*</td>
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<tr>
<td>vir/d</td>
<td>-0.214</td>
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<tr>
<td>vir/m</td>
<td></td>
<td>-0.518*</td>
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### B:

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<th>vir</th>
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<td>0.50</td>
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<td>67-4/d</td>
<td>-0.60</td>
<td>0.80**</td>
<td>0.50</td>
</tr>
<tr>
<td>168-8/d</td>
<td>-0.50</td>
<td>0.90**</td>
<td>0.60</td>
</tr>
<tr>
<td>135/d</td>
<td>-0.50</td>
<td>1.00**</td>
<td>0.60</td>
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<tr>
<td>97/d</td>
<td>-0.50</td>
<td>0.90**</td>
<td>0.40</td>
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<td>191/d</td>
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<td>0.80**</td>
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<td>0.70*</td>
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<td>1.00**</td>
<td>0.80**</td>
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<td>0.90**</td>
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<td>135/m</td>
<td>-0.70*</td>
<td>0.90**</td>
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<td>mel+2vir</td>
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Appendix 7.5.

Newman-Keuls probability matrix based on the ANOVA for bPF. Any significant probability (p< 0.05) is shaded in grey. Bold numbers at the head of each column indicate each genotype's average bPF value.
Appendix 7.6.

Newman-Keuls probability matrix based on the ANOVA for SSF. Any significant probability (p<0.05) is shaded in grey. Bold numbers at the head of each column indicate each genotype's average SSF value.

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</tr>
<tr>
<td>235R11/d</td>
<td>0.32139</td>
<td>0.27112</td>
<td>0.35088</td>
<td>0.69210</td>
<td>0.47791</td>
<td>0.04938</td>
<td>0.88263</td>
<td>0.40519</td>
<td>0.80471</td>
<td>0.00018</td>
<td>0.00011</td>
<td>0.25673</td>
<td>0.79345</td>
<td>0.19669</td>
<td>0.35134</td>
<td>0.34180</td>
<td>0.197616</td>
<td></td>
</tr>
<tr>
<td>97/m</td>
<td>0.00390</td>
<td>0.00172</td>
<td>0.00113</td>
<td>0.09458</td>
<td>0.71954</td>
<td>0.53054</td>
<td>0.21159</td>
<td>0.00101</td>
<td>0.35191</td>
<td>0.00003</td>
<td>0.00002</td>
<td>0.74854</td>
<td>0.05736</td>
<td>0.19659</td>
<td>0.51204</td>
<td>0.00034</td>
<td>0.00008</td>
<td></td>
</tr>
<tr>
<td>2mel</td>
<td>0.04752</td>
<td>0.09971</td>
<td>0.08003</td>
<td>0.54436</td>
<td>0.02236</td>
<td>0.00000</td>
<td>0.00000</td>
<td>0.00000</td>
<td>0.00000</td>
<td>0.00000</td>
<td>0.00000</td>
<td>0.00000</td>
<td>0.00000</td>
<td>0.00000</td>
<td>0.00000</td>
<td>0.00000</td>
<td>0.00000</td>
<td>0.00000</td>
</tr>
<tr>
<td>3mel</td>
<td>0.94939</td>
<td>0.96101</td>
<td>0.92149</td>
<td>0.53616</td>
<td>0.00000</td>
<td>0.00000</td>
<td>0.00000</td>
<td>0.00000</td>
<td>0.00000</td>
<td>0.00000</td>
<td>0.00000</td>
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<td>0.00000</td>
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<td>0.00000</td>
<td>0.00000</td>
<td>0.00000</td>
<td>0.00000</td>
</tr>
<tr>
<td>mel+2we</td>
<td>0.90713</td>
<td>0.96129</td>
<td>0.94689</td>
<td>0.36657</td>
<td>0.00169</td>
<td>0.00000</td>
<td>0.16541</td>
<td>0.90750</td>
<td>0.07256</td>
<td>0.01327</td>
<td>0.33158</td>
<td>0.00375</td>
<td>0.46306</td>
<td>0.18767</td>
<td>0.00098</td>
<td>0.01777</td>
<td>0.72960</td>
<td></td>
</tr>
</tbody>
</table>
Appendix 7.7.

Newman-Keuls probability matrix based on the ANOVA for IPI. Any significant probability (p < 0.05) is shaded in grey. Bold numbers at the head of each column indicate each genotype's average IPI value.
Appendix 7.8. Results of the discriminant analysis for the *MEL* group.

<table>
<thead>
<tr>
<th>Traits</th>
<th>F (2, 27)</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>IPI</td>
<td>13.7</td>
<td>&lt;=0.0001</td>
</tr>
<tr>
<td>PF</td>
<td>9.58</td>
<td>0.0007</td>
</tr>
<tr>
<td>CPP</td>
<td>4.82</td>
<td>0.016</td>
</tr>
<tr>
<td>bCCP</td>
<td>4.18</td>
<td>0.026</td>
</tr>
<tr>
<td>bPF</td>
<td>0.78</td>
<td>0.47</td>
</tr>
<tr>
<td>SSF</td>
<td>8.32</td>
<td>0.0015</td>
</tr>
</tbody>
</table>

\[ \lambda = 0.107 \]
\[ F = 7.53 \]
\[ d.f. = 12.44 \]
\[ P = <=0.001 \]

Appendix 7.9. Results of the discriminant analysis for the *vir/d* group.

<table>
<thead>
<tr>
<th>Traits</th>
<th>F (6, 63)</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>IPI</td>
<td>3.4</td>
<td>0.0057</td>
</tr>
<tr>
<td>PF</td>
<td>6.55</td>
<td>&lt;=0.0001</td>
</tr>
<tr>
<td>CPP</td>
<td>10.11</td>
<td>&lt;=0.0001</td>
</tr>
<tr>
<td>bCCP</td>
<td>1.71</td>
<td>0.133</td>
</tr>
<tr>
<td>bPF</td>
<td>2.75</td>
<td>0.0195</td>
</tr>
<tr>
<td>SSF</td>
<td>7.56</td>
<td>&lt;=0.0001</td>
</tr>
</tbody>
</table>

\[ \lambda = 0.089 \]
\[ F = 5.25 \]
\[ d.f. = 36, 257 \]
\[ P = <=0.001 \]
Appendix 7.10. Results of the discriminant analysis for the hybrids (*vir*/*m* and *mel+2vir*).

<table>
<thead>
<tr>
<th>Traits</th>
<th>F (6, 58)</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>IPI</td>
<td>4.21</td>
<td>0.0014</td>
</tr>
<tr>
<td>PF</td>
<td>23.29</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>CPP</td>
<td>23.20</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>bCCP</td>
<td>2.57</td>
<td>0.028</td>
</tr>
<tr>
<td>bPF</td>
<td>3.89</td>
<td>0.003</td>
</tr>
<tr>
<td>SSF</td>
<td>18.22</td>
<td>&lt;0.0001</td>
</tr>
</tbody>
</table>

$\lambda$ = 0.027

<table>
<thead>
<tr>
<th>F</th>
<th>8.34</th>
</tr>
</thead>
<tbody>
<tr>
<td>d.f.</td>
<td>36, 235</td>
</tr>
<tr>
<td>P</td>
<td>&lt;0.0001</td>
</tr>
</tbody>
</table>

$\lambda$ = 0.027

Appendix 7.11. Results of the discriminant analysis for all the transformants (excluding *mel+2vir*).

<table>
<thead>
<tr>
<th>Traits</th>
<th>F (4, 94)</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>IPI</td>
<td>15.08</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>PF</td>
<td>7.46</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>CPP</td>
<td>2.26</td>
<td>0.0680</td>
</tr>
<tr>
<td>bCCP</td>
<td>7.67</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>bPF</td>
<td>9.64</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>SSF</td>
<td>3.29</td>
<td>0.014</td>
</tr>
</tbody>
</table>

$\lambda$ = 0.274

<table>
<thead>
<tr>
<th>F</th>
<th>5.82</th>
</tr>
</thead>
<tbody>
<tr>
<td>d.f.</td>
<td>24, 311</td>
</tr>
<tr>
<td>P</td>
<td>&lt;0.0001</td>
</tr>
</tbody>
</table>


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RYNER, L.C., GOODWIN, S.F., CASTRILLON, D.H., ANAND, A., VILLELLA, A., BAKER, B.S., HALL, J.C., TAYLOR, B.J., and WASSERMAN,
considerable within-species diversity. Nucleic Acids Res. 16: 8207-8211.


nuclear protein common to many cell types, results in specific physiological and behavioral defects. *Genetics* 135: 419-442.


