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Chapter 4 - Isolation and Preliminary Characterisation of cnox1Am and barhAm genes from Acropora millepora - no clear functional correlation with the Hox11-like genes or the Bar family

4.1 Introduction

The Bar and Hox11/Tlx families are defined by the presence of a threonine residue at homeodomain position 47, a feature not found in other homeodomain families. Although many members of the Hox11 family are as yet largely uncharacterised, most of the characterised members function in nervous system patterning. Prior to the start of this project an A. millepora genomic clone was isolated in our laboratory, and shown to contain a Tlx/Hox11-related gene, that was most similar to the predicted Drosophila gene cg13424 and is hereafter known as cnox1Am. In addition, a preliminary EST analysis of Acropora (Kortschak et al., 2003) led to the identification of a Bar-related gene, referred to here as barhAm. As members of these gene families function in nervous system patterning in higher animals, their temporal and spatial expression patterns were investigated.

4.1.1 cnox1Am and cg13424 - Related to the Hox11 and Mnx families

Preliminary BLASTx analysis indicated that Drosophila CG13424 was the closest sequence in the database to the partial sequence available for the Acropora CnoxlAm from the EST; the predicted homeodomains have 72%/80% identity/similarity. Comparison of the CG13424 protein with others clearly placed it into the Tlx/Hox11 group, with no obvious matches to other homeobox gene families, thus the implication was that CnoxlAm was likewise a Tlx/Hox11 family member. However, analysis of the full-length CnoxlAm protein indicates similarities not only to the Tlx/Hox11 group, but also to HB9/Mnr2 proteins from chicken and zebrafish, which define a subfamily of the Mnx class of homeodomain containing transcription factors. Additionally, several clear orthologs of CG13424 have subsequently been added to the database (see Fig 4.2).
So far only 3 members of the Mnr2 family have been described, chick Mnr2 and zebrafish Mnr-2a and -2b, and it appears this subfamily of Mnx genes has been lost in mammals. In the chick, studies on the effect of ectopic expression of Mnr2 have shown that it plays a major role in specifying various aspects of motor neuron identity and in interneuron fate decisions (Tanabe et al., 1998). Related genes of the HB9 family have been identified not only in vertebrates (e.g., mouse, Xenopus and chicken), but also in invertebrates (e.g., sea urchin, amphioxus and Drosophila) (Saha et al., 1997; Bellomonte et al., 1998; Ferrier et al., 2001; Broihier and Skeath, 2002; Odden et al., 2002). In vertebrates, HB9 genes are involved in the first stages of pancreas morphogenesis and in the differentiation of β-cells. In invertebrates (which lack a pancreas) embryonic expression of hb9 is in the primitive trunk endoderm (Bellomonte et al., 1998; Ferrier et al., 2001; Broihier and Skeath, 2002; Odden et al., 2002). The relevance of this highly conserved expression remains to be investigated. In both vertebrates and invertebrates, hb9 genes are essential for motoneuron consolidation (Arber et al., 1999; Thaler et al., 1999) and are required for neuronal migration and axonal outgrowth in mouse, chicken and Drosophila (Tanabe et al., 1998; Arber et al., 1999; Thaler et al., 1999; Broihier and Skeath, 2002; Odden et al., 2002; Williams et al., 2003). The N-terminal domain of MNR2 is required both in vivo and as a transcriptional repressor in in vitro cell-based reporter assays (William et al., 2003). In mouse, HB9 has been shown to repress its own expression (Arber et al., 1999; Thaler et al., 1999). The precise mechanism of MNR2- and HB9-mediated transcriptional repression remains unclear. MNR2, like many other HD proteins (Muhr et al., 2001), possesses a well-conserved Eh1 motif that, in other contexts, can recruit Groucho class co-repressors (Smith and Jaynes, 1996). However, deletion of the Eh1 motif in MNR2 does not affect its role in motor neuron generation and when the HD of MNR2 is spliced to a powerful Groucho recruitment domain, induction of motoneurons in vivo is poor (William et al., 2003). It therefore seems that interaction with Groucho co-repressors is not essential for the repressive function of MNR2 (and by implication HB9), and it may be that MNR2 repressor activity requires the recruitment of Ctbp class co-repressors (William et al., 2003). Evidence has suggested that in some Drosophila proteins, Eh1-recruitment and Ctbp-recruitment domains might cooperate to mediate repressor function (Hasson et al., 2001; Barolo et al., 2002).
Tlx/Hox11 genes are orphan homeobox genes that play critical roles in the regulation of early developmental processes in vertebrates. The name of this family of genes is confusing because they are not true Hox genes and are physically located outside the Hox gene clusters. In addition, the designation Tlx, (for T-cell leukaemia homeobox-containing gene) (Raju et al., 1993) is not accurate as expression of these genes is not restricted to T-cells (Andermann and Weinberg, 2001), and the same name has subsequently been used to identify vertebrate homologs of the Drosophila zinc-finger tailless gene (Yu et al., 1994). The core DNA motif recognised by HOX11 has been determined to be TAAGTG or TAATTG using in vitro selection techniques (Tang and Breitman, 1995) and direct interactions between the homeodomain of HOX11 and the regulatory regions of specific genes is most likely essential for its role as an oncogene (Owens et al., 2003). Mouse HOX11 has both transcription activation and repression roles, and its repressive function is independent of DNA-binding; it is most likely mediated via protein-protein interactions with other transcription factors (Owens et al., 2003).

In humans, the HOX11 gene was originally identified due to its association with the breakpoint in specific chromosome translocations in patients with acute T-cell leukaemia, and has since been found to be essential for spleen development during embryogenesis (Dube et al., 1991; Hatano M et al., 1991; Kennedy et al., 1991; Lu et al., 1991; Heidari et al., 2002). The mouse ortholog Hox11, is involved in splenogenesis, but is also expressed in the developing hindbrain, spinal cord and the neurons of the developing cranial sensory ganglia (Dear et al., 1993; Raju et al., 1993; Roberts et al., 1995). The presumptive chicken ortholog exhibits a similar expression pattern (Logan et al., 1998), while the potential Xenopus ortholog, Xhox11 is expressed in regions similar to the mouse, but not the spleen (Patterson and Krieg, 1999). Zebrafish possesses three Tlx/Hox11 genes; Tlx-1, Tlx-3a and -3b; Tlx-3a and -3b most likely represent the result of a duplication of an ancestral Tlx-like gene. Similar to the expression of other vertebrate Tlx/Hox11 genes, Tlx-1 is expressed early in regions of the brain and is also present in splenic primordial tissue, although it differs with that of its orthologs in vertebrates as it is not present in the cranial sensory ganglia or spinal cord (Langenau et al., 2002). Tlx-3a and Tlx-3b appear to have complementary expression patterns in the developing nervous system – Tlx-3a is expressed in the
cranial ganglia, enteric neurons and some non-neural tissues while Tlx-3b is expressed in the dorsal root ganglia (Langenau et al., 2002).

4.1.2 BarhAm – A cnidarian member of the Bar class of transcription factors

The Bar class of homeodomain-containing transcription factors have been described from a wide range of organisms. Two atypical characteristics define and distinguish the Bar class - the presence of a threonine residue at homeodomain position 47 (T47) rather than an isoleucine or valine residue, and the presence of a tyrosine residue (Y49) rather than the near-universal phenylalanine (F49) residue in helix 3 (Burglin, 1994). Note that in the Hox11/Tlx family, a threonine at position 47 is also present.

Bar class genes were first identified in Drosophila, after investigation of the locus involved in the Bar mutation in Drosophila melanogaster and the related Om(1D) mutation of Drosophila ananassae, which impedes ommatidium differentiation. In both cases, the loci responsible contained a novel homeobox gene of the Bar class, BarH1 (Kojima et al., 1991). Characterisation of the locus revealed a second Bar gene, BarH2 and both Drosophila Bar genes have since been found to have multiple roles in eye development including the formation of pigment and cone cells (Higashijima, 1992; Higashijima et al., 1992). In addition, the BarH1 and BarH2 regulate microchaetae formation in the anterior notum and are important in the development of distal leg segments (Sato et al., 1999; Kojima, 2000).

In the mouse, four Bar class genes have been identified, Barhl1, Barhl2, Barx1 and Barx2. While the BARX proteins had previously been reported to be closely related to the Drosophila Bar proteins, it is now accepted that they belong to a different Bar subfamily and the BARHL1/2 proteins are orthologous to the BarH1/2 proteins of Drosophila. The Barhl1 gene is restricted to the developing CNS, where it may play a role in cell fate determination (Bulfone et al., 2000), and to the hair cells in the inner ear (Li et al., 2002). Although expression data are not yet available for murine Barhl2, expression of the rat ortholog, MBH1 is specific to the eye and CNS (Saito et al., 1998). The Barx1 and Barx2 genes have overlapping expression patterns in the central nervous system, including in the telencephalon and spinal cord, while in other non-neural tissues
(for example the developing facial structures), their expression patterns are complementary (Tissier-Seta et al., 1995; Jones et al., 1997).

Two Bar class homeoproteins have also been isolated from the chick; Barx-1 and Barx1b (Barlow et al., 1999; Nakamura et al., 2001). The homeodomains of both chick Bar proteins are identical to each other, and to mouse Barx1 (Tissier-Seta et al., 1995; Barlow et al., 1999; Nakamura et al., 2001), however their NH₂ and COOH termini show no conservation (Nakamura et al., 2001). In both chick and mouse, Barx-1 expression is present in the stomach, limbs and early facial structures (Tissier-Seta et al., 1995; Barlow et al., 1999). Chick Barx1b is also expressed in similar regions, but expression extends to the smooth muscle cells of the upper digestive system (Nakamura et al., 2001).

Members of the Bar class of homeobox genes have also been found in Xenopus (XBH1, XBH2, Xbr1a) (Patterson et al., 2000), sponge (prox2) (Seimiya et al., 1994), and the medaka fish (OlBar) (Poggi et al., 2002). Prior to this study, only one cnidarian member of the Bar class of genes had been isolated (cnox3; Chlorohydra viridissima (Schummer et al., 1992)). However, other cnidarian genes have been given the name cnox3 since, even though they are not related to the Bar class, such as in Hydra vulgaris (Gauchat et al., 2000), Podocoryne carnea (unpublished; AC# BAC56129), and Eleutheria dichotoma (Kuhn et al., 1996). Preliminary analysis implied that the Chlorohydra Bar ortholog cnox3, was more highly expressed in the head region than along the body axis, and on this basis a role during head regeneration has been suggested (Schummer et al., 1992).

4.1.3 Statement of goals

Two partial sequences encoding cnidarian orthologs of the Tlx/Hox11 and Bar homeobox genes were recently isolated from the coral Acropora millepora. The cnox1Am locus was identified in a genomic clone by Dr Julian Catmull, while a ~380bp Bar-like fragment was identified in a preliminary EST analysis of A. millepora. In view of the roles of related genes in higher animals, these coral genes were characterised at the level of expression patterns. In parallel, the Drosophila homolog of cnox1Am
(cg13424) was studied. Characterisation of the Tlx/Hox11-related genes cnox1Am and cg13424 is described in sections 5.2.1 – 5.2.4 while characterisation of the Bar ortholog barhAm is presented in sections 5.2.6 – 5.2.9.

4.2 Results

4.2.1 Isolation and preliminary characterisation of cnox1Am and cg13424

4.2.1.1 The cnox1Am cDNA

The cnox1Am cDNA clone was isolated using a portion of the ORF generated by PCR from a genomic clone isolated by Dr Julian Catmull. The complete cDNA was obtained from a λZAP-II Acropora millepora cDNA library following a number of repeat screens of approximately 5 x 10^5 plaques each, from a number of different libraries. This ultimately led to the identification of a single cDNA clone from the post-settlement library of ~1kb comprising 30bp of 5' untranslated region, 666bp of open reading frame and 363bp of 3' untranslated region (Figure 4.1). Conceptual translation of the open reading frame of cnox1Am results in a putative protein, CnoxlAm of 222 amino acids. Translation initiation is predicted to begin at base pair 31 of the cDNA clone – while no in-frame upstream termination codons in the cDNA sequence, initiation was deduced to most likely to begin from this codon from comparison with a genomic clone containing cnox1Am, which contains the first of a series of in-frame stop codons at 42bp upstream of the putative translation start site. The homeodomain begins 89 amino acid residues from the N-terminus of the Cnoxl-Am protein. A motif similar to the Eh1 domain found in other homeoproteins (Smith and Jaynes, 1996) is present 59 amino acid residues N-terminal of the HD (see Figs 4.1 and 3.8).

4.2.1.2 The CnoxlAm protein - Assignment to the Hox11/Tlx family using phylogenetic analysis

The homeodomain of CnoxlAm is located at approximately the midpoint of the protein at amino acid residues 90 – 149 and is related to the vertebrate Tlx/Hox11 and the Mnx
homeodomain families as well as to Drosophila CG13424 and its orthologs (see Fig 4.2). Relationships between these homeodomain families are complex and have not previously been investigated. Although the invertebrate CG13424-related sequences are clearly related to the Hox11-family, these latter are distinguished by the presence of four highly conserved motifs designated TH1 – TH4 (Cheng and Mak, 1993), which function in transcriptional activation (Zhang et al., 1996). These regions are not present in CnoxlAm, however, Eh1 motifs (see Fig 3.8) are a common feature of CnoxlAm, the CG13424-like, Hox 11 and Mnx family proteins.

To better understand the evolutionary position of CnoxlAm and CG13424, maximum likelihood phylogenetic analyses were carried out on the homeodomains using MolPhy Version 2.3 (Adachi and Hasegawa, 1996). Phylogenetic analyses (Fig 4.2A) implied that CnoxlAm is likely to be orthologous to the CG13424-like proteins which have been identified from *Drosophila*, the honey bee (*Apis mellifera*) and the sea squirt (*Ciona intestinalis*). This group forms a clade that is closely related to, but distinct from, the Tlx/Hox11-like proteins. Although related, the Mnx family were clearly distinct from CG13424 and Tlx/Hox11-types (Fig 4.2A).

4.2.1.3 The cnoxlAm and cg13424 genomic loci

Comparison of genomic sequence to the *cnoxlAm* cDNA sequence revealed the presence of a single intron of 933bp located 29bp 5' of the homebox (Fig 4.1). In comparison, the *Drosophila cg13424* locus contains two introns - one (891bp) located 17bp 5' of the homebox, and another (156bp) between nucleotides 132 and 133 of the homeobox (corresponding to amino acid positions 44 and 45 of the homeodomain). Virtual northern analysis indicated the size of the *cnoxlAm* transcript to be approximately 1kb (data not shown), which is similar to the size of the *cnoxlAm* cDNA clone. This implies that the transcription start site must be proximal (within 100-200bp) – a size discrepancy that could be accounted for by the inaccuracy of agarose gels) to the first exon. TATA and CAAT boxes located at −68 and −161bp respectively fit the spacing requirements for a start site located at approximately −60bp.
Figure 4.1: The cnoxlAm cDNA. In both (A) and (B) red shading represents the homeodomain and yellow shading the Eh1 motif. (A) Schematic representation of the cnoxl-Am. The start (ATG) and stop (TGA) codons are shown at the top. (B) The nucleotide sequence of the cnoxlAm cDNA and the predicted amino acid sequence of the protein.
Figure 4.2: Comparison of the Acropora Cnox1Am protein to other related proteins. (A) Cnox1Am and related homeodomains were analysed by Maximum-Likelihood phylogenetic analysis in MolPhy version 2.3 (Adachi and Hasegawa, 1996) using the Dayhoff model of protein evolution and local rearrangement of the NJ trees. Numbers against branches indicate the percentage of 1000 bootstrap replicates supporting topology. The asterisk indicates the Cnox1Am protein. The Msx3-Am homeodomain from Acropora served as an outgroup. (B) A Boxshade alignment of the homeodomain sequences used in the phylogenetic analyses. Identical residues are shaded black and conserved substitutions are shaded grey. The column to the right of the alignment indicates the overall identity and similarity of each protein with the Cnox1Am homeodomain. The species name and GenBank Accession number of each protein used are as follows: Drosophila melanogaster cg13424 (NP_611491), Hox11 (Z22959), engrailed (P02836); Apis mellifera cg13424-like (XP_393940); Ciona intestinalis (sea squirt) cg13424-like (AK174982); Gallus gallus Tlx1 (O93366), HB9 (AAC64925); Mus musculus (mouse) Tlx1 (P43345), Tlx3 (CAI25153), Hox11 (P43345); Human Tlx1 (NP_05512); Danio rerio (zebrafish) Hox11 (NP_705937), Tlx3b (NP_739572), HB9 (AAS07018), Mnr2a (AY445045); Branchiostoma floridae (amphioxus) Mnx (AAG33015); Discocelis tigrina (planarian) H6 (AJ300664); Caenorhabditis elegans (Nematode) ceh19 (NM_068928).
While upstream promoter sequences controlling gene transcription are generally located within approximately 100bp of the transcription start site (Lewin, 1997), without additional evidence from functional studies it is difficult to ascertain the regions responsible for the regulation of a gene. In closely related species, the regulation of a developmentally important gene is likely to have been conserved. In this way it is possible to identify putative regulatory regions by comparing the non-coding sequence upstream of genes, or their introns in related species and determining if any stretches of sequence are conserved. These methods have been remarkably effective in identifying mammalian regulatory sequences (Pennacchio and Rubin, 2001) and preliminary studies in *Drosophila* suggest their usefulness will extend to the study of insect regulatory sequences (Berman et al., 2004; Sinha et al., 2004). A number of transcription factor databases, such as MatInspector 2.2 and TFSEARCH are available to examine potential transcription factor binding sites for a given sequence, and programs such as BLASTalign allows for pairwise alignment of two nucleotide sequences to identify conserved segments. *D. melanogaster* cg13424 is located only 4774bp downstream of another gene, cg33008. For this reason, 4700bp of the *Drosophila melanogaster* cg13424 putative promoter sequence was compared with the equivalent region of *Drosophila pseudoobscura*. In addition, the first introns in the cg13424 loci were compared between these species. BLASTalign analysis revealed a number of short stretches of DNA, ranging from 33 to 122bp with 80% - 95% similarity - eight in the 5' UTR and four in the first intron. TRANSFAC® 6.0 (Heinemeyer et al., 1998) analyses of the arthropod database revealed that all but one of these stretches contained binding sites for a number of transcription factors, and binding sites for chorion factor 2 (CF2-II) heat shock factor (HSF), deformed (Dfd) and broad complex (BR-C) were conserved in equivalent positions between species (see Fig 4.3).

4.2.1.4 Spatial expression patterns of cnox1Am and cg13424

The spatial expression patterns of both *cnox1Am* and *Drosophila* cg13424 were analysed by in situ hybridisation (*Acropora, Drosophila*), double in situ hybridisation (*Drosophila*) and in situ hybridisation/22c10 antibody staining (*Drosophila*).
Figure 4.3: Transcription factor binding sites in the conserved between the first intron and putative promoter region of *D. melanogaster* and *D. pseudoobscura* cg13424 genes. (A) Conserved transcription factor binding sites in the first intron of the cg13424 gene of both species. The relevant species is shown at the left of the sequence, and numbers refer to the position from the first base of the intron. (B) Conserved transcription factor binding sites in the 4700bp putative promoter sequence of the cg13424 gene from both species. Numbering refers to the base position relative to the start codon. Blue boxes represent Broad Complex (BR-C) binding sites, green Deformed (Dfd) binding sites, red Heat Shock Factor (HSF) binding sites and yellow Chorion factor (CF2-II) binding sites. Overlapping sites are shown by dotted lines.
A digoxigenin (DIG)-labelled riboprobe generated from linearised plasmid containing the complete cnoxlAm was used in the Acropora in situ experiments. Only post-settlement stages of Acropora were used for in situ experiments, on the basis of the virtual northern results (data not shown). In situ hybridisation showed cnoxlAm to be expressed in a spatially restricted pattern, although staining did not appear to be cell type specific. Occasionally, Acropora embryos appear mushroom shaped (see Fig 4.4A(i) – (iv)) although it is not known which stage of development this represents. It was initially thought this stage represented the intermediate progression from pre-settlement planulae to post-settlement embryos as the planula shorten and thicken along their oral-aboral axis. However, ‘mushroom’ embryos are not seen frequently enough in developing embryo collections to be known with certainty to represent an intermediate stage (personal observations). It is now thought that this may be an abnormality of certain embryos around the time of settlement. In mushroom-shaped embryos, cnoxlAm expression is restricted to the ectoderm covering the entire oral surface, and is completely absent from the aboral ‘stalk’ region (Fig 4.4A(i) – (iv)). In the, flattened disc post-settlement stage cnoxlAm continues to be expressed in the ectoderm of the entire oral surface but not in the basal surface ectoderm (Fig 4.4A(v)a – (v)b); this is most apparent in transverse sections (Fig 4.4B (i) – (iii)).

In situ hybridisation experiments were also conducted on Drosophila embryos using a cg13424 riboprobe. The complete cg13424 cDNA was obtained from ResGen™ (Invitrogen Corporation) via the Flybase website (http://flybase.bio.indiana.edu) and was sequenced prior to manipulation. A 1kb portion (base 21 to 1020) of the cg13424 ORF was PCR amplified and subcloned into pGEM-T. The cg13424/pGEM-T clone was then linearised to act as template for production of DIG-labelled and Fluorescein(FL)-labelled riboprobes. In situ hybridisation was performed using the DIG-labelled cg13424 riboprobe and NBT/BCIP as a substrate.

In order to facilitate interpretation of the CG13424 expression patterns, double in situs were also attempted using a combination of either DIG-labelled cg13424 riboprobe with FL-labelled even skipped riboprobe or FL-labelled cg13424 riboprobe with DIG-labelled even skipped or DIG-labelled bagpipe riboprobes, kindly provided by Dr David Hayward and Dr Masha Smallhorn respectively. Primary antibodies conjugated to different enzymes were also used; either anti-FL antibody conjugated to alkaline
Figure 4.4: Spatial expression pattern of *cnox1Am* in post-settlement polyps. *In situ* hybridisation pattern observed in post-settlement polyps when hybridised with a DIG-labelled *cnox1Am* probe. Asterisks mark the oral pore and arrows mark the limits of *cnox1Am* expression. 4.4(A)(i) - (iv) show expression in the mushroom shaped polyps which may be an abnormality of development (see section 4.2.1.4) but which clearly show expression in the oral surface. 4.4(v)a and (v)b show expression in the oral side and aboral side respectively. B(i), (ii) and (iii) show expression in cross-sections which show the expression in the oral ectoderm, which is absent from the aboral ectoderm.
phosphatase (AP), or an anti-DIG antibody conjugated to AP or horseradish peroxidase (HRP). Substrates used to detect message were Vector Red (Vector Laboratories) or NBT/BCIP (Vector Laboratories) used in reactions catalysed by AP, and DAB used in reactions catalysed by HRP. In addition, direct visualisation of the fluorescein riboprobe after hybridisation was attempted, without the use of an antibody or colour substrate. Unfortunately, no combination was found which allowed for the visualisation of both cg13424 with either even skipped or bagpipe (results not shown).

Further attempts to relate the expression of the cg13424 transcript to known tissues involved in situ/antibody double staining experiments performed using DIG-labelled cg13424 riboprobe and the 22c10 antibody, which stains neurons of the peripheral nervous system (Zipursky et al., 1984). For these experiments, Drosophila embryos that had been fixed in paraformaldehyde for 12 – 15 min (‘short fix’) were used, as the longer fixation times (30-60 mins; ‘long fix’) generally used for in situ hybridisation lead to loss of the epitopes recognised by the 22c10 antibody. Successful in situ/antibody double staining was achieved following overnight hybridisation of DIG-labelled cg13424 riboprobe followed by incubation of the embryos with both antiDIG-AP and 22c10 primary antibodies simultaneously (see section 2.2.2.2). Colours were developed using first the NBT/BCIP substrate to detect the cg13424 signal, followed by addition of goat anti-mouse-HRP secondary antibody and finally detection of 22c10 protein with the use of DAB as a substrate.

The results of the in situ hybridisation and in situ hybridisation/22c10 Ab double staining experiments are shown as Figs 4.5 and 4.6. The in situ data imply that expression is initiated around stage 9 in the maxillary bud (Fig 4.5A). By stage 12, expression can be seen in the thoracic segments T1, T2 and T3 (Fig 4.5C); the expression domain continues to expand towards the posterior in stages 14 and 15 (Fig 4.5D-E), until in stage 16 all segments show staining in the body wall (Fig 4.5F). The double in situ/22c10 antibody results clearly shows that the cells expressing CG13424 lie deeper than the peripheral nervous system, which is in the surface ectoderm of the Drosophila body wall. Axons and sensory cells of the peripheral nervous system are clearly seen to pass over the top of the cg13424 stained cells (See Fig 4.6). The three obvious cg13424 stained cells (See Fig 4.6(A)) are located in a position where there are
Figure 4.5: Spatial expression pattern of \textit{cg13424} at various stages of \textit{Drosophila} development. The stages are noted to the left of each series row. The first column of each shows a lateral view, the middle column a dorsal view and the far right column a ventral view. Expression initiates at stage 9 in the maxillary bud and extends to the thoracic segments (T1, T2, T3) by stage 12. In later stages (14 - 16), expression is seen in the lateral transverse muscles of the body wall.
Figure 4.6: Coexpression of cg13424 and 22c10 in the body wall of late stage *Drosophila* embryos. In each, cg13424 cells are stained with NBT/BCIP (purple/blue), while 22c10 is stained with DAB (light brown). Fig 4.6A(ii) is an enlarged version of A(i) as B(ii) is an enlarged version of B(i). The three arrowheads in A(ii) show the three characteristically shaped cg13424-expressing cells. The arrow in B(ii) shows the axon of the 22c10 stained peripheral nerves passing over the cg13424-expressing cells.
no sense organs, i.e. in abdominal segments A1-7 between the 1cg5 and dbd neurons. The position and alignment of cells expressing cg13424 are reminiscent of the lateral muscle group which consists of three muscles that run dorso-ventrally – the lateral transverse muscles (Landgraf et al., 1997). The above results imply that cg13424 is specifically expressed at the dorsal attachment point of these muscles.

RNA interference (RNAi) was proposed in order to determine the knockout phenotype of Drosophila cg13424 null mutants. Initiation of this work is presented as Appendix C, and is ongoing.

4.2.2 BarhAm – a second cnidarian member of the Bar class

4.2.2.1 The barhAm cDNA

Virtual northern analysis of barhAm indicated expression in post-settlement and adult stages, represented by a single band of ~1.5kb (data not shown). A screen of 5x10^5 clones of a λZAP-II post settlement cDNA library using the complete barhAm EST as a probe yielded four positive clones, three of which covered different regions of the same gene, together spanning ~1.4kb and thus correlating with the results obtained from virtual northern blotting. The contig derived from the three cDNA sequences contained a putative ORF of 669bp with 159bp 5’ UTR and 555bp of 3’ UTR (see Fig 4.7). Conceptual translation of the ORF of barhAm gives a putative protein BarhAm of 223AA. In the absence of confirmatory evidence, the assignment of the start codon is tentative. The sequence of the only other known cnidarian Bar family member, cnox3 from C. viridissima is incomplete, therefore it is not possible to make inferences about translation start sites etc by comparison with this gene. A number of unsuccessful attempts were made to isolate barhAm genomic clones, using the longest cDNA clones as probes. The reasons for failure are not clear; one possibility is that the coding sequence may be highly interrupted.
Figure 4.7: The *barhAm* cDNA. In both (A) and (B) red shading represents the homeodomain. (A) Schematic representation of the *barh-Am ORF*. The start (ATG) and stop (TGA) codons are shown at the top. (B) The nucleotide sequence of the *barh-Am* cDNA and the predicted amino acid sequence of the protein.
4.2.2.2 The BarhAm protein

The *Acropora barhAm* gene encodes a protein containing a *Bar* class homeodomain in the C-terminal portion of the protein, at amino acid residues 117 - 176. This homeodomain is more similar to the vertebrate Barx genes than to BarH1/2 from *Drosophila* (see Fig 4.8). As might be expected, the BarhAm homeodomain is most similar to that in Cnox3 from *Chlorohydra* (62/77% identity/similarity). Although BarhAm clearly falls into the Bar family, the homeodomain does not have the characteristic tyrosine residue at position 49, shared by all other known Bar class homeoproteins except for sponge prox2 and *Xenopus* Xbrlα (see Fig 4.8B). Outside of the homeodomain BarhAm does not share any of the features that are present in some other members of the Bar class. For example BarH1 and BarH2 (*Drosophila*), Barh1 (mouse), XBH1 and XBH2 (*Xenopus*), Bar (medaka) and cBarx2b (chick) each contain single Eh1 motifs upstream of the homeodomain; MBH1 (rat) contains two such motifs. Mouse Barx1 and Barx2 both contain a basic region of 17 amino acids C-terminal of the homeodomain (Barx basic region/BBR) (Edelman et al., 2000), and Barx2 also possesses a leucine zipper, polyalanine tract and an acidic domain (Jones et al., 1997). Both rat MBH1 and *Drosophila* BarH1/2 proteins contain strings of alanine, glutamine and histidine residues (Higashijima, 1992; Saito et al., 1998).

4.2.2.3 Evolutionary relationships amongst the Bar class of homeodomains

To better understand the evolutionary position of BarhAm, maximum likelihood phylogenetic analyses were carried out using MolPhy Version 2.3 (Adachi and Hasegawa, 1996) based on the homeodomain sequence. Phylogenetic analysis separates most members of the Bar class into two major clades – one containing proteins most similar to the Barx class, and the other containing proteins more similar to BarH from *Drosophila*. The two cnidarian sequences (BarhAm and Cnox3) form a well supported clade that is more closely related to the Barx than to the Barh clades. The length of the BarhAm branch suggests that the protein is more derived than its *Chlorohydra* counterpart, which is unusual; in general, *Acropora* proteins are less
Figure 4.8: Comparison of the Acropora BarhAm protein to other Bar-related proteins. (A) BarhAm and related homeodomains were analysed by Maximum-Likelihood phylogenetic analysis in MolPhy version 2.3 (Adachi and Hasegawa, 1996) using the Dayhoff model of protein evolution and local rearrangement of the NJ trees. Numbers against branches indicate the percentage of 1000 bootstrap replicates supporting topology. The asterisk indicates the BarhAm protein. The CG13424 (accession #NP_611491) and Vnd (CAA60619) homeodomains from Drosophila served as outgroups. (B) A Boxshade alignment of the homeodomain sequences used in the phylogenetic analyses. Identical residues are shaded black and conserved substitutions are shaded grey. The column to the right of the alignment indicates the overall identity and similarity of each protein with the BarhAm homeodomain. The black dot represents the distinguishing threonine residue at position 47 (T47) and the star the tyrosine residue (Y49) conserved in most Bar-type homeodomains. The species name and GenBank Accession number of each protein used are as follows: Mus musculus Barx1 (AAG18573), Barx2 (AAH12684), Barh1 (NP_031552); Chlorohydra viridissima cnx3 (CAA45910); Rattus norvegicus MBH1 (O88181); Xenopus laevis XBH1 (AAG14450), XBH2 (AAG14451), Xbrl1 (AAB03566); Drosophila melanogaster Barh1 (Q24255), Barh2 (Q24256); Anopheles gambiae Barl (XM_320094); Caenorhabditis elegans ceh30 (NP_508524).
A

B

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Figure 4.9: Spatial expression patterns of *barhAm* in pre-settlement and post-settlement stages. Asterisks mark the position of the oral pore in each plate. A(i) and (ii) show expression in a whole pre-settlement larvae and a cross-section of a similarly aged larvae respectively. B shows staining in a cross-section of a 'mushroom' stage polyp, while C-D show expression in post-settlement polyps. C(i) and (ii) are polyps of the same age showing expression exclusively around the oral pore, while D(i) - (iii) are cross-sections of polyps of the same age as those shown in C. Expression is restricted to the ectoderm around the oral pore, although 'bleeding' of stain can make the expression appear more extensive (as indicated by arrowheads).
derived than are their orthologs from hydrozoan cnidarians. Within the Barh clade, the arthropod and vertebrate sequences are clearly resolved (see Fig 4.8A).

4.2.2.4 Spatial expression pattern of barhAm

The spatial expression pattern of barhAm was analysed by in situ hybridisation. A digoxigenin (DIG)-labelled riboprobe was generated from linearised plasmid containing the complete barhAm cDNA. While virtual northern results indicated expression only post-settlement (data not shown), in situ hybridisations were conducted on both pre- and post-settlement material, as certain collections contained representatives of both stages. Surprisingly, expression in pre-settlement was seen, in apparent contradiction to the virtual northern results.

Expression in pre-settlement larvae is seen in the ectoderm immediately surrounding the oral pore (Fig 4.9A(i)-(ii)). In sections of 'mushroom' stage post-settlement material, expression is restricted to the ectoderm at the cup-like oral pore (Fig 4.9B), and this specific pattern persists as the coral settles and flattens, (Figs 4.9C (i)-(ii), D(i)-(iii)). Virtual northern analysis indicates that barhAm is also expression in the adult colony, albeit at lower levels; unfortunately, the calcified skeleton greatly complicates the application of in situ hybridisation methods on adult.

4.3 Discussion

4.3.1 Structural features of the Cnox1Am and BarhAm proteins

The Bar and Hox11/Tlx families are typical members of the Antennapedia superfamily in that they contain a glutamine residue at position 50 in the third helix of the homeodomain (Q50) which results in preferential binding at sites containing the core sequence CATTA (Scott et al., 1989; Kappen et al., 1993). However, these same families are atypical in that they are the only known homeodomains with a threonine residue at position 47 (T47) within helix3 – in other homeodomains, isoleucine or
valine is most commonly found in this position, but asparagine, leucine or histidine are
also known (Laughon, 1991; Dear et al., 1993). Position 47 is one of the four positions
in helix 3 known to confer DNA binding specificity (Burglin, 1994), proven to be the
case for the HOX11 HD in vitro (Dear et al., 1993); most likely the specificity
determined by the threonine in this position extends to roles in vivo. A further
characteristic of Bar class homeodomains is the tyrosine residue at position 49 (Y49)
replacing the near-universal phenylalanine residue. It has been suggested that together,
these two positions (T47 and Y49) could play a role in influencing DNA-binding after a
posttranslational modification, such as phosphorylation changes the protein
conformation (Jones et al., 1997). As the Acropora BarhAm homeodomain has F49
rather than Y49 residue, this raises the question of whether this is an ancestral or a
derived feature. The fact that Chlorohydra Cnox3 HD has Y49 suggests that Acropora
may contain a second Bar class protein with the F49/Y49 substitution. This is also
indicated by the derived nature of BarhAm in comparison to Cnox3, shown by the
length of the BarhAm branch in the maximum likelihood analysis (see Fig 4.8).

With the exception of the Eh1 motif (see sections 4.2.1.2 and 4.2.2.2), the Bar and
Hox11 proteins exhibit no significant homology outside of the homeodomain. All Barx
family members contain an Arg/Lys-rich region N-terminal of the HD, a feature shared
by some members of the NK2 family. This region appears to be required for serum
response factor (SRF) binding (Chen and Schwartz, 1995), and a direct interaction
between SRF and Barx2b has been demonstrated (Herring et al., 2001). The vertebrate
members of this class contain additional motifs which may reflect the need for
increased complexity of action of Bar proteins in vertebrates, relative to their activity in
'simpler' animals such as cnidarians and Drosophila.

Phylogenetic analyses of Cnox1Am and BarhAm places both homeodomains in specific
classes – Cnox1Am as a member of the Hox11/Tlx class and BarhAm as a member of
the Barx subfamily of the Bar class. It is interesting to note that there are no clear
vertebrate orthologs of Drosophila cg13424 in the database. The presence of an
ascidian ortholog implies that loss of members of this gene family has occurred in
vertebrate species. No cnidarian homologs of the BarH subfamily of the Bar class of
genes have been isolated, although the possibility of cnidarians possessing this
subfamily cannot be discounted. Vertebrates contain members of both subfamilies,
while *Drosophila* has only those of the BarH type. Clearly both the BarH and Barx subfamilies were present in the last common ancestor of higher metazoans, and that the latter has been lost in the arthropod lineage, but maintained in the vertebrate lineage. Alternately if cnidarians were conclusively shown to possess only Barx-type genes, it could be hypothesised that while the last common ancestor had both, it is possible that the Barx-type were ancestral, and that the BarH subclass originated after the cnidarian-bilaterian split.

### 4.3.2 Expression patterns of cnox1Am, cg13424 and barhAm

The *Hox11/Tlx* and *Bar* genes are both expressed in the nervous system in all organisms studied thus far, although each also have diverse roles in other tissues. On the basis of apparent conservation of function in the nervous system, these genes were characterised in *Acropora* in the expectation that they would also play roles in nervous system development or patterning in cnidarians. However, both *cnox1Am* and *barhAm* were more generally expressed in the ectoderm around the time of settlement. While expression in the nerve net cannot be discounted, it is apparent that expression of neither gene is limited to any one cell type; rather, they are both uniformly expressed in spatially restricted patterns. *BarhAm* expression is seen as a single band in the ectodermal layer surrounding the oral pore and is initiated at settlement. During this time, the nervous system of *Acropora* is completely remodelled (E Ball, pers comm.), as has also been described in hydroids (Martin, 2000). Settlement initially involves the receipt of settlement cues which induce the planula larvae to attach to the substratum at its aboral end and contract along the oral-aboral axis to form a flattened disc, giving the characteristic post settlement morphology. It is possible that *cnox1Am* might function in regulation of this process, as it is exclusively expressed in the ectoderm on the oral surface of the embryo, from approximately the time of settlement. The expression pattern of the Chlorohydra gene *cnox3* is similarly restricted to the head region, and it has been suggested that this gene plays a role in head regeneration (Schummer et al., 1992). The role of *barhAm* in *Acropora* is unclear; it can be speculated that its restricted expression pattern might reflect a role in specifying a specific cell type(s) to ensure the correct functioning of the oral pore.
The expression pattern of cg13424 was investigated in Drosophila embryos in an attempt to determine the function of this uncharacterised gene. Here, the expression pattern was very specific, in what are thought to be the lateral transverse muscles of the body wall, based also on the morphology of the cg13424-specific cell staining. The fly gene was not expressed in the nervous system. Thus, although clearly related in sequence, the fly and coral genes have completely different expression patterns, implying unrelated functions. However, there are many known examples of Drosophila genes having diverged function (eg (Strecker et al., 1986)), and it would be of interest to determine the expression patterns of cg13424-related genes in a wider variety of animals. The phylogenetic analyses presented here indicate that cg13424 and cnoxlAm define a family of homeobox genes that are clearly related to, but distinct from the Hox11/Tlx type; from this preliminary characterisation their functions do not appear to be related.

4.3.3 Regulation of cnoxlAm and cg13424

Recently, a second Drosophila genome (Drosophila pseudoobscura) was sequenced by the National Human Genome Research Institute (NHGRI) to aid in the identification of conserved regions, and guide the annotation of functional sequences in D. melanogaster (Berman et al., 2004). A secondary use for the data was in identifying conserved regulatory regions of genes. Although conservation of DNA sequence cannot always accurately determine functional regulatory sequences, conservation of binding-site clustering between D. melanogaster and D. pseudoobscura has been used to discriminate functional binding-site clusters from those with no function, as demonstrated by several recent studies (Bergman et al., 2002; Berman et al., 2004; Grad et al., 2004; Sinha et al., 2004). For this reason, a comparison of the 5’ upstream sequence and first intron of both D. melanogaster and D. pseudoobscura cg13424 genes was compared. It was found from a search of the 4.7kb 5’UTR and first intron (see section 4.2.1.3), only four of the binding sites were shared between the two species; those for heat shock factor (HSF), deformed (Dfd), Chorion factor 2 (CF2-II) and broad (Br;BR-C) (Fig 4.3). For any of these proteins to play a role in the regulation of cg13424, it would be necessary for these genes to be expressed in analogous positions, i.e. the maxillary bud in early stages and in the lateral transverse muscles in later stages.
Based on published expression data alone, it appears that of these four transcription factors Dfd and CF2-II are possible regulators of cg13424.

The binding of HSF homotrimers to heat shock elements (HSEs) is cooperative, but requires close proximity of HSEs, and a minimal complete binding site for HSF is made up of three contiguous units (Fernandes et al., 1995). Results of the TF search of the 4.7 kb promoter region and first intron of D. melanogaster and D. psuedoobscura cg13424 genes do not identify more than two contiguous HSEs indicating that none of the HSF binding sites identified can operate as a functional HSF promoter element. Br-C (Broad Complex) is essential for the progression of larave through metamorphosis, and those which lack BR-C die without pupriating (Stewart et al., 1972; Kiss et al., 1978). It is expressed in numerous tissues including the forming CNS, the developing eye and is required for the attachment of the thoracic muscles to the body wall (DiBello et al., 1991; Emery et al., 1994; Sandstrom et al., 1997; Brennan et al., 1998). However, while extensive, the expression of BR-C does not seem to coincide with CG13424 and can thus be discounted as a possible regulator. Dfd is a Hox gene in the antennapedia complex expressed in the eye-antennal and labial imaginal discs, and also in the mandibular and maxillary segments of the head (see Hughes and Kaufman, 2002); it is therefore possible Dfd plays a role in regulating cg13424 in the maxillary bud, but regulation in the muscles of the body wall at later stages is not possible. CF2-II however is expressed in the developing muscles of the embryo where it first appears at stage 12; later it is expressed in all muscle lineages including skeletal, visceral and cardiac, and is in the nuclei of all skeletal muscles (Bagni et al., 2002). Thus, the combinatorial actions of both Dfd and CF2-II might specify the region of cg13424 expression in early and late embryonic stages respectively. Interestingly, in Drosophila, CF2-II expression is dependant on the highly conserved MADS-box gene MEF2; a MEF2 homolog has recently been isolated from the hydrozoan jellyfish Podocoryne carnea suggesting the possibility of a conserved muscle patterning cascade (Bagni et al., 2002; Spring et al., 2002).

In orthologous genes, intron positions are often conserved. However, Drosophila cg13424 contains two introns, whereas Acropora cnox1Am contains only one. The first in both of these genes is located at seemingly equivalent positions, located 29bp 5' of the homeobox in Acropora, and 17bp 5' of the homeobox in Drosophila while the
second *Drosophila* intron is located between amino acids 44/45 of the homeodomain. A plausible evolutionary scenario is that the ancestral *cnox1/cg13424*-type gene contained an intron close to the 5' end of the homeobox, and that the additional intron that is present in the arthropod lineage has accumulated during evolution. Alternately, it may be that secondary loss is responsible for the reduced intron density in *Acropora*; further sampling of *cnox1/cg13424* genes from other organisms will help to resolve this issue.

### 4.4 Conclusions

In this Chapter, the isolation and expression patterns of two coral genes, a Barx-type gene (*barhAm*) and Hox11/Tlx-related gene (*cnox1Am*), and the preliminary characterisation of a *Drosophila* gene related to the latter are described. Like most members of the corresponding protein families, threonine residues are present at HD position 47 in both BarhAm and Cnox1Am. However in the BarhAm HD F49 is present, rather than Y49 as in most other Bar class proteins. There is no significant conservation outside of the homeobox in either BarhAm or CnoxlAm, with the exception of an Eh1 motif N-terminal of the homeodomain in CnoxlAm which is also present in all members of the Hox11/Tlx family.

Both *barhAm* and *cnox1Am* are expressed around the time of settlement; *cnox1Am* is only present in post-settlement and adult stages, while *barhAm* expression appears to initiate immediately prior to the settlement stage. Both genes are expressed in the ectoderm in spatially restricted patterns. While vertebrate homologs of both *barhAm* and *cnox1Am* are expressed in the nervous system during development, the expression of the *Acropora* genes is not cell type restricted. The *Drosophila* homolog of *cnox1Am*, *cg13424*, is expressed from approximately stage 9, and is restricted to the lateral transverse muscles of the body wall in later stages (14 – 16). Analysis of the promoter region and first intron revealed that the Dfd and CF2-II proteins are potential regulators of *cg13424*. 
4.5. Future Directions

The work described here on barhAm, cnox1Am and cg13424 suggests several lines of further investigation. While Acropora lacks obvious muscle, many cnidarians (eg Podocoyne) have well developed striated and smooth muscles that are derived from a mesoderm-like structure, the entocodon (Spring et al., 2002). Hence it may be informative to characterise cnox1Am-like genes from cnidarians with well-defined muscles, as well as cg13424 family members from other “higher” animals. In terms of understanding the function of the Drosophila gene cg13424, it is important that loss of function phenotypes are examined, and the most promising way of addressing this is via RNAi expression under control of the GAL4-UAS system. The identification of downstream targets of this range of genes can now also be attempted via the use of yeast one-hybrid technology.
Chapter 5 - DNA-Binding Characteristics of Cnidarian PaxC and PaxB Proteins In Vivo and In Vitro: No Simple Relationship With the Pax6 and Pax2/5/8 Classes

5.1. Introduction

5.1.1 Common Molecular Mechanisms of Eye Development

To a surprising extent, common molecular mechanisms appear to underlie the early morphogenesis of eyes across the animal kingdom. Classically, eyes were thought to have polyphyletic origins, and to have evolved at least 40 times independently in the various animal lineages - an idea that can be traced back to Darwin’s *Origin of Species* (Darwin, 1859; Salvini-Plawen and Mayr, 1977). However, the discovery that the *Drosophila* eyeless (ey) gene is orthologous and functionally interchangeable with the mammalian and *Xenopus* Pax6 genes (Quiring et al., 1994; Halder et al., 1995; Onuma et al., 2002) led to recognition that aspects of the molecular basis of eye specification are similar despite major morphological differences between fly and vertebrate eyes, and suggested a monophyletic origin of eyes. Subsequently, conservation not only of Pax6 function, but also of much of the genetic network of eye specification and patterning has been demonstrated between *Drosophila* and mammals (Wawersik and Maas, 2000; Kumar and Moses, 2001). In higher animals, clear Pax6 orthologs have been universally implicated in eye specification, and have been cloned from a diverse range of animals including the squid *Logilo* (Tomarev et al., 1997) and ribbon worm *Lineus* (Loosli et al., 1996).

Not only are the Pax6 proteins conserved across deep evolutionary time, so too are functional aspects of their regulatory elements. Autoregulation is a general feature of Pax6-like genes, and in the case of the eyeless gene of *Drosophila* this is mediated by the two intronic enhancer elements that are required for expression in the nervous system and eye. These enhancer elements share a significant degree of sequence identity with other *Drosophila* species and with *Anopheles gambiae* (Hauck et al., 1999; Adachi et al., 2003). Although lacking significant sequence identity in the non-coding
regions, these *Drosophila* enhancers can recapitulate the major features of mouse *Pax6* expression during development when placed in an appropriate reporter construct (Xu et al., 1999). Regulation of mouse *Pax6* is complex; transcription can start from any one of three promoters under the control of six enhancer elements - four of these regulate different aspects of *Pax6* expression in the developing eye (Williams et al., 1998; Kammandel et al., 1999; Xu et al., 1999; Griffin et al., 2002). These enhancer elements are recognisable across a broad range of non-mammalian vertebrates including quail, pufferfish and *Xenopus*, and the pufferfish enhancer elements for example, can substitute for their mouse equivalents (Kammandel et al., 1999). Identity between vertebrates and invertebrates in the non-coding regions of *Pax6* genes is limited to short motifs (Morgan, 2004) that closely match the *Pax6* consensus binding sites in all of the genes studied (*Xenopus*, Mouse, Zebrafish, Human, Ciona, *Drosophila* and *Caenorhabditis*) and these are often at equivalent positions relative to the first exon. This suggests that an ancestral control mechanism that has subsequently been ‘fine-tuned’ in different lineages (Morgan, 2004).

5.1.2 Pax Genes

The defining characteristic of Pax genes is they encode a 128 amino acid DNA binding motif, the paired-domain (PD). The term paired box (from which the name pax originates) was coined due to the fact that the motif was first recognised in the *Drosophila* segmentation gene *paired* (Bopp et al., 1986; Frigerio et al., 1986). In addition to the paired domain, many *Pax* genes also encode another DNA-binding domain, a paired-type homeodomain (HD), and/or a small octapeptide-motif which binds Groucho-class corepressors (Eberhard et al., 2000). *Pax* genes have been cloned from many representatives of a diverse range of phyla, including the cnidarian *Acropora millepora* (Catmull et al., 1998). In many cases, those *Pax* genes which contain a homeobox are expressed in the nervous system during development.

The paired domain is responsible for sequence specific DNA-binding via two helix-turn-helix subdomains, termed PAI and RED, separated by a short linker (reviewed in (Callaerts et al., 1997)). The paired domain binds as a monomer to a non-palindromic recognition sequence in the two major grooves on the one side of the DNA helix.
The PAI subdomain recognises the more extensive 5' consensus half-site motif, whereas the RED subdomain interacts with the 3' consensus sequence. Specificity of binding is achieved through the co-ordinated actions of these two half sites (Czemy et al., 1993) and is due to particular amino acids in the paired domains themselves. For example, positions 42, 44 and 47 in the N-terminal region of the paired domain, were shown to be responsible for differences in DNA binding specificity between Pax5 and Pax6 (Czemy and Busslinger, 1995). Mutation of these three specific residues in Pax6 to those of Pax5 resulted in a complete switch of the DNA-binding specificity from Pax 6 to Pax5, proving that these three residues are the primary determinants of DNA binding specificity (Czemy and Busslinger, 1995).

The second DNA-binding domain, the paired-type homeodomain, is able to form homo- and hetero- dimers on DNA, and binding of one homeodomain protein has been shown to increase the affinity of a second by up to 300-fold (Wilson et al., 1993). Most homeodomains recognise almost identical sites, and although distinct and independent from the paired domain, recent studies have indicated that the absence of the homeodomain greatly decreases the affinity of paired domain binding. This suggests that the paired domain and homeodomain must interact cooperatively to activate their targets (Jun and Desplan, 1996). Homeodomain proteins have been shown to cooperatively dimerise on palindromic binding sequences known as P2 or P3 sites on the basis of the number of nucleotides between the palindromic half sites [TAAT (N2 or N3) ATTA] (Wilson et al., 1993; Jun and Desplan, 1996). The homeodomains encoded by different Pax gene classes have characteristic preferences for either the P2 or P3 sites.

The diversity of Pax gene function contrasts sharply with their conservation at the amino acid level and is puzzling in light of their in vitro DNA-binding specificity. The solution to how Pax genes achieve their many roles may lie in the ability of the corresponding proteins to use multiple combinations of their helix-turn-helix motifs in order to recognise more targets. In this way, Pax genes are able to achieve a wide variety of functions (Jun and Desplan, 1996). Members of the Pax2/5/8 group contain an 'incomplete' homeodomain, which is unable to bind to DNA, and yet has been shown to be conserved in animals from sea urchin to humans, suggesting a non-DNA-binding function for this protein; the partial homeodomain of Pax-5 has been shown to be able to bind TATA-binding protein (TBP) and Retinoblastoma protein (Rb)
The C-terminal domain of several Pax proteins contains a proline-serine/threonine (PST)-rich region with cell-type independent transactivation activity, and an adjacent inhibitory sequence (Dorfler and Busslinger, 1996). The octapeptide motif mediates transcriptional repression by recruiting Groucho class co-repressors (Eberhard et al., 2000).

5.1.3 Specification of Mammalian and Drosophila Eyes

The development of the Drosophila eye begins in the eye-antennal imaginal disc from a single layer epithelium. Initially, the eye primordium is subdivided into dorsal and ventral (D/V) compartments via wingless (wg) which activates the Notch signalling pathway and leads to elevated Notch levels on both sides of the compartment boundary and functions to control eye growth (Dominguez and de Celis, 1998; Cavodeassi et al., 1999; Irvine and Rauskolb, 2001). The initiation of the D/V patterning mechanism also functions to set up a midline organizer and ‘firing centre’ from which the morphogenetic furrow (MF) forms, leading to the formation of photoreceptor cells (ommatidia) of the eye. The morphogenetic furrow is a coordinated wave of cell-shape regulation and patterning which moves from posterior to anterior across the eye field, initiated by a pulse of Hedgehog expression at the posterior margin, and which also requires the presence of decapentaplegic (dpp), and wg at the lateral margins which inhibits ectopic furrow initiation (Dominquez and Hafen, 1997; Borod and Heberlein, 1998; Baker, 2001). The ommatidia are placed in a precisely spaced array as the furrow moves with normal cell proliferation both anterior and posterior to the furrow dependent on the Egr/Ras pathway that is activated by wg expression (Baker and Yu, 2001; Yang and Baker, 2003). Prior to the D/V and A/P patterning mechanisms however, is the expression of a network of eye specification genes at the posterior margin of the eye disc where differentiation will later initiate (Quiring et al., 1994; Czerny et al., 1999). This complex regulatory network consists of what are known as the eye specification genes, eyeless (ey), twin of eyeless (toy), sine oculis (so), eyes absent (eya), dachshund (dac), eye gone (eyg) and optix (see Gehring and Ikeo, 1999; Wawersik and Maas, 2000; Kumar and Moses, 2001). These genes do not function in a simple linear hierarchy, but rather as a complex interwoven regulatory network (Fig 5.1); loss of function of any of these seven genes in the eye primordium results in a reduction or
deletion of the adult eye, whereas ectopic expression of any of these genes (with the exception of so) in the eye/antennal imaginal disc results in ectopic retinal development (Halder et al., 1995; Shen and Mardon, 1997; Bonini and Fortini, 1999; Czerny et al., 1999; Seimiya and Gehring, 2000). Toy, ey and eyg are expressed first, while expression of eya, so and dac begins immediately prior to morphogenetic furrow formation, although these genes also have an earlier role independent of toy and ey in the embryonic development of the whole visual system. Ey, eya, so and eyg are responsible for growth of the early eye disc, ey, eya, so, eyg and dac are required for initiation of the morphogenetic furrow, eya and so for photoreceptor differentiation, and ey for rhodopsin gene expression during pupal eye development (see (Treisman, 1999)). What genes act upstream of the network of this eye specification cascade is unclear, although it has been theorised to be a combinatorial code of HOX genes (see (Treisman, 1999)).

![Complex regulatory network of eye specification genes](image)

**Fig 5.1**: Representation of the complex regulatory network of the eye specification genes. Twin of eyeless (toy) initiates the cascade, with the remaining gene products involved in feedback regulation. (Adapted from Nordström, 2003).

The two pairs of Pax6-related genes, eyeless/twin of eyeless and eyegone/twin of eyegone, exist as paralogous pairs on *Drosophila* chromosomes 4 and 3 respectively (Flybase). Eyeless and its paralog twin of eyeless both encode canonical Pax6 products, and twin of eyeless functions upstream of eyeless by directly regulating a specific enhancer element, activating eyeless expression, which in turn activates expression of so and eya (Halder et al., 1998; Czerny et al., 1999; Hauck et al., 1999; Niimi et al., 1999). Twin of eyeless is more similar to vertebrate Pax6 proteins than eyeless with
regard to overall sequence conservation and DNA-binding function, (suggesting that *eyeless* is insect-specific), but the two genes share a similar expression pattern in the developing visual system (Czerny *et al.*, 1999). *Eyegone*, like its paralog *twin of eyegone*, encodes a Pax6-like protein with a truncated paired domain, lacking most of the N-terminus (Jun *et al.*, 1998; van Heyningen and Williamson, 2002; Jang *et al.*, 2003). The function of Toe is unclear although it has recently been reported that while Eyg is essential for eye growth, it seems dispensable for eye specification, thus there may be functional redundancy between Eyg and Toe with respect to the latter (Jang *et al.*, 2003; Dominguez *et al.*, 2004). In addition, it has been shown that Notch signalling in the eye acts through Eyg and, while overexpression of Eyg can fully rescue Notch mutants, Toe overexpression results only in a partial rescue (Dominguez *et al.*, 2004). Eyg binds to a consensus site through the C-terminal (RED) subdomain of the paired domain which corresponds to the consensus binding site for a mammalian Pax6 isoform PAX6(5a) (Epstein *et al.*, 1994; Jun *et al.*, 1998). In human and mice, the PAX6(5a) isoform is generated by alternative splicing of the PAX6 primary transcript, and overexpression of human PAX6(5a) induces strong overgrowth *in vivo* whereas the canonical PAX6 variant hardly effects growth showing that these two isoforms are required for different functions – PAX6 for differentiation of the eye, and PAX6(5a) for growth (Dominguez *et al.*, 2004). Like Eyg, PAX6(5a) recognises DNA exclusively through the RED region of its paired domain (Jun *et al.*, 1998). Therefore, the PAX6 and its fly counterparts, Ey and Toy, act principally to specify eye fate whereas the PAX6(5a) isoform and its functional conterpart in the fly Eyg (and perhaps also Toe) function in growth of the eye. PAX6(5a) can functional substitute for Eyg in *Drosophila*, as can PAX6 for ey (Dominguez *et al.*, 2004).

5.1.4 Cnidarian Pax Genes

The evolutionary origins of *Pax* genes are unclear; no *Pax* genes have yet been identified outside of the Metazoa, thus there are no obvious 'outgroups' for phylogenetic analyses. In addition, alternative splicing and multiple roles during development complicate the identification of ancestral functions. As the Cnidaria are the simplest animals at the tissue level of organisation, considerable attention has
focused on defining their Pax gene complement and expression patterns in the hope
that this might lead to insights into ancestral functions and Pax gene evolution.

The first Pax genes identified in Cnidarians were PaxA and PaxB from the sea nettle
Chrysaora quinquecirrhha (a scyphozoan) and Hydra littoralis (a hydrozoan; (Sun et al.,
1997). Phylogenetic analyses and preliminary DNA binding assays demonstrated that
three of the cnidarian paired domains (sea nettle PaxA and -B and Hydra PaxB) bound
to Pax5/6 sites and implied that both the cnidarian PaxA and PaxB genes were more
closely related to the Pax2/5/8 and Pax6 classes rather than to Pax1/9 and Pax3/7.
Thus, it was proposed that modern Pax2/5/8 and Pax6 genes evolved from an ancestral
gene similar to cnidarian PaxB (Sun et al., 1997). To date, PaxA and/or PaxB genes
have been identified in a broad range of medusazoan cnidarians, including the
hydrozoans Podocoryne (PaxB) (Groger et al., 2000), Polyorchis penicillatus (PaxB)
(Nordstrom, 2003) the scyphozoans Aurelia aurita (PaxA) (Nordstrom, 2003),
Chrysaora (PaxB) and Cladonema (PaxA/B) (Sun et al., 1997; Sun et al., 2001) and the
cubozoan Chiropsalmus sp (PaxB) (Nordstrom, 2003).

Our laboratory has identified four Pax genes from the anthozoan cnidarian Acropora
millepora, PaxAam (am = Acropora millepora), PaxBam, PaxCam and PaxDam, with
PaxCam and PaxDam apparently unique to Acropora. One of the two initial genes
discovered, PaxAam is clearly homologous to the Hydra and sea nettle PaxA genes,
whereas the other, PaxCam appears to be relatively distantly related to the cnidarian
PaxB genes (Catmull et al., 1998). The PaxAam and PaxCam paired domains most
resemble those of the vertebrate Pax2/5/8 class, although the homeodomain of PaxCam
is somewhat more closely related to the Pax6 type than are the cnidarian PaxB
homeodomains, suggesting that PaxC rather than PaxB might correspond to a Pax-6
precursor. This hypothesis is supported by the fact that the homeodomain of Hydra
PaxB has only 55% identity with mammalian Pax6 (compared to 70% identity of
PaxCam with Pax6) and PaxB proteins typically contain an octapeptide motif, whereas
true Pax6 proteins do not. The fact that the PaxAam and PaxCam proteins share several
distinctive substitutions in their paired domains led to the suggestion of a common
origin via a duplication event (Catmull et al., 1998). Subsequently two more Pax genes
were cloned from Acropora; PaxBam is orthologous to PaxB isolated in other
Cnidarians, whereas PaxDam (like PaxCam) appears to be unique to Acropora (Miller,
PaxDam clearly belongs to the Pax3/7 class, while PaxBam appears to be an ancestral Pax2/5/8 gene although it is shown in this work that the PaxBam protein can bind to Pax6 binding sites (Miller, 2000). Of the four Acropora Pax genes, three (PaxBam, -Cam and -Dam) encode complete homeodomains and PaxBam also contains an octapeptide motif (typical of the Pax2/5/8 class). The PaxAam protein contains neither homeodomain or octapeptide, suggesting that PaxAam may be related to Drosophila pox neuro (Miller, 1999; Miller, 2000). The possession of common splice sites has frequently been used to support common ancestry, and the fact that several splice sites are shared between the Cnidarian and triploblastic metazoan Pax genes supports the monophyly of the Pax gene family (Miller, 2000).

5.1.5 Eyes in Cnidaria

Despite the absence of a central nervous system with which to process images, distinct eyes ranging in complexity from simple eye-spots to complex lens eyes are present in many representatives of three of the four cnidarian classes and photosensitivity is considered to be a general property of the phylum. The most sophisticated eyes are present in the most motile of cnidarians, the cubozoans (box jellyfish). The adult cubozoan eye may consist of up to 11,000 sensory cells and has an epidermal cornea, spherical lens and a retina with distinct sensory, pigmented nuclear layers (Brusca and Brusca, 1990). In contrast, the larvae of Tripedalia cystophora display photosensitivity in the complete absence of a nervous system; the ocelli which act as photoreceptors have no neural connections to any other cells, but each has a well-developed motor cilium (Nordstrom et al., 2003b). Some cnidarians, including Hydra, lack any obvious photoreceptors but clearly react to light (Tardent and Frei, 1969) and generally cnidarian larvae respond to light (Svane and Dolmer, 1995). Members of the cnidarian class Anthozoa (of which Acropora is a member) lack the motile medusa stage that is characteristic of the other cnidarian classes and also lack eyes, however they do display photosensitive behaviour. For example, coral polyps are extended at night and retracted during the day, and coral larvae display a variety of phototactic behaviours (reviewed in Harrison and Wallace, 1990). As previously mentioned Acropora remains the only cnidarian species in which PaxC and PaxD genes have been identified despite considerable efforts by several laboratories (eg Sun et al., 1997; Groger et al., 2000;
Sun et al., 2001; Nordstrom, 2003) suggesting that either these genes have been lost in other lineages or evolved in the anthozoan lineage after the Anthozoa/Medusozoa split. One problem with the hypothesis that PaxC represents a Pax6 precursor is that this implies that a PaxC gene should be involved in specifying jellyfish eyes, although to date there is no evidence that this is the case.

5.1.6 Statement of Goals

To better understand the relationship of the cnidarian genes with the Pax6 and Pax2/5/8 classes, we studied the DNA-binding specificity of PaxCam and PaxBam Paired domains in vivo and in vitro. This work was done in collaboration with Dr Serge Plaza and Prof Walter Gehring of Biozentrum at the University of Basel, Switzerland. The results reported in this chapter were a necessary extension of my honours project, and have been published (Plaza et al., 2003). In order to put the new data in context, some of my honours work has been included in this chapter. The expression of PaxBam in Drosophila was performed during my PhD enrolment, but the corresponding PaxCam work was conducted during my honours year (sections 4.2.1 and 4.2.5), as were the in vitro DNA binding assays (described in sections 4.2.2 - 4.2.4). The yeast one-hybrid experiments (section 4.2.6) were conducted during my PhD enrolment. The Drosophila work and aspects of the yeast one-hybrid analyses were performed by Dr Serge Plaza.

5.2 Results

The results indicate that both PaxBam and PaxCam proteins bind to EY targets in vivo and in vitro, and thus indicate that the relationship between these cnidarian proteins and the Pax6 and Pax2/5/8 classes of bilateral animals is unlikely to be simple. The literature suggests that Pax gene loss may be an ongoing process within the Cnidaria. We suggest that in non-anthozoan cnidarians, PaxB may have acquired the roles of PaxC or alternatively, that within the Anthozoa PaxC may have arisen from a PaxB-like ancestor to fulfil more restricted roles.
5.2.1 Expression of PaxCam in Drosophila imaginal discs results in a dominant negative-like phenotype

In order to test the ability of PaxCam to generate ectopic eyes, several independent Drosophila lines carrying the PaxCam cDNA transgene under the control of the yeast GAL4-UAS regulatory sequence were generated via P-element-mediated germ line transformation (Brand and Perrimon, 1993). The UAS sequence upstream of the cDNA results in the transcription of the cDNA when yeast GAL4 is expressed in Drosophila cells. A UAS-Drosophila eyeless cDNA line was used as a positive control for the effects of induced EY expression. The UAS-PaxCam fly lines were crossed with the dpp-GAL4 driver, permitting expression of the PaxCam cDNA in the wing, leg and eye/antenna discs. In the case of the UAS-ey lines, GAL4-directed expression of EY in any of these imaginal disc types resulted in ectopic eyes in the corresponding adult structure. GAL-4 driven expression of PaxCam however, did not only result in ectopic eye formation, but also appeared to interfere with the development of adult structures arising from the disc in which it was expressed (wing, leg and eye). The legs were malformed and truncated, the wings did not develop correctly and the eyes were reduced in size (Fig 5.3a).

5.2.2 Chimeric PaxCam constructs encoding the EY C-terminal domain result in eye formation in Drosophila

Comparing the PaxCam C-terminal sequence with a range of PAX6 proteins from other animals suggested that the Acropora protein may lack the C-terminal transactivation domain and that the phenotype seen with ectopic expression of full length PaxCam might be due to a dominant negative effect with respect to EY targets in vivo. To test this hypothesis, a series of UAS-constructs were generated via splicing by overlap extension (Clackson et al., 1991) in which the C-terminal region of EY was transposed onto regions of PaxCam encoding the DNA-binding domains and vice versa, generating four constructs in total (see Fig 5.2). Crosses of fly lines carrying these UAS-domain swap constructs with the dpp-GAL4 driver line showed that UAS lines in which the EY C-terminal region was present were capable of triggering ectopic eye development in the wing disc. This effect was seen to be independent of the presence of the
homeodomain, i.e. with constructs in which the PaxCam PD and the EY C-terminal
domain were present. The reciprocal constructs - that is EY constructs featuring the
PaxCam C-terminal region - were incapable of inducing ectopic eyes. Eyes induced in
response to expression of PaxCam/EY fusion proteins were always significantly smaller
than those resulting from EY misexpression, but were otherwise morphologically
normal. In scanning electron micrographs regular ommatidia and inter-ommatidial
bristles are clearly visible (Fig 5.3f).

5.2.3 Ectopic sine oculis expression is induced by PaxCam/EY constructs

The Drosophila experiments described above implied that the PaxCam protein bound to
EY targets in vivo and this was tested directly by examination of the effect of PaxCam
on sine oculis (so) expression in vivo. sine oculis is one of the best characterised direct
targets of EY; the EY protein activates expression of so by binding to an eye-specific
enhancer called so10 in the so gene (Niimi et al., 1999). Figure 5.4 shows LacZ
staining patterns in wing discs from fly lines in which expression of the enhancer trap
so-LacZ (Cheyette et al., 1994) was driven by various EY, PaxCam or PaxCam/EY
chimeric constructs. Both of the constructs in which the EY C-terminal domain was
present (Fig 5.4c-d) activated so expression, albeit at significantly lower levels that did
EY (Fig 5.4b). Constructs consisting of the EY PD and the PaxCam HD and C-terminal
region did not drive significant levels of so expression (data not shown).

5.2.4 In vitro binding properties of Pax-Cam and Pax-Bam PDs

The 128bp fragment in the so enhancer found to be a direct target of EY (so10) by band
shift assays and DNA footprinting experiments (Niimi et al., 1999; Punzo et al., 2002)
was used in vitro to investigate the DNA-binding specificity of PaxBam and PaxCam
PDs. In addition, the ability of the PaxBam and PaxCam PDs to bind to consensus
Pax2/5/8 and Pax6 binding sites were also examined. In each case, the recombinant
PDs bound specifically and with high apparent affinity to the labelled oligonucleotides,
and although the method does not permit quantitation of the interaction, no major
differences were apparent between the PDs in affinity for the oligonucleotides (Fig 5.5).
Fig. 5.2: Schematic representation of constructs expressed in Drosophila imaginal discs. (A) The EY protein, (B) Pax-Cam, (C-F) EY / Pax-Cam chimeric proteins, (G) Pax-Bam / EY chimera.
Fig. 5.3: Phenotypes resulting from expression of Pax-Cam constructs in Drosophila wing imaginal discs. The constructs shown in Fig. 5.2 were expressed under GAL4-UAS control in wing discs. Expression of the eyeless cDNA leads to the formation of ectopic eye tissue seen as the red-pigmented structure (a), whereas expression of the Pax-Cam cDNA does not result in eye formation (b). (c-d) Expression of chimeric constructs (c and d in Fig. x) encoding the C-terminal region of EY result in eyes that are smaller than those induced by EY. (e) Expression of Pax-Cam in the wing disc causes severe abnormalities. (f) At the SEM level, the morphology of the eyes induced by the Pax-Cam / EY constructs can be seen to include regular ommatidia and inter-ommatidial bristles.
Fig. 5.4: β-galactosidase expression in sine oculis (so)-lacZ line driven by dppGAL4-UAS Pax-Cam constructs. Wing discs are shown in which the constructs shown in Fig. 5.2 were expressed. (a) Pax-Cam does not induce significant so-lacZ expression, (b) positive control by misexpression of EY. (c-d) Both domain swap constructs encoding the C-terminus of EY (i.e. constructs c and d in Fig. 1) induced so-lacZ expression, whereas the reciprocal constructs did not (data not shown).
Fig. 5.5: DNA-binding assays using the Pax-Cam and Pax-Bam Paired domains. The binding of recombinant Pax-Cam (right panel) and Pax-Bam (left panel) PDs to (a) a consensus Pax-6 binding site (5'-AGGTTACGCTCTGATGTCAG-3'), (b) a consensus Pax-2/5/8 binding site (5'-CTAGTCATGCGTATGCTGTTCCAG-3'), and (c) a known EY target (the so10 oligonucleotide 5'-GCAAAACAGCTATTAATTCCCTCCTGCTGGCACAAT-3') was determined by electrophoretic mobility shift assays. In each case, the concentration of oligo-nucleotide was held constant, and the concentration of the PD varied from 2.6 x 10^-5 M to 7.9 x 10^-10 M (corresponding to 8-fold dilution between lanes); the right hand lane in each case is a negative control in which no protein was added. Open and closed triangles on the left of the figure indicate the positions corresponding to free and bound probe respectively.
5.2.5 Expression of a PaxBam/eyeless chimera in Drosophila

Because the PaxBam PD bound the same range of target sites \textit{in vitro} as did that of PaxCam, the morphogenetic properties of a PaxBam/EY domain swap construct were examined in \textit{Drosophila} imaginal discs. To avoid potentially complicating protein-protein interactions mediated by the Pax-Bam octapeptide motif, and to enable direct comparison with the Pax-Cam/EY phenotypes, the construct encoded only the N-terminal region and PD of PaxBam, and the region of EY C-terminal of the PD (Fig 5.2G). When expressed either in the leg or wing discs, the PaxBam/EY construct was able to induce ectopic eyes, albeit with lower efficiency than was the corresponding PaxCam/EY construct (Fig 5.6 – note the extremely weak wing disc phenotype). Thus the \textit{in vivo} data are consistent with the \textit{in vitro} DNA-binding experiments, indicating that the PDs of both PaxCam and PaxBam bind EY targets and, in the presence of the EY C-terminal domain, can initiate compound eye morphogenesis in the fly.

5.2.6 PaxCam binds to the \textit{sine oculis} eye-specific enhancer region in a yeast one-hybrid system

The yeast one-hybrid system described by Mastick \textit{et al} (1995) was used to examine the interaction of PaxCam constructs with a defined EY target. This method was employed as the \textit{Drosophila} transgenic experiments implied that PaxCam/EY chimeras were capable of activating expression of EY targets, such as \textit{sine oculis}, \textit{in vivo}. To better understand this interaction, we examined the ability of the corresponding chimeras to bind to the \textit{sine oculis} eye-specific enhancer region in a yeast one-hybrid assay. Yeast activator constructs expressing the PaxCam/EY chimeras corresponding to those used in \textit{Drosophila} were cloned into the \textit{BamHI} site of the activator plasmid pBM258T (Mastick \textit{et al}., 1995) and used in conjunction with the \textit{so10-HIS3} reporter in pHR307a previously described (Niimi \textit{et al}., 1999). The insert fragment was generated by PCR using the domain swap constructs as templates with \textit{BglII} restriction sites introduced at either end of the primers to facilitate cloning into pBM258T. Prior to expression, the PCR fragment was cloned first into pGEM-T to enable verification of the sequence,
Fig. 5.6: Comparison of phenotypes resulting from expression of Pax-Bam / EY and Pax-Cam / EY constructs in leg and wing imaginal discs. The Pax-Bam / EY construct (shown schematically in Fig. 5.2G) was capable of inducing eye morphogenesis in the leg disc only (a); typical results of expressing this construct in the wing disc are shown as (b). The corresponding Pax-Cam / EY construct (shown schematically in Fig. 5.2D) displayed stronger morphogenetic properties in both the leg (c) and wing (d) discs.
Fig. 5.7: Pax-Cam binds to the sine oculis eye-specific enhancer so10 in a yeast one-hybrid system. Expression of the proteins indicated on the right of the figure was driven by a galactose-inducible promoter that is strongly repressed by glucose. Each panel (a, b and c) represents the same colonies plated onto different media, the composition of which is indicated at the bottom of the figure. Panel a: growth control experiment. The presence of histidine (His) allows all of the colonies to grow; the medium lacks tryptophan (Trp) and uracil (Ura) to select for maintenance of the so10-His3 reporter and pBM258T activator plasmids respectively. Panel b: negative control experiment; no growth is observed on medium lacking histidine in the presence of glucose since activator proteins are not produced. Panel c: In the presence of galactose, yeast colonies are able to grow on media lacking histidine if the protein produced binds to the so10 target and activates transcription of the HIS3 reporter gene. Lane 1: pBM258T empty vector as negative control; lane 2: pBM258T Eyeless expressing vector as positive control. Lanes 3 to 6: various PaxClEy chimeras cloned into pBM258T as indicated to the right of the figure. Lanes 4, 5 and 6 correspond to constructs c, d and f respectively in Fig 1. Note that no growth is observed with all constructs on the empty His plasmid lacking the so10 sequence (not shown and Niimi et al., 1999).
then excised using *BglII* and ligated into the *BamHl* site of pBM258T (n.b. *BamHl* and *BglII* have compatible ends). Cloning into pBM258T was complicated by the size of the vector (~14kb); once achieved, insert orientation was verified by PCR. Yeast cells (strain YM4271) transformed with the activator plasmid were selected on media lacking uracil, and these were then transformed with the appropriate reporter construct. The basis of this system is that double transformants are only capable of growth on galactose media (galactose drives expression of the activator construct) lacking histidine if the chimeric protein constructs are capable of binding the *solO* region and activating expression of the reporter gene (*HIS3*). Yeast transformation and expression were performed by Dr Serge Plaza in the Biozentrum (University of Basel). Results of these experiments are shown as Figure 5.7.

### 5.3 Discussion

Our initial goal was to test the hypothesis that *PaxCam* represents a precursor of the Pax6 class by examining the morphogenetic properties of the PaxCam protein expressed in *Drosophila* imaginal discs. Although PaxCam was unable to initiate eye morphogenesis in imaginal discs, this effect appears to result from the lack of a transactivation domain and hence an inability to activate transcription in *Drosophila*. Expression of chimeric PaxCam proteins containing the C-terminal region of EY in imaginal discs resulted in eyes that were morphologically normal, but smaller than those induced by EY misexpression. The PaxCam/EY chimeras conferred a number of phenotypic characteristics normally associated with Pax6 proteins, including activation of a *so-lacZ* construct in vivo. Similarly PaxCam/EY chimeras were able to activate transcription of a HIS3 reporter by binding to the *so10* fragment in a yeast one-hybrid system. The C-terminal region of the PaxCam protein is much shorter (only 81 AA residues C-terminal of the HD) than that in EY and PAX6 proteins in general (the EY C-terminal region is 387 AA residues; that of PAX6 is 152), and contains no obvious transcription activation domain (Czerny and Busslinger, 1995; Tang *et al.*, 1998). It is therefore likely that PaxCam functions primarily as a transcriptional repressor, as does the mammalian PAX6-related protein PAX4 (Smith *et al.*, 1999).
Whilst the experiments in which PaxCam/EY chimeras were expressed in Drosophila support the hypothesis that PaxCam is a precursor of the Pax6 class, the in vitro DNA-binding properties of a second Acropora Pax protein, PaxBam led us to question this assumption. The PaxCam and PaxBam PDs bound the same range of sequences in vitro, including a known EY target site – a footprinted sequence in the so eye-specific enhancer. The PDs of the cnidarian Pax proteins appear to have relatively low DNA-binding specificites; published data for the Cladonema and Chrysaora PaxB proteins (Sun et al., 2001) are broadly consistent with the Acropora data. Although the specificity of the Acropora PaxAam PD has not yet been determined, binding to EY targets is not a universal property of cnidarian PDs, as the Acropora PaxDam PD does not bind to these same sites in vitro (Nordstrom et al., 2003a). The DNA-binding behaviour of the PaxBam PD in vitro led us to examine the morphogenetic properties of a PaxBam/EY chimera in Drosophila. The PaxBam/EY chimera was able to induce ectopic eyes in the leg disc and to a limited extent in wing discs, but with lower efficiency than the corresponding PaxCam/EY construct. Phylogenetic analyses clearly show that cnidarian PaxB and PaxC both belong to the Pax supergroup which also includes the Pax6 and Pax2/5/8 classes (Balczarek et al., 1997; Catmull et al., 1998; Groger et al., 2000; Miller, 2000). Although we have previously suggested otherwise, (Catmull et al., 1998; Miller, 2000), the results presented here suggest that there is unlikely to be a simple correspondence between the cnidarian PaxB and PaxC genes and the Pax2/5/8 and Pax6 classes in higher animals.

Clearly, Pax proteins are an ancient class of transcription factors (Hoshiyama et al., 1998) that are absent from fungi and plants (Galliot et al., 1999) but diversified very early in animal evolution (Miller, 2000). Other groups investigating the evolution of the pax family have suggested an evolutionary scheme, based solely on comparisons between the HDs and PDs of Pax proteins from a variety of Metazoa. Their scheme, which accommodates both vertebrates and arthropods, is based on the assumption that the failure to isolate PaxC and PaxD sequences from other cnidarians is due only to incomplete libraries or the fact that primers used are not compatible with the target sequence. The scheme allows only for a minimum of domain acquisitions and losses and proposes that PaxA/PoxN and PaxC/Eyg form two subgroups separate from the other four groups of Pax genes proposed by Noll, 1993; (Pax1/9, Pax3/7, Pax4/6 and Pax2/5/8), and that the PaxA/PoxN and PaxC/Eyg groups have since been lost in
vertebrate evolution but retained in certain cnidarian and arthropod lineages (Nordstrom, 2003). The PaxA/PoxN genes were proposed to be founded by a PD-containing gene which lacked a HD. Upon acquisition of the HD the PaxC/Eyg-like group of genes were formed (note that Eyg has lost part of its PD). The acquisition of an octapeptide and subsequent duplication then gave rise to the PaxB and PaxD genes. Loss of the octapeptide or part of the HD of the PaxB-like gene gave rise to Eye/Toe/Pax6 genes, and the Pax2/5/8/sparking genes respectively, while the PaxD-like gene gave rise to the Pax1/9/Poxmeso group and the Pax3/7/paired/GooseberryN group (Nordstrom, 2003) (See Fig 5.8). If it is assumed that undersampling or other issues are not the cause of the lack of PaxC and PaxD genes found in other cnidarians other than Acropora, it can be supposed that there has been ongoing loss of Pax genes throughout the Cnidaria. Using this scenario, the PaxB and PaxC types are likely to post-date the Cnidaria/Bilateria split; PaxC either originated within the common cnidarian ancestor or within the Anthozoa after the Anthozoa/Medusozoa (Hydrozoa, Scyphozoa and Cubozoa) split. In either case, the specificity associated with true Pax6 genes presumably arose after the Cnidaria/bilateral Metazoa split.

From our data we cannot exclude the possibility that the specificity of PaxB and PaxC proteins is influenced by regions other than the PD; it is quite possible that the activity and specificity of the PD is influenced by the overall protein environment. Therefore the fact that the in vitro experiments described here were carried out with PDs alone, and the in vivo work was carried out on Acropora PDs in the context of the EY protein, is one major limitation in interpreting the results. Although their DNA binding characteristics are similar, the two proteins are likely to have distinct roles – PaxCam presumably functions primarily as a repressor of transcription, whereas sequence comparisons imply that PaxB proteins may be able to act either as transactivators (via the C-terminal domain) or repressors (via the octapeptide) depending on context. The presence of complete HDs in PaxB proteins distinguishes these from the Pax2/5/8 class proper; presumably the full HD enables PaxB proteins to also act via their HD to regulate specific gene expression. In addition to common roles throughout the Cnidaria, the functional flexibility of PaxB proteins may have enabled them in some cnidarians to effectively fulfil the roles of PaxCam in Acropora. Either the roles of PaxCam may have been subsumed by PaxB in medusozoans (the non-anthozoan cnidarians), or in contrast to the Nordstrom evolutionary scheme, PaxCam may have derived from a
Figure 5.8: Suggested scenario of Pax evolution. The left column indicates genes in the Bilateria, the middle column the last common ancestor of bilaterians and cnidarians, and the right column the cnidarian Pax proteins. Paired domains are shown as grey shaded boxes, homeodomains by open boxes and octapeptides by black circles and the appropriate protein names shown. Complete or partial domain loss is shown by the delta symbol and the domain abbreviation. The question mark on cnidarian PaxD indicates the ambiguity of the octapeptide. Initially, an ancestral PD-containing gene founded the PaxA and PoxNeuro genes. Capture of the HD led to the development of the PaxC and loss of part of the paired domain led to Drosophila Eyg. After capture of the octapeptide and duplication in the ancestral organism, the cnidarian PaxB and PaxD genes were created. Further duplication events in bilaterians gave rise to Sparkling/Pax-2/5/8 and Eyeless/Pax-4/6 families and PoxMeso/Pax-1/9 and Paired, Gooseberry (Neuro)/Pax-3/7 families. Thus following this scenario, the last common ancestor of cnidarians and bilaterians had at least four Pax genes (adapted from Nordstrom et al, 2003).
PaxB-like precursor to fulfil more specific roles. One prediction of the above model is that we might expect the expression patterns of PaxB genes in non-anthozoan cindarians to correspond to the sum of the patterns of PaxCam and PaxBam in Acropora. Unfortunately, expression data are available only for Acropora PaxCam and Podocoryne PaxBPC. PaxCam has a very specific pattern of expression, in a subset of presumed neurons in the planula larva (Miller, 2000). At the same stage in Podocoryne, PaxBPC is expressed throughout the entire ectoderm (Groger et al., 2000). In Podocoryne polyps, PaxBPC expression is restricted to ectodermal cells that are either interstitial cells or neurons (or both) and in medusae, the (endodermal) pattern of PaxBPC expression is again consistent with a role in nerve cell differentiation (Groger et al., 2000). However, testing these ideas requires expression data for more Pax genes in a variety of cindarians.