



# Comparison of pretectal genoarchitectonic pattern between quail and chicken embryos

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Regionalization of the central nervous system is controlled by local networks of transcription factors that establish and maintain the identities of neuroepithelial progenitor areas and their neuronal derivatives. The conserved cerebral Bauplan of vertebrates must result essentially from conserved patterns of developmentally expressed transcription factors. We have previously produced detailed molecular maps for the alar plate of prosomere 1 (the *pretectal region*) in chicken (Ferran et al., 2007, 2008, 2009). Here we compare the early molecular signature of the pretectum of two closely related avian species of the family Phasianidae, *Coturnix japonica* (Japanese quail) and *Gallus gallus* (chicken), aiming to test conservation of the described pattern at a microevolutionary level. We studied the developmental pretectal expression of *Bhlhb4*, *Dbx1*, *Ebf1*, *Gata3*, *Gbx2*, *Lim1*, *Meis1*, *Meis2*, *Pax3*, *Pax6*, *Six3*, *Tal2*, and *Tcf7l2* (*Tcf4*) mRNA, using *in situ* hybridization, and PAX7 immunohistochemistry. The genoarchitectonic profile of individual pretectal domains and strata was produced, using comparable section planes. Remarkable conservation of the combinatorial genoarchitectonic code was observed, fundamented in a tripartite anteroposterior subdivision. However, we found that at corresponding developmental stages the pretectal region of *G. gallus* was approximately 30% larger than that of *C. japonica*, but seemed relatively less mature. Altogether, our results on a conserved genoarchitectonic pattern highlight the importance of early developmental gene networks that causally underlie the production of homologous derivatives in these two evolutionarily closely related species. The shared patterns probably apply to sauropsids in general, as well as to more distantly related vertebrate species.

**Keywords:** prosomere 1, precommissural pretectum, juxtacommissural pretectum, commissural pretectum, Galliformes, Phasianidae, forebrain, diencephalon

## INTRODUCTION

Starting from early induction processes taking place at the neural plate stages, the partitions of the central nervous system are established sequentially by positional information created by morphogens secreted from various organizers. Prior to neural tube closure the major brain subdivisions – forebrain, midbrain, and hindbrain – are molecularly delimited one from another. Each of these partitions becomes regionalized subsequently by expressing specific combinations of genes, including transcription factors and cell-signaling, cell-cycle-regulating, and cell-adhesion molecules, which work in a networked way to generate different position-dependent molecular identities, specific neuronal, and

glial populations and the resulting differential histogenesis of each molecularly individualized sector of the neural wall. Among the different types of genes, those coding for transcription factors (which represent collectively the core of the developmental network) play a key role in this process, since they specify, modifying, or stabilizing, the molecular codes active during the developmental process of any given brain region. The diffusible morphogens released from the secondary organizers establish gradient concentration fields of signals whose interpretation by competent cells can change the primary molecular identity of the local histogenetic area (the neural progenitor cells) by up/down-regulation of downstream transcription factors (Puelles, 1995, 2001; Echevarria et al., 2003; Puelles et al., 2004; Kiecker and Lumsden, 2004; Davidson, 2006; Davidson and Erwin, 2006, 2009; Guillemot, 2007; Sánchez-Arrones et al., 2009). Supporting the idea that specific genetic codes identify each neural partition (and establish the respective boundaries), several comparative studies of gene expression patterns have revealed that homologous neural regions largely share a common molecular code at early stages of development across vertebrates (Fernandez et al., 1998; Hauptmann and Gerster, 2000; Puelles et al., 2000; Bachy et al., 2001; Murakami et al., 2001; Hauptmann et al., 2002; Puelles and Rubenstein, 2003; Ferran et al., 2007, 2008;

**Abbreviations:** AAB, alar–basal boundary; CNS, central nervous system; CoP, commissural pretectum; Di, deep intermediate zone; DMB, di-mesencephalic boundary; Hab, habenula; Hyp, hypothalamus; i, intermediate zone; JcP, juxtacommissural pretectum; MCPC, magnocellular nucleus of the posterior commissure; mes, mesencephalon; mi, middle intermediate; oi, outer-intermediate zone; OT, optic tectum; p1, prosomere 1; p2, prosomere 2; p3, prosomere 3; p1MT, p1 medial terminal nucleus; PcP, precommissural pretectum; pe, periventricular zone; PG, pineal gland; rh, rhombencephalon; RP, roof plate; Tel, telencephalon; TG, tectal gray; Th, thalamus; TPB, thalamo-pretectal boundary; tt, tecto-thalamic tract; su, superficial zone; v, ventricular zone.

García-López et al., 2008; Abellán and Medina, 2009; Bardet et al., 2010; Morona et al., 2010). A shared molecular pattern underpinning conserved morphogenesis may be held to represent the topological and molecularly conserved Bauplan of the neural tube, common to all vertebrates (Puelles and Rubenstein, 1993, 2003; Puelles, 1995, 2001; Puelles et al., 2004, 2007). Within this paradigm, combined gene expression patterns *properly correlated to their topological position* are highly relevant for dissecting the molecular code of each region, domain, layer, or nucleus. This is a scientific endeavor recently defined as the study of cerebral genoarchitecture (Ferran et al., 2009; Puelles, this volume). It is proving to be extremely useful for identifying homologous regions and derivatives between different vertebrates (Puelles and Medina, 2002; Puelles and Rubenstein, 2003; Ferran et al., 2007, 2008, 2009).

The pretectal region is the alar plate of prosomere 1 (p1), placed between the thalamus and the midbrain (Puelles et al., 2007). Building upon the pioneering studies on the chicken pretectum done by Rendahl (1924) and Kuhlenbeck (1939), several recent contributions have improved our knowledge of the anatomy of this region, both in terms of its subdivision into histogenetic domains and the characterization of its various derivatives (Redies et al., 1997, 2000; De Castro et al., 1998; Yoon et al., 2000; Ferran et al., 2007, 2008, 2009; Puelles et al., 2007). We defined a number of molecular characteristics of the pretectal identity codes from stage HH13 to posthatched chicken (Ferran et al., 2007, 2009). Specific molecular markers identify the rostral or pretecto-thalamic boundary (*Pax3*, *Meis1*) and the caudal or pretecto-mesencephalic boundary (*Pax6*, *Meis2*). In addition, the pretectal region was subdivided molecularly into three anteroposterior domains described as *precommissural pretectum* (PcP; *Bhlb4*, *Ebf1*), *juxtacommissural pretectum* (JcP; *Six3*, *Tal2*, *Lim1*), and *commissural pretectum* (CoP, *Pax7*, *Tal2*, *Lim1*). Several molecularly distinct dorsoventral subdomains of the three primary AP domains were detected as well (Ferran et al., 2007, 2009). Importantly, these molecular partitions were largely consistent with the fate map studies of García-López et al. (2004). The same tripartite molecular subdivision was found in the pretectum of the mouse (Ferran et al., 2008) and the *Xenopus laevis* frog (Morona et al., 2010).

Our present aim was to analyze the degree of conservation of pretectal genoarchitecture at a microevolutionary scale. For this purpose, we studied two closely related avian species (*Coturnix japonica* and *Gallus gallus*) that belong to the family Phasianidae, from the order Galliformes (Kimball et al., 1999; Crowe et al., 2006; Kan et al., 2010a,b; Shen et al., 2010). We chose to perform our study at the intermediate stages Q26/HH26 and Q28/HH28 (Q = Ainsworth et al., 2010 quail stages; HH = Hamburger and Hamilton, 1951 chicken stages). These stages represent in the chick the transition between early neurogenesis (HH20–HH27) and incipient mantle layer transformation into pronuclei (HH28–HH32; Crossland and Uchwat, 1982; Martínez, 1987; Puelles et al., 1987; Ferran et al., 2009). These embryos are still far from the period when definitive nuclei start to be distinguished (HH33–HH36). These important “hinge stages” have not been analyzed in depth previously. Therefore, we obtained a molecular map of selected markers for these developmental stages in both avian species, studying the previously defined molecular codes for each anteroposterior pretectal domain, as well as the relative pattern in which the incipient mantle strata became segregated molecularly. We included detailed studies of expression patterns for

some genes that were not previously analyzed, or had been only superficially explored at these stages (*Ebf1*, *Gata3*, *Meis1*, *Meis2*, *Pax3*, and *Tcf712* [*Tcf4*]). We found in both species remarkable conservation at the studied stages of the combinatorial genoarchitectonic codes across the tripartite anteroposterior subdivision, and virtually the same combinatorial code with regard to radial stratification.

## MATERIALS AND METHODS

The present research conforms to the stipulations of the European Community (86/609/EEC) and the Spanish Government (Royal Decree 223/1998) on care and use of laboratory animals.

## ANIMALS

Fertilized quail (*Coturnix coturnix japonica*) and chicken eggs (*Gallus gallus domesticus*) from a local farm were incubated at 37°C in a forced-air incubator until the desired embryonic stage (Q26/HH26 to Q28/HH28). Around 60 embryos were staged according to Ainsworth et al. (2010) for quails and Hamburger and Hamilton (1951) for chickens. The embryos were fixed overnight by immersion in 4% paraformaldehyde in phosphate-buffered saline (PBS, pH7.4) at 4°C. Some embryos used for *in situ* hybridization (with or without immunochemical counterstain) were processed as whole-mounts (or as brain halves, each half serving for 1–2 contrasting markers). The rest of the embryo heads were fixed overnight, washed 24 h in cold PBS, cryoprotected overnight in 20% sucrose in PBS, and finally embedded in 15% gelatin and 20% sucrose in PBS. The blocks were cryostat-sectioned 18 µm-thick in topologically true horizontal and transversal section planes relative to the length axis of p1 (pretectum). The sections were adhered onto SuperFrost-Plus slides (Menzel-Gläser, Braunschweig, Germany), separated into four or five parallel series. Each series was processed subsequently for *in situ* hybridization with a different mRNA probe. Some series were immunoreacted after the *in situ* hybridization reaction.

## RT-PCR

RNA was extracted from fresh dissected brains of *G. gallus* embryos at stages HH24, HH30, or HH33 and *C. japonica* at stage Q26 using Trizol reagent (10296-028, Invitrogen). The RNA was treated with DNase I (18068-015, Invitrogen) during 15 min at room temperature, and then the enzyme was inactivated at 65°C. The RNA was reverse-transcribed into single-stranded cDNA with Superscript II reverse transcriptase and oligo dT anchored primers (11904-018, SuperScript First-Strand Synthesis System for RT-PCR; Invitrogen). The resulting first-strand cDNA (0.5 µl of the reverse transcription reaction) was used as a template for PCR, performed with Taq polymerase (M8305, Promega). Specific primers listed below were designed to clone gene fragments from *Bhlhb4*, *Ebf1*, *Gbx2*, *Pax3* from *G. gallus* sequences; and *Bhlhb4*, *Dbx1*, *Ebf1*, *Gata3*, *Lim1*, *Meis1*, *Meis2*, *Pax3*, *Pax6*, *Pax7*, *Six3*, and *Tcf712* from *C. japonica* sequences (*G. gallus* sequences were used for primer design).

CQBhlhb4F1: 5'-ATGGCCGAGCTCAAGTCGCT-3';  
 CQBhlhb4F2: 5'-TGGGCAAGTCGGCAGAGAG-3'.  
 CQBhlhb4R: 5'-TCAAGGCTTGTCGCTGCAGT-3'.  
 CQEbf1F1: 5'-CAGTCAATGTTGATGGCCAT-3';  
 CQEbf1F2: 5'-GCACAACAATTCCAAGCAT-3';  
 CQEbf1R: 5'-AGGAGAAGTTTTCGGTCTCA-3'.

*CGbx2F*: 5'-AGCGACCTCGACTACAGCTC-3';  
*CGbx2R*: 5'-ATTCACAAGACGGGAGTTGG-3'.  
*CPax3F*: 5'-GCCGCCGCGATGACCACG-3';  
*CPax3R*: 5'-GAGCGAGACCGGAAAATAACACCA-3';  
*QDbx1F*: 5'-GTCCCCGCTACACAAGGCAC-3'.  
*QDbx1R*: 5'-CTTCCTGCTCCAGGTATTCG-3'.  
*QGata3F*: 5'-TGGAACCTCAGCCCTTTTTC-3'.  
*QGata3R*: 5'-GTAAAGGAGCTGCTCTTGG-3'.  
*QLim1F*: 5'-GGAGCAAAGTGTTCACCTG-3'.  
*QLim1R*: 5'-CGGTTCTGGAACCACACCTGG-3';  
*QMeis1F*: 5'-GTGTTCCGCAAACAGATCCG-3';  
*QMeis1R*: 5'-CCTCCATGCCCATATTCATGC-3'.  
*QMeis2F*: 5'-ATGGCGCAAAGGTACGATGAG-3';  
*QMeis2R*: 5'-TTGCGACTGATTTACAAGAT-3'.  
*QPax3F*: 5'-TCGGCGGCAGCAAACCCAAG-3'.  
*QPax3R*: 5'-GGCTCCTGCCTGCTTCCTCC-3'.  
*QPax6F*: 5'-GCAGTATTACGAAACTGGC-3'.  
*QPax6R*: 5'-GGGTTGCATAGGCAGGTTGT-3'.  
*QPax7F*: 5'-GATGTTACAGCTGGGAGATCC-3'.  
*QPax7R*: 5'-ACAGGATTCATGTGGTTG-3'.  
*QSix3F*: 5'-GTGGCCAGCGTCTGCGAGAC-3'.  
*QSix3R*: 5'-GTTAAAGGAGCTGCTCTTGG-3'.  
*QTcf7l2F*: 5'-CACCCGCACCATGTACACCC-3'.  
*QTcf7l2R*: 5'-CCTTCACCTTGTATGTAGCG-3'.

PCR conditions were as follows: 5 min at 94°C, then 35 cycles (30 s at 94°C, plus 1 min at  $T_m$  temperature -57°C, and 1 min at 72°C), followed by 20 min at 72°C. The PCR products were cloned into pGEM-T Easy Vector (Promega) and sequenced (SAI, University of Murcia). *C. japonica* and *G. gallus* sequences were submitted to GenBank (accession numbers: HQ436513, *Bhlhb4 C. japonica*; HQ436514, *Dbx1 C. japonica*; HQ436515, *Ebf1 C. japonica*; HQ436512, *Ebf1 G. gallus*; JF297589, *Gata3 C. japonica*; HQ436511, *Gbx2 G. gallus*; JF719578, *Lim1 C. japonica*; JF304296, *Meis1A.2 C. japonica*; HQ436516, *Meis2A.1 C. japonica*; HQ436517, *Pax3 C. japonica*; HQ436518, *Pax6 C. japonica*; HQ436519, *Pax7 C. japonica*; JF297588, *Six3 C. japonica*; HQ436520, *Tcf7l2 C. japonica*).

#### IN SITU HYBRIDIZATION

Digoxigenin-11-UTP (Roche, Lewes, UK) was used for RNA probe synthesis. Plasmid information is provided in **Table 1**.

The whole-mount *in situ* hybridization protocol used was that of Shimamura et al. (1994), whereas the ISH reaction on cryostat-sectioned material followed the protocol of Hidalgo-Sánchez et al. (2005). After ISH, the sections were washed, and the hybridized product was detected immunocytochemically using anti-digoxigenin antiserum coupled to alkaline phosphatase (Roche Diagnostics, Mannheim, Germany). Nitroblue tetrazolium/bromochloroindolyl phosphate (NBT/BCIP) was used as chromogenic substrate for the final alkaline phosphatase reaction (Boehringer, Mannheim, Germany).

#### IMMUNOHISTOCHEMISTRY

After *in situ* hybridization, immunocytochemical reaction for PAX7 was performed in whole-mounts according to Ferran et al. (2007). Hybridized cryostat sections were processed for

**Table 1 | List of gene markers used, sequence of the gene in the construct, and origin of the plasmid construction.**

Gen	Species	NCBI number/ size/position	Laboratory/references
Bhlhb4	<i>G. gallus</i>	NM_204504.2 (663bp;77-739)	Present results
Bhlhb4	<i>C. japonica</i>	HQ436513 (666bp;1-666)	Present results
Dbx1	<i>G. gallus</i>	XR_026947.1 (1511bp; 1-1511)	Ferran et al. (2007)
Dbx1	<i>C. japonica</i>	HQ436514 (441bp;1-441)	Present results
Ebf1	<i>G. gallus</i>	HQ436512 (1055bp;1-1055)	Present results
Ebf1	<i>C. japonica</i>	HQ436515 (1004bp;1-1004)	Present results
Gbx2	<i>G. gallus</i>	HQ436511 (977bp, 1-977)	Present results
Gata3	<i>G. gallus</i>	NM_001008444.1 (779bp;758-1533)	EST clon: ChEST663o17; Boardman et al. (2002)
Gata3	<i>C. japonica</i>	JF297589 (844bp; 1-844)	Present results
Lim1	<i>G. gallus</i>	NM_205413.1 (1348bp; 105-1453)	Matsunaga et al. (2000)
Lim1	<i>C. japonica</i>	JF719578 (454bp; 1-454)	Present results
Tal2	<i>G. gallus</i>	XM_424886.2 (898bp; 17-915)	EST clon: ChEST45a19; Boardman et al. (2002)
Meis1A.2	<i>G. gallus</i>	FJ265709.1 (1230bp;1-1230)	Sánchez-Guardado et al. (2011)
Meis1A.2	<i>C. japonica</i>	JF304296 (787bp; 1-787)	Present results
Meis2A.1	<i>G. gallus</i>	HQ436521 (1231bp;1-1231)	Sánchez-Guardado et al. (2011)
Meis2A.1	<i>C. japonica</i>	HQ436516 (905bp;1-905)	Present results
Pax3	<i>G. gallus</i>	NM_204269.1 (1514bp; 12-1525)	Present results
Pax3	<i>C. japonica</i>	HQ436517 (344bp;1-344)	Present results
Pax6	<i>G. gallus</i>	NM_205066.1 (1327bp; 553-1880)	Ferran et al. (2007)
Pax6	<i>C. japonica</i>	HQ436518 (845bp;1-845)	Present results
Pax7	<i>C. japonica</i>	HQ436519 (744 bp;1-744)	Present results
Six3	<i>G. gallus</i>	NM_204364 (798bp; 406-1204)	Bovolenta et al. (1998)
Six3	<i>C. japonica</i>	JF297588 (647bp; 1-647)	Present results
Tcf712	<i>G. gallus</i>	AB040438.1 (2345bp; 1-2345)	Matsunaga et al. (2000)
Tcf712	<i>C. japonica</i>	HQ436520 (827bp;1-827)	Present results

immunocytochemistry in a similar way. After several washes in 0.1 M PBS with 0.75% Triton X-100 (PBT), sections were treated with 3% hydrogen peroxide in PBT for 15 min in the dark to inactivate endogenous peroxidase activity. After several rinses in PBT, they were blocked with 0.2% gelatin (PBGT) and 0.1 M lysine during 1–4 h. The anti-PAX7 primary antibody diluted in PBGT was incubated for 48 h at 4°C (1:20; Developmental Studies Hybridoma Bank, Iowa City, IA, USA). After washes in PBT and PBGT, the tissues were incubated with biotinylated goat anti-mouse IgG (1:200 in PBT, 1.5 h; Vector Laboratories, Burlingame, CA, USA), passing thereafter to streptavidin/horseradish peroxidase complex (Vectastain-ABC kit; Vector Laboratories; 1:350, 1 h). The brown peroxidase reaction was performed with a 15-min soak in 3,3-diaminobenzidine [Sigma, St. Louis, MO, USA; 50 mg/100 ml in 0.05 M Tris–HCl buffer (pH 7.6)], plus 5–10 min with added 0.03% hydrogen peroxide. The reaction was stopped with 0.05 M Tris buffer. Sections were coverslipped with Eukitt.

## IMAGING

Digital microphotographs of the whole-mounts were obtained with a Zeiss Axiocam camera (Carl Zeiss, Oberkochen, Germany), and a ScanScope digital scanner (Aperio, Technologies, Inc.; Vista, CA, USA) was used for the cryostat sections (exported as TIF files). The images were corrected for contrast and brightness in Photoshop 7.0. In order to compare different gene expression patterns, images were artificially pseudocolored (from blue to red or green) and superimposed using Photoshop 7.0. The plates were labeled using Adobe Photoshop IllustratorCS2 (Adobe Systems, San José, CA, USA). Representative sections from comparable chicken and quail embryos were measured using Image-ProPlus (Media Cybernetics Inc.). Image J software was used to delineate contours around areas to be compared quantitatively, obtaining the areas (in arbitrary units) of the whole pretectum at several sections levels, as well as the individual precommissural, juxtacommissural, and commissural domains. Considering volume proportional to sectioned surface, the promediated values of the individual domains were calculated as percentage of the total pretectal area, giving a rough estimate of relative proportions. No statistic was possible due to the small number of data.

## RESULTS

### PRELIMINARY REMARKS ON THE SELECTION OF SPECIES, GENES, AND DEVELOPMENTAL STAGES

The goal of the present work was to make a comparative study of the spatio-temporal distribution of key mRNAs in the pretectal region of two closely related species. Since a previous detailed study was made in *G. gallus* (Chicken; Ferran et al., 2007, 2009), we decided to use *C. japonica*, a member of the same avian family (Phasianidae). This species is commonly used as tissue-donor for quail–chicken grafting experiments (Figure 1A). Regarding the stages, we aimed to characterize early combinatorial molecular codes and test pattern profiles defining the general pretectal Bauplan during initial steps of pronuclear generation (after radial migration of postmitotic neurons into the mantle layer). Therefore, we decided to perform our study between stages Q26/HH26 and Q28/HH28, thus also expanding our previous chicken studies. Finally, the genes chosen as markers were selected accord-

ing to our previous knowledge of their importance for delimiting boundaries or a specific domain, layer, or nuclear derivative (Ferran et al., 2007, 2009).

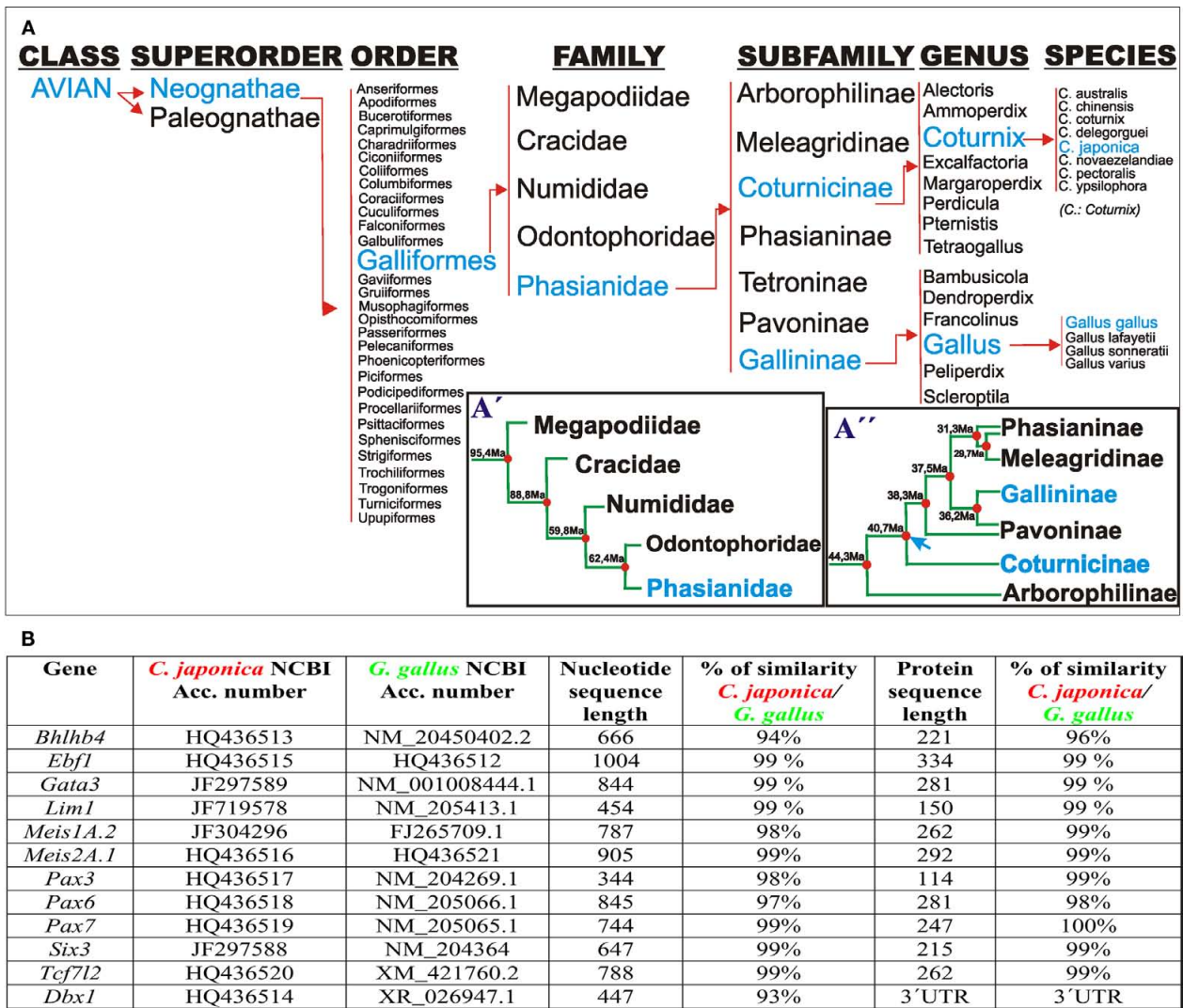
### DIFFERENCES IN TRANSCRIPT SEQUENCES BETWEEN *C. japonica* AND *G. gallus*

*Coturnix japonica* and *G. gallus* underwent 40 million years of independent evolution as members of the Coturnicinae and Gallinae subfamilies, respectively (Figure 1A). Therefore, our first aim was to assess any differences at the nucleotide and aminoacidic level of the studied genes. For this purpose, we cloned the complete coding sequence of *Bhlhb4*, obtained partial coding sequences from *Ebf1*, *Gata3*, *Lim1*, *Meis1*, *Meis2*, *Pax3*, *Pax6*, *Pax7*, *Six3*, and *Tcf7l2*, we also studied the 3'UTR of *Dbx1* from *C. japonica*, comparing them to available *G. gallus* sequences. We found >93 and >96% (often ~99%) similarity at the nucleotide and protein levels, respectively, for all the analyzed *C. japonica* and *G. gallus* genes (Figure 1B). In addition, we compared the four published gene sequences from *C. coturnix* obtained from GenBank, finding a similar level of conservation in *G. gallus* and *C. japonica* (data not shown). These results indicate a high level of mRNA sequence conservation between *C. japonica* and *G. gallus*, which is likely to represent the general degree of divergence among the subfamilies of the family Phasianidae.

### PRETECTAL BOUNDARIES AND DOMAINS ARE DEFINED BY THE SAME GENOARCHITECTONIC CODES IN *C. japonica* AND *G. gallus*

A major conclusion of our previous studies was that the combined expression patterns of three genes were enough to define the antero-posterior pretectal boundaries and its internal tripartition (Ferran et al., 2007, 2008). Here we corroborated that the same molecular code applies to *C. japonica*. In both species, pretectal expression of *Pax3* stops rostrally at the thalamo-pretectal boundary (TPB; p2/p1), whereas that of *Pax6* ends caudally at the pretecto-mesencephalic limit (p1/m1); these patterns jointly define the pretectal region (alar p1; Figures 2A,C,D,E,H,I). In addition, expression of *Six3* neatly distinguished in both cases the intermediate pretectal domain (JcP; Figures 2B,E,G). Therefore, the combination of markers *Six3/Pax3/Pax6* also identifies the rostral and caudal pretectal domains (PcP and commissural pretectum, or CoP, respectively). As previously found in the chicken, the quail CoP domain showed extensive PAX7 immunoreaction (Figures 2E,I).

Next, we analyzed *Gbx2*, *Meis1*, and *Pax3* expression patterns in cryostat sections in order to define more precisely the molecular TPB. In both species, *Gbx2* is strongly expressed in the thalamic mantle zone, stopping caudally at the TPB; *Gbx2* is absent at the overlying habenular region (Figures 2J,O,O',Y and 3M–P,AE,AF). In addition, we observed a periventricular patch expressing *Gbx2* in the rostral pretectum (see below; asterisk in Figures 2O and 3N,P,AF). *Pax3* expression is largely restricted to the pretectal ventricular zone, though its signal appears as well in the mantle of the CoP domain (see below). *Pax3* transcripts generally stop rostrally at the TPB (Figures 2A,D,E,K, O',PY and 3I–L). However, at the dorsal-most levels, *Pax3* expression expands into the habenular dorsal p2 domain, and is accordingly not useful as a p2/p1 limit marker at that specific position (blue arrowhead; Figures 2A,D,E). *Meis1* appears expressed in the pretectum but not in the thalamus. Its domain stops precisely at the rostral pretectal boundary,



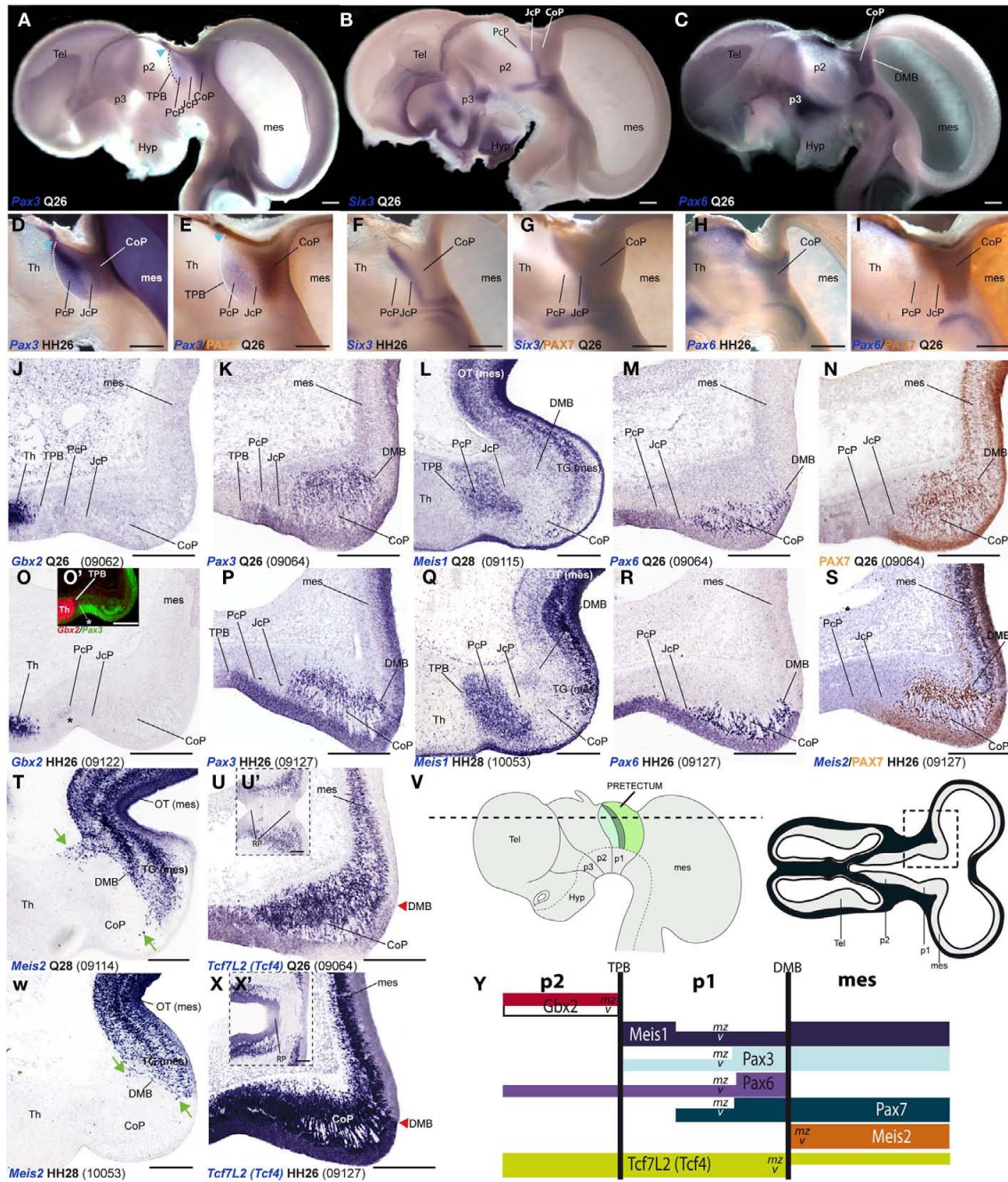
**FIGURE 1 | (A)** Taxonomic classification of Galliformes according to Crowe et al. (2006). **(A')** Postulated phylogenetic relationships between families of the order Galliformes. The timetree represents the mean times of independent evolution, estimated from several studies, according to Pereira and Baker

(2009). **(A'')** Phylogenetic relationships among members of the family Phasianidae according to Kan et al. (2010a,b). Blue arrow: time point of Gallinae and Coturnicinae split. **(B)** Comparison of the coding and protein sequences of the listed genes between *G. gallus* and *C. japonica*.

including its dorsal-most part, showing no expansion into p2, in contrast to *Pax3*. Moreover, *Meis1* shows strong expression in the ventricular and mantle zones of the PcP domain in both species (see below; **Figures 2L,Q,Y** and **3A–D**). Altogether, these genes allow the precise identification not only of the early TPB, but also of the different mantle derivatives abutting at this boundary from either side (*Meis1*, pretecal; *Gbx2*, thalamus; **Figure 2Y**).

Additionally, we compared markers that recognize the pretecal- or diencephalo-mesencephalic boundary (DMB). To this end, we studied *Pax6*, *Meis2*, *Tcf712* (previously named *Tcf4*), and *PAX7* expression patterns. *Pax6* is consistently expressed in the ventricular and mantle zones of the caudal pretecal (CoP), with a clearcut caudal end at the DMB in both animal models (**Figures 2C,H,M,R,Y**). We found that *Meis2* expression, which appears extensively at the

mesencephalic ventricular and mantle zones, stops rostrally at the mesencephalic side of the DMB boundary up to stage Q26/HH27. From Q27/HH28 onward, *Meis2* expression emerges in some periventricular and superficial cells of CoP (see below; green arrows **Figures 2S,T,W,Y**). *Tcf712* was previously reported to be expressed in the alar plate of the caudal diencephalon (thalamus and pretecal), stopping at the DMB up to stage HH24 (Ferran et al., 2007). From HH25 onward, we found that *Tcf712* transcripts expand progressively caudalward within the mesencephalic alar mantle. However, *Tcf712* signal remains restricted to the diencephalon at the level of the ventricular zone, stopping caudally at the DMB; therefore, this marker continues being useful for identifying this boundary (**Figures 2U,X,Y**). Finally, as previously described in chicken, *PAX7* is widely expressed in the ventricular and mantle zones of the caudal



**FIGURE 2 | Pairwise comparison of gene patterns in the developing pretecal region of Q26 quail or HH26 chicken embryos (as identified at left bottom corner of each panel).** The markers shown are *Pax3* (A,D,E,K,P), *Six3* (B,F,G), *Pax6* (C,H,I,M,R), *PAX7* (E,I,N,S), *Gbx2* (J,O,O'), *Meis1* (L,Q), *Meis2* (S,T,W), and *Tcf7L2* (U,U',X,X'). (A–I) Whole-mount *in situ* hybridization and immunoreaction processed material identify the main pretecal boundaries and define an anteroposterior tripartition. *Blue arrowhead* in (A,D,E): *Pax3* expression extends continuously from the pretecal into the thalamic habenular region. (J–X) Horizontal sections at corresponding levels in both species illustrating distribution along the radial dimension of the studied markers. The pairwise comparable panels are organized one on top of the other (J,O,K,P, ...). The numbers between brackets at the bottom of each panel give the code identifying the specimen in our collection (data

from the same specimen may appear in different panels or plates). Inset (O') compares in pseudocolor the patterns of *Gbx2* and *Pax3*, demonstrating that the cell group marked with an asterisk (O,O') lies caudal to the thalamo-pretecal boundary. Insets (U',X') show horizontal sections passing through the roof of the pretecal, where *Tcf7L2* is not expressed. *Green arrows* in (T,W): *Meis2* positive cells at the CoP. The schemata in (V) represent a lateral view of the brain indicating the horizontal section plane used, and the position in such sections of the pretecal region that appears magnified in the panels (dashed box). (Y) Schema of gene combinations mapped in the midbrain (mes) and diencephalic prosomeres p1 and p2. Each gene is represented by a color-coded bar. The upper and lower halves of each bar represent gene expression in the mantle (mz) or ventricular (v) zones respectively. Scale bars = 300  $\mu$ m.

pretectum (CoP). Although PAX7 is detected as well in the midbrain tectum, its expression is downregulated caudal to the DMB in a rostral alar midbrain territory intercalated between the pretectum and the optic tectum proper, the primordium of the tectal gray (TG; Ferran et al., 2007, 2009; see also García-Calero et al., 2002 and Puelles et al., 2007). Therefore, both species showed a sharp boundary at the DMB through the combination of PAX7 immunohistochemistry and *Meis2/Tcf712* mRNA *in situ* hybridization (Figures 2I,N,S,Y).

As regards the ventral boundary of the pretectum, marked in chicken by *Pax6* expression (neither *Pax3* or PAX7 reach ventrally the alar–basal boundary), it also was comparable in the quail (Figures 2C,H and 8J). In both cases, moreover *Tcf712* expression reaches dorsally the roof–alar plate boundary (Figures 2U',X' and 8K).

### CONSERVED MOLECULAR PROFILE OF THE PRECOMMISSURAL PRETECTUM DURING EARLY MANTLE HISTOGENESIS

Ventricular (*v*), periventricular (*pe*), intermediate (*i*), and superficial strata (*su*) can be distinguished in the PcP at the stages analyzed (Rendahl, 1924; Senn, 1970, 1979; Ferran et al., 2009). During the pronuclear period, the thick intermediate stratum becomes subdivided into *deep* (*di*), *middle* (*mi*), and *outer* (*oi*) intermediate layers (Ferran et al., 2009).

*Bhlhb4*, *Dbx1*, *Ebf1*, *Meis1*, and *Pax3* mRNA expression mark the PcP derivatives, therefore we compared these patterns with those of *Gbx2*, *Lim1*, and *Six3* in order to clearly identify the rostral and caudal boundaries of PcP (Figure 3). *Meis1* appears expressed at the PcP ventricular zone and all corresponding mantle layers in both avian species at stages Q26/HH26–Q27/HH27, but at Q28/HH28 its signal clearly decreased in the *pe* mantle layer (Figures 3A–D, 4A–F,Y–AD, and 7). A pioneering group of *Meis1*-positive PcP cells apparently moving incipiently into the JcP domain was clearly observed at HH26 (green arrowhead, or asterisk, in Figures 3B,D,F,AK,AL). At stages Q27/HH27–Q28/HH28 there is a population of cells expressing strongly *Meis1* in the deep part of the intermediate layer (*di*), which, when compared to *Six3* signal (a JcP marker), are revealed as PcP cells that have translocated into the JcP domain. This is the primordium of the dorsocaudal pretectal nucleus, which later also penetrates the CoP (Ferran et al., 2009). *Pax3* mRNA was only detected at the *v* and *pe* layer of the PcP mantle zone in both species (Figures 3I–L, 5G,V, and 7). We observed a previously undescribed stream of *Gbx2* positive cells localized in the periventricular layer of the PcP (Figures 2O, 3N,N',P,P',AE,AF', and 7). This *Gbx2* signal was visualized at dorsal levels of the PcP from stage Q26/HH26 onward, and it increased progressively at later stages (asterisk in Figures 2O', 3N,P,AF). *Bhlhb4* mRNA expression was strongly present at the PcP *pe* and *i* strata in both species. As described for *Meis1*, we observed a few *Bhlhb4* positive cells originated from the PcP *di* layer that apparently translocate into the JcP domain (green arrow, Figures 3Q–T,AH,AI and 7). *Dbx1* transcripts appear in the PcP *pe* layer and in a salt and pepper pattern at the *v* stratum, in both species. The expression of this gene does not stop at the thalamo-pretectal limit, but continues in p2 within the habenular region (Figures 3U–X and 7). *Ebf1* mRNA expression was observed throughout the PcP mantle layers, but not at the ventricular stratum. At dorsal PcP levels, *Ebf1* expression extends also into the habenular region of p2, but more ventrally it clearly stops at the boundary between thalamus and pretectum. As reported above

for *Meis1* and *Bhlhb4*-positive cells, comparison of the expression patterns of *Ebf1* and *Lim1* (JcP marker) showed some *Ebf1*-positive cells from the PcP *di* layer entering the JcP domain (green arrow, or asterisk; Figures 3Y–AD,AM and 7). Finally, a cluster of *Gata3*-expressing cells was observed at the PcP *oi* and *su* strata from both species (Figures 4S–V,AQ–AT).

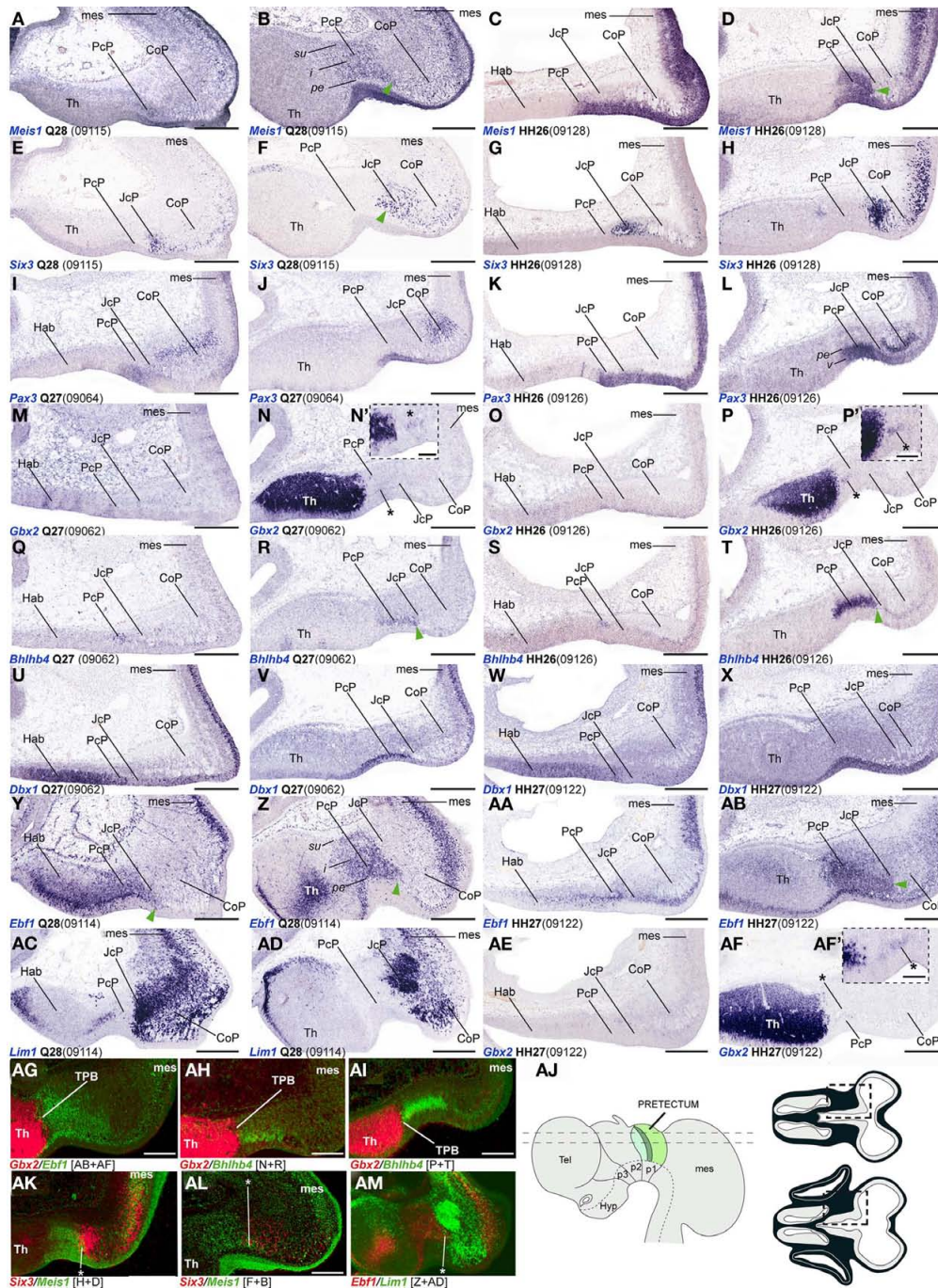
### CONSERVED MOLECULAR PROFILE OF THE JUXTACOMMISSURAL PRETECTUM DURING EARLY MANTLE HISTOGENESIS

To analyze the JcP region, we studied the *Six3*, *Gata3*, *Lim1*, and *Tal2* gene expression patterns. We had previously demonstrated in chicken that *Six3* is a selective cell marker for the mantle zone of the JcP domain from HH17/18 onward (Ferran et al., 2007). Here we first compared the expression patterns of *Six3* and *Meis1* to double-check the rostral JcP boundary along the full radial dimension (Figures 4A–L,Y–AJ). Then, we compared *Six3* with the *Lim1*, *Gata3*, and *Tal2* markers – all expressed throughout the JcP and CoP domains – to identify the caudal JcP boundary within these patterns (Figures 4G–X,AE–AV, 6E–L,Q–X, and 7). Between Q26/HH26 and Q28/HH28 all JcP mantle layers show *Six3* expression (Figures 4G–J,AE–AH and 7), but the *i* stratum, in most of the sections, and the *su* stratum, at dorsal levels, already have started to downregulate *Six3* expression at Q27/HH27 (asterisk in Figures 4I,J,AE–AH). This change in the pattern seems correlated with the onset of *FoxP1* expression at the same loci (Data not shown; see Discussion). *Lim1* is strongly expressed in all JcP mantle derivatives of both species, with a particularly compact appearance of the *oi* layer from the *i* stratum, which mainly represents the primordium of the lateral spiriform nucleus (asterisk Figures 4M–R,AK–AP; Ferran et al., 2009). JcP expression of *Gata3* mRNA, which has not been described previously, is similar to that of *Lim1* in both species (Figures 4S–V,AQ–AT). Nevertheless, a relatively stronger *Gata3* signal at the JcP *pe* and *di* strata distinguishes them from the CoP ones (Figures 4U–X,AS–AV and 7). *Tal2* signal is strong at the JcP *pe* and *di*, but decreases slightly at the *oi* layer from the dorsal and lateral subdomains; this signal is absent at the ventral JcP and CoP subdomains of both species (Figures 6E–H,Q–T and 7). *Pax3* and *Pax6* are expressed in the JcP ventricular zone, and PAX7 only appears in overlap at the level of the most dorsal JcP subdomains (Figures 5 and 7).

### CONSERVED MOLECULAR PROFILE OF THE COMMISSURAL PRETECTUM DURING MANTLE HISTOGENESIS

In order to study radial segregation of strata and layers in the CoP domain, we compared *Gata3*, *Lim1*, *Pax3*, *Pax6*, *Pax7*, and *Tal2* mRNA expression, adding occasional data from the *Bhlhb4*, *Dbx1*, *Ebf1*, *Meis1*, *Meis2*, and *Six3* expression patterns. The genes of the PAX family are variously combined within CoP, generating by themselves a molecular code that identifies the four strata observed from the ventricle to the pial surface (*v*, *pe*, *i*, *su*). The *v* zone is characterized by the expression of *Pax3*, *Pax6*, and *Pax7*, though *Pax7* expression is absent at the most ventral alar plate levels (arrowhead; Figures 5 and 7). The *pe* stratum expresses strongly *Pax6* and *Pax7*, and shows a weak *Pax3* signal (Figures 5 and 7).

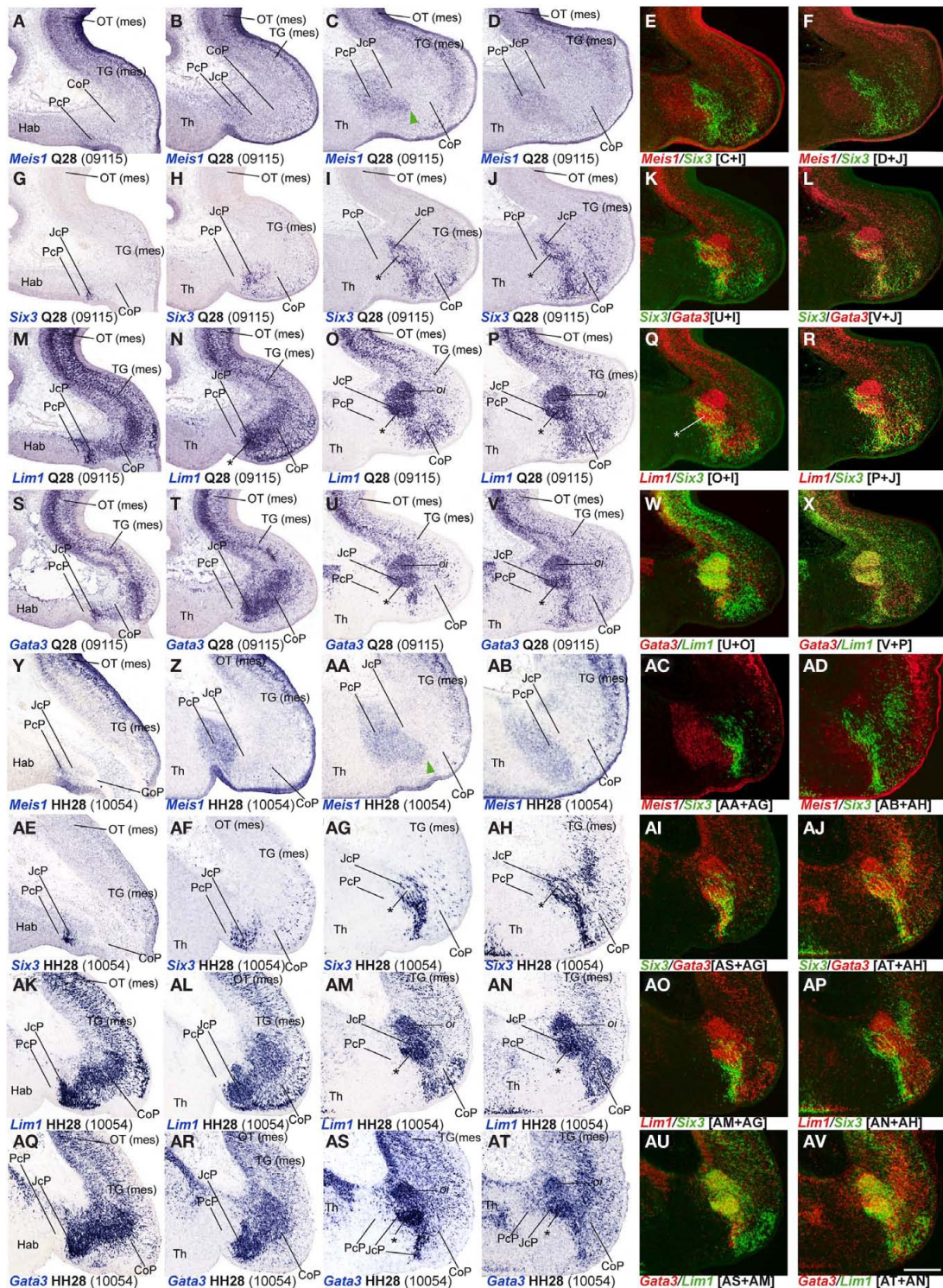
Starting at stages Q26 in quail and HH27 in chicken, the CoP *i* stratum started to be segregated in three parts, the deep, middle and outer intermediate layers (*di*, *mi*, *oi*). The thin *di* layer shows *Pax7*- and *Pax3*-positive cells (Figures 5I,J,N,O).



**FIGURE 3 | (A–AM) Comparisons of *in situ* hybridization data for PcP markers at HH26/27-Q27/28 in horizontal cryostat sections.** From top to bottom of the plate, each row of four panels generally contains data on one gene: *Meis1* (A–D), *Six3* (E–H), *Pax3* (I–L), *Gbx2* (M–PAE,AF), *Bhlhb4* (Q–T), *Dbx1* (U–X), *Ebf1* (Y–AB), *Lim1* (AC–AD). Note that material from different specimens (identifying codes in brackets) was pooled into this plate. Each adjacent pair of panels from left to right corresponds to two different section levels in one species (in dorsoventral order), to be compared with the next two panels of corresponding section levels in the other species [e.g., for *Meis1*, (A,B) are two quail section levels to be compared with (C,D), showing the same marker at equivalent levels in the chick]. Patterns comparable between quail and chicken are thus shown horizontally. The panels illustrating the other markers in

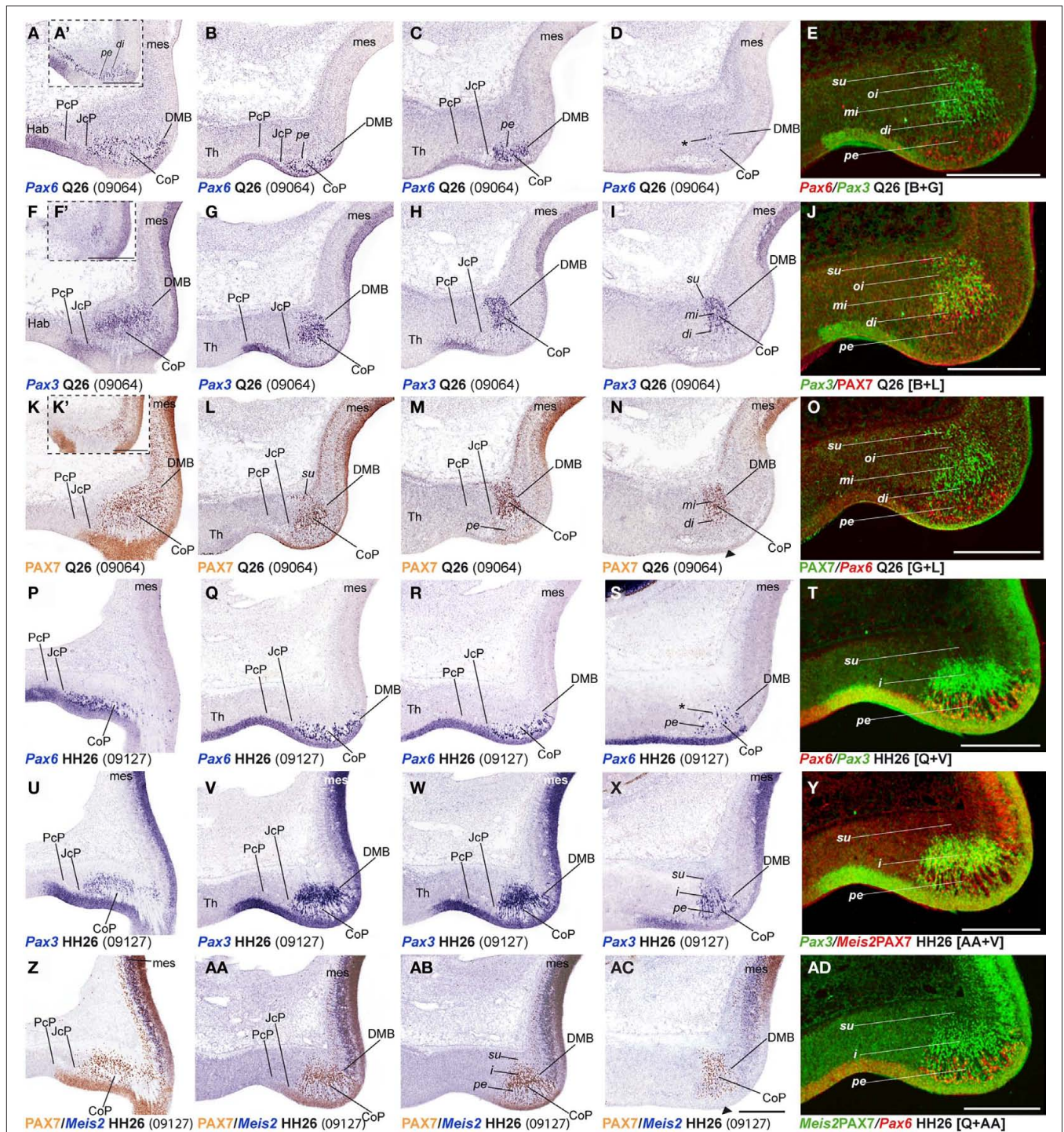
successive rows represent corresponding section levels (though sometimes a different specimen is compared), so that different gene patterns are compared vertically. *Green arrowheads* in (B,D,F,R,T,Y,Z,AB) and *asterisks* in (AK–AM) indicate small groups of cells expressing a PcP marker that lie within the molecularly identified JcP domain, suggesting a tangential migratory displacement. *Asterisks* in (N/N',P/P',AF/AF'): *Gbx2* positive cells within the PcP. (AG–AI,AK–AM) Pseudocolored superposition of the indicated pairs of marker domains from adjacent sections, to illustrate boundary relationships between them. (AJ) Schematic view of the brain indicating the two horizontal section planes illustrated in this plate, and the position in such sections of the pretectal region that appears magnified in the panels (dashed boxes). Scale bars = 300 μm for (A–AM) and 100 μm for (N',P',AF').





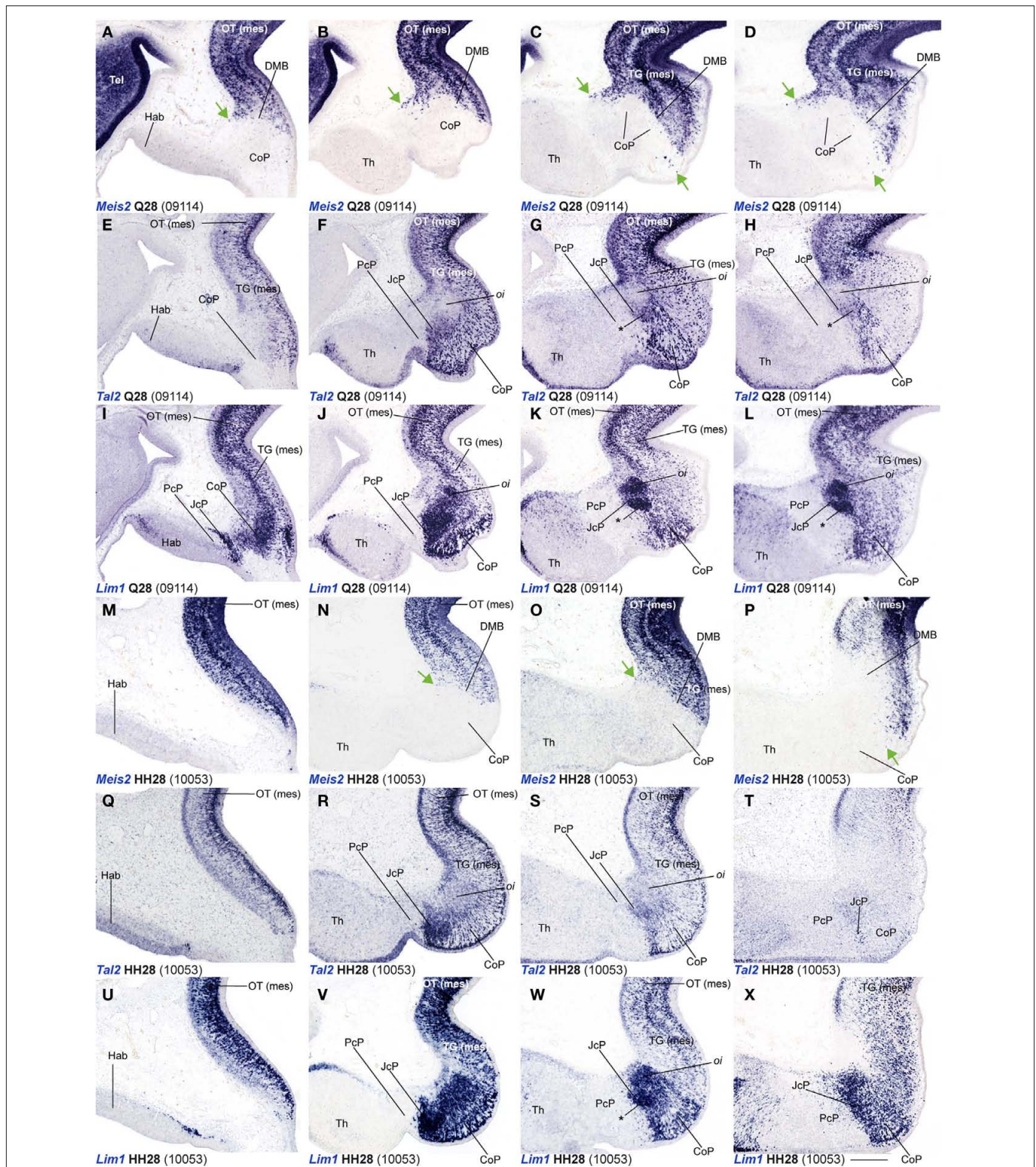
**FIGURE 4 | (A–AV) Comparisons of *in situ* hybridization data for four JcP markers at Q28/HH28 in horizontal cryostat sections at four section levels (in each case a single specimen).** The markers are identified at the bottom left corner of each panel. Each row of panels [e.g., (A–F)] shows from left to right four section levels (dorsal to ventral sequence) reacted for a particular marker, followed by two pseudocolored images comparing the same marker, but paired with another at the two ventral-most levels studied. The first four rows show different

markers in the quail, and the last four rows contain comparable material for the chick, so that panels taken along the vertical columns represent essentially the same section level and are comparable across the species. *Green arrowheads* in (C,AA) mark cells expressing a PcP marker that are found inside the differentially specified JcP domain. *Asterisks* in (I,J,O,PU,V,AG,AH,AM,AN,AS,AT) identify the developing lateral spiriform nucleus. Numbers between brackets: code of the specimen in our collection. Scale bar = 300 μm.



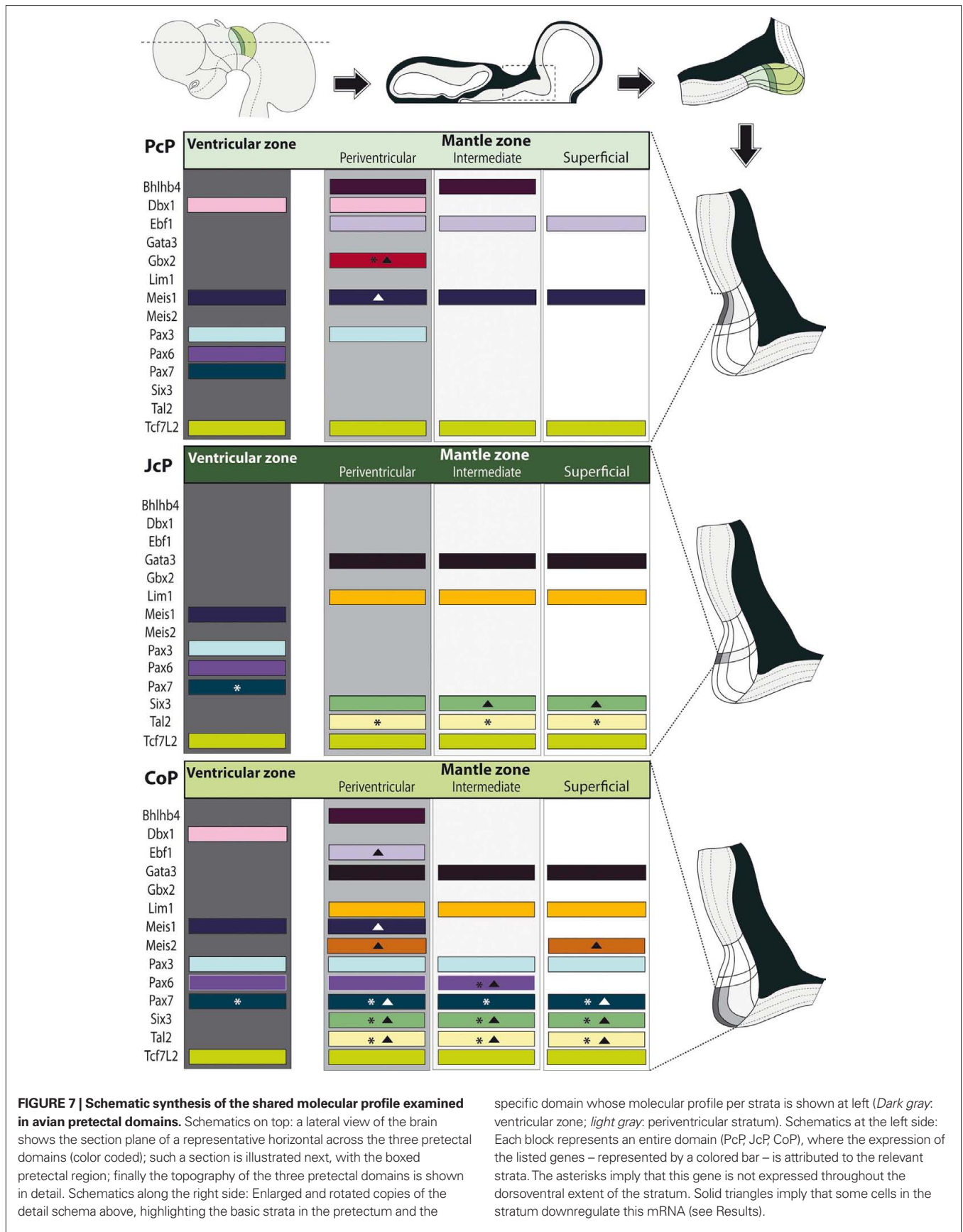
**FIGURE 5 | (A–AD) Comparisons of *in situ* hybridization/immunochemical data for three CoP markers at Q26–HH26 in horizontal cryostat sections at four section levels (in each case a single specimen).** The markers are identified at the bottom left corner of each panel. Each row of panels [e.g., (A–E)] shows from left to right four section levels (dorsal to ventral sequence) reacted for a particular marker, followed by a pseudocolored image comparing the same marker, but paired with another at the second level studied. The first three rows show different markers in the quail, and the last three rows contain

comparable material for the chick, so that panels taken along the vertical columns represent essentially the same section level and are comparable across the species. Insets (A',F',K') show a more dorsal section level, illustrating details of expression of the corresponding marker. *Black arrowheads* in (N,AC) indicate ventral levels of CoP ventricular zone lacking PAX7 immunoreaction. *Asterisks* in (D,S) identify *Pax6* positive cells in the CoP deep intermediate layer. Numbers between brackets: code of the specimen in our collection. Scale bar in (AC) applies as well to (A–D,F–I,K–N,P–S,U–X,Z–AB). Scale bars = 300  $\mu$ m.



**FIGURE 6 | (A–X) Comparisons of *in situ* hybridization data for three markers labeling JcP and CoP at Q28/HH28 in horizontal cryostat sections at four section levels (in each case a single specimen).** The markers are identified at the bottom left corner of each panel. Each row of panels [e.g., (A–D)] shows from left to right four section levels (dorsal to ventral sequence) reacted for a particular marker. The first three rows show different markers in the quail, and the last three rows contain comparable material for the chick, so that

panels taken along the vertical columns represent essentially the same section level and are comparable across the species. *Green arrows* in (A–D, N–P) indicate apparently ectopic cells expressing *Meis2* (midbrain marker) at the CoP *su* and *pe* strata. *Asterisks* in (G, H, K, L, W) identify the developing lateral spiriform nucleus, exhibiting high expression levels of *Lim1* and a low *Tal2* signal. Numbers between brackets: code of the specimen in our collection. Scale bar = 300 μm.



Additionally, at stage Q26/HH27 an incipient patch of *Pax6* expression was observed at the dorsal-most portion of the CoP *di* layer (Figures 5A,A'). At the ventral-most levels we detected a few cells of the same layer with strong *Pax6* expression, which possibly constitute the primordium of the magnocellular nucleus of the posterior commissure (MCPC; Figures 5D,S). The overlying *mi*, *oi*, and *su* CoP mantle layers showed strong *Pax7* and *Pax3* signal (Figures 5F–O,U–AD). Interestingly, the specimens used to study PAX family genes belonged precisely to the same stage, according to the respective quail and chicken developmental tables (Hamburger and Hamilton, 1951; Ainsworth et al., 2010). This allowed us to detect two variant aspects between the two species. First, the relative size of the pretectum and its CoP domain was larger in chicken than in quail (Figures 9A,B). Secondly, quail specimens at an equivalent stage in contrast seemed to be relatively slightly more advanced in development, since their layering process was more advanced than in the chicken (Figure 9C; and see Discussion). In quail embryos at Q26, we observed an incipient segregation of the intermediate stratum into *di*, *mi*, and *oi* layers, a structural pattern that was present only in chicken from HH27 onward (Figures 5A,A',I,J,N,O,P,X,Y,AC,AD).

Leaving apart *Gata3*, which had not been studied previously in this region, *Lim1* and *Tal2* were expressed in the CoP as previously described (Ferran et al., 2007). To determine precisely the antero-posterior boundaries of the CoP, we combined these expression patterns with *Six3*, thus obtaining the anterior CoP boundary, and with *Meis2*, highlighting the caudal limit of CoP or DMB boundary (Figures 4G–L,Q,R and 6E,F,K,L,Q,R,W,X). Comparing *Lim1* with *Gata3* and *Tal2*, we noticed that *Lim1* and *Tal2* are both present at the CoP *pe* stratum (although *Tal2* signal disappears at the ventral subdomains). In addition, only a few *Gata3* cells were observed at the *pe* of the dorsal CoP levels, though their number increases ventralward (Figures 4M–X,AK–AV and 6E–Q,X). The *di*, *mi*, and *oi* layers of the *i* mantle stratum express *Lim1* and *Gata3*, but *Tal2* was virtually absent there. The compact group of cells developing at the *oi* layer (the prospective pretectal-subpretectal complex; Ferran et al., 2009) and at the *su* stratum (terminal nuclei) express strongly *Lim1* and *Gata3* mRNAs (Figures 4M–X,AK–AV and 7).

*Dbx1* shows a salt and pepper pattern in the *v* zone of the CoP, and a stronger and wider expression domain in the ventral CoP subdomains in both species (Figures 3U–X). *Ebf1*, *Bhlhb4*, and *Meis1* are expressed in the *pe* stratum of the CoP (Figures 2L, 3S,T,Y–AB, 4B–F,Z–AD, and 7). *Meis2* is expressed in a small group of cells at the CoP *pe* stratum and in a major cluster at the *su* stratum close to the DMB (green arrow, Figures 2T,W, 6A–D, M–P, and 7). Interestingly, the latter *Meis2* expression was observed in a larger number of cells in the quail than in chicken of the same stage, possibly suggesting again a relatively more advanced developmental status of quail embryos. Finally, *Six3* – the main JcP marker – is also expressed in some groups of cells at the CoP *pe* layer, a pattern that best visible at the level of ventral subdomains (CoV). Moreover, the CoV *Six3* expression expands into the *i* mantle layers (Figures 3E–H and 4F–J,AD–AH).

#### SHARED MOLECULAR PROFILE IN THE DORSOVENTRAL SUBDOMAINS

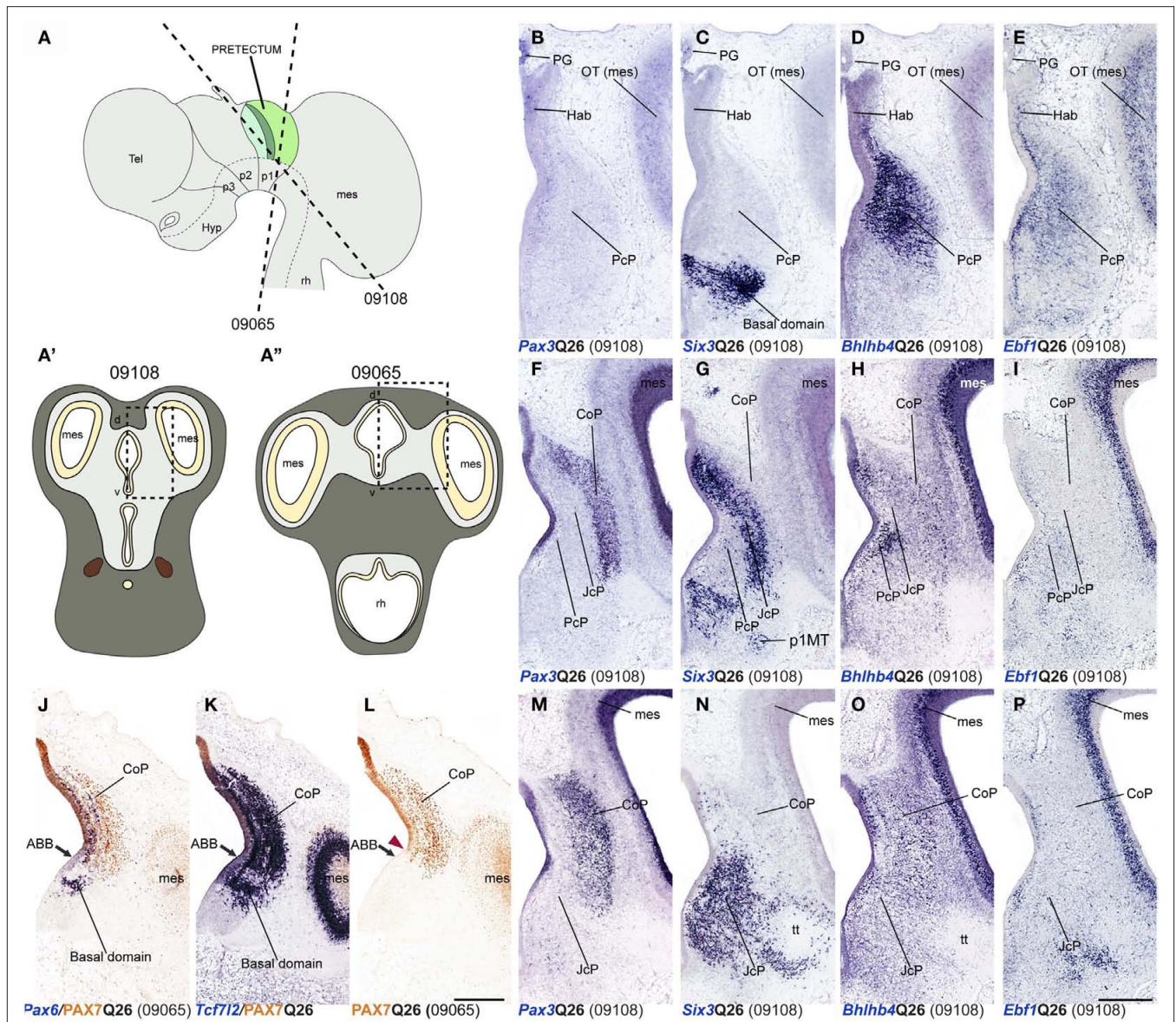
Using horizontal and transversal (coronal) sections, we were able to make a first approach to examine conservation of genoarchitectonic pattern in pretectal dorsoventral subdomains. We found

strikingly identical patterns in both species. *Pax6* signal at the level of the *vz* reaches the alar–basal boundary throughout the pretectum (Ferran et al., 2009). PAX7 expression is restricted to dorsal and lateral CoP subdomains (Figures 2, 3, 5, 7, and 8A,J–L). *Six3* signal is observed in all DV subdomains of JcP and its transcripts also appear in the mantle zone of the ventral CoP subdomains. *Tcf712* signal was distributed from the most dorsal basal plate domains up to the alar–roof boundary (Figures 2U,U',X,X', 4I–J,AG,AH, and 8C,G,K,N). *Bhlhb4* and *Ebf1* were observed in all DV parts of PcP (Figures 8D,E,H,I,O,P). *Pax3* was found in nearly all pretectal DV subdomains, except the ventral-most ones (Figures 8B,F,M). *Meis1* DV expression stopped ventrally at the alar–basal boundary of all pretectal domains in both quail and chicken (Figures 4A–E,Y–AD).

## DISCUSSION

### EQUIVALENT DEVELOPMENTAL STAGES FROM *C. japonica* (QUAIL) AND *G. gallus* (CHICKEN) SHOW DIFFERENT PRETECTAL SIZE AND DEGREE OF DIFFERENTIATION

We staged embryos according to Ainsworth et al. (2010) for quail, and Hamburger and Hamilton (1951) for chicken. Both staging tables were generated based on standard criteria about useful external anatomical landmarks, and their stages are essentially comparable one-to-one, at least in the period examined here (Hamburger and Hamilton, 1951; Ainsworth et al., 2010). However, we found that identically staged embryos from chicken and quail are not fully equivalent as regards pretectal development (in fact, brain development in general). First, the pretectal region of chicken embryos at stages HH26–HH28 was significantly larger by roughly 30% than that of quail embryos at stages Q26–Q28 (Figure 9A). Differences among close species in the proportions of adult brain regions have been suggested to be caused by changes in cell-cycle rates and/or the timing of neurogenesis, as well as by alterations of brain patterning (Finlay and Darlington, 1995; Striedter, 2005; Menuet et al., 2007; Charvet and Striedter, 2008; Charvet et al., 2010; Sylvester et al., 2010). Our results strongly suggest that differences in patterning are not likely to be responsible for the observed variation in pretectum size. Therefore, further studies should address the effect of cell-cycle rates and timing of neurogenesis upon the observed size differences. It is of course well known that, under comparable temperature conditions, the incubation period up to hatching of quail is shorter by 84 h than the chicken one. However, comparative data about the relative length of neuroepithelial cell cycles in these two species are not available. Our results suggest that differences in brain proportions between these closely related species begin during early development, as described for mesencephalic differences between parrots and galliformes (McGowan et al., 2010). Secondly, we found that the molecularly identified quail pretectal strata developing at the CoP domain appeared slightly more mature – by representing a larger proportion of the total domain volume than the corresponding chicken counterparts at equivalent stages (Figure 9C). These differences reflect an unavoidable inaccuracy of current staging tables in predicting detailed microscopic traits heterochronic relative to macroscopic morphologic features. More complex and detailed distinctive molecular features of particular organs will be needed to upgrade the standard staging tables (e.g., as done for chicken primitive streak stages by López-Sánchez et al., 2005).



**FIGURE 8 | Dorsoventral extent of seven pretectal markers in the quail, studied by *in situ* hybridization/immunoreaction in transversal cryostat sections at Q26 (two specimens). (A,A',A'')** Schemata showing a lateral view with two variant transversal section planes used (see specimen code in brackets) and the respective aspect of such transversal sections, in order to illustrate the brain region analyzed in both specimens. [Block comprising (B–I,M–P)] comparisons of four markers (*Pax3*, *Six3*, *Bhlhb4*, *Ebf1*) at three

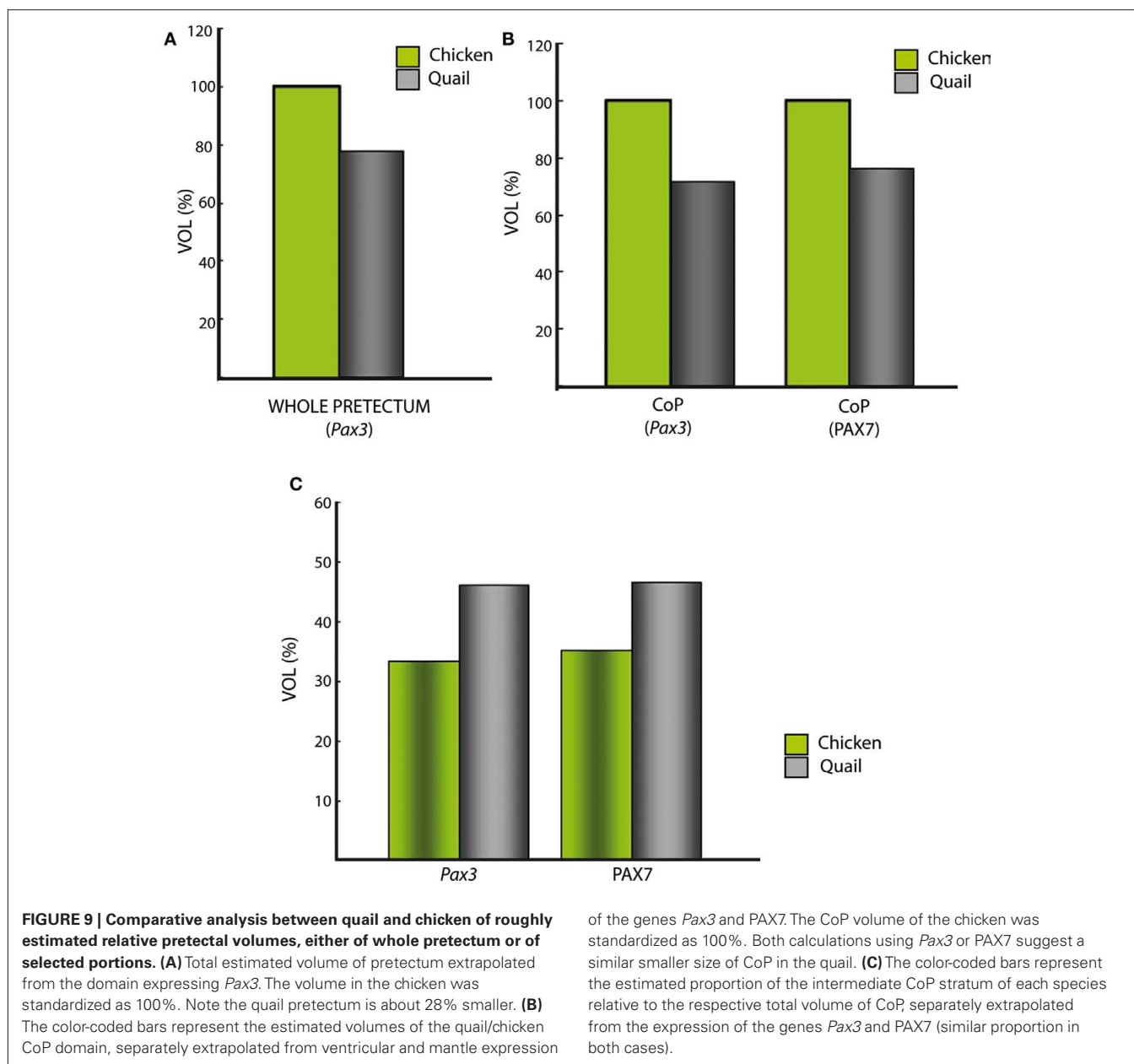
section levels from rostral to caudal from a single specimen (levels arranged top to bottom along the columns). (J–L) Three adjacent transversal sections at CoP level, allowing comparison of *Pax6*, *PAX7*, and *Tcf7l2* dorsoventral patterns in a different quail specimen. The red arrowhead in (L) shows the ventral-most limit of *PAX7* expression, slightly above the alar–basal boundary (ABB; black arrow). Numbers between brackets: code of the specimen in our collection. Scale bar = 300 μm.

**CONSERVED GENOARCHITECTONIC PROFILE OF THE PRETECTAL DOMAIN IN PHASIANIDAE**

We present a detailed comparative characterization of molecularly distinct pretectal progenitor areas from chicken (*G. gallus*) and quail (*C. japonica*) at early stages of mantle development (Q26/HH26–Q28/HH28). We found a strikingly conserved combinatorial genetic code that patterns this region in both species, strongly suggesting that this corresponds to the ancestral pretectal code for members of the family Phasianidae. We had previously defined the molecular code identifying each region, domain and subdomain

(progenitor areas), as well as their respective mantle derivatives, in chicken embryos (Ferran et al., 2007, 2008, 2009). The present study adds information for new gene expression patterns and expands our previous knowledge for some domains, in some cases modifying previously established notions.

The rostral pretectal boundary (TPB) is identified by the rostral limit of expression of *Bhlhb4*, *Ebf1*, *Meis1*, and *Pax3* genes in both species (Ferran et al., 2007, 2008, 2009; present results). Here we confirmed that the thalamic expression of *Gbx2* is a complementary boundary marker for the TPB at the studied stages. Similar thalamic



*Gbx2* expression was previously observed in mouse, chicken (older stages), zebrafish, and *Xenopus* (Bulfone et al., 1993; Niss and Leutz, 1998; Martínez-de-la-Torre et al., 2002; Kikuta et al., 2003); however, the strict complementarity of thalamic *Gbx2* with pretectal *Pax3* was demonstrated only recently in *Xenopus* (Morona et al., 2010). We nevertheless observed as well a patch of *Gbx2* signal within the rostral *pe* stratum of PcP. It can be speculated that this result is due to some thalamic cells migrating tangentially into the neighboring pretectum. This possibility acquires relevance because observations in a mutant mouse line in which the gene *Gbx2* was deleted (*Gbx2*<sup>CreER/-</sup>; R26R mutants) was thought to produce abnormally such a displacement of thalamic cells into the pretectum (Chen et al., 2009). Our present data suggest instead the possibility that such a movement may be constitutive in the wild type.

While *Meis1* delineates the TPB (the p2/p1 interprosomic boundary) across the whole alar plate, up to the diencephalic roof, the *Dbx1*, *Ebf1*, and *Pax3* expression domains present in PcP extend across the pretecto-habenular limit into the habenular region of prosomere 2, as is also true of *Pax6* and *PAX7*, whose CoP domains similarly extend rostralward at this dorsal level (Figures 2A,C,D,E,H,I, 3U,W,Y,AA, and 5A,P; Ferran et al., 2007, 2008, 2009). We think that the primary pattern causing these shared properties of the habenular region and the cited dorsocaudal pretectal regions is probably *Pax6*. At earlier stages this gene is first expressed throughout the diencephalic alar plate. Subsequently it becomes progressively downregulated at both sides of the zona limitans intrathalamica (probably a SHH-mediated effect; Hashimoto-Torii et al., 2003; Kiecker and Lumsden, 2004; Vieira et al., 2005;

Vieira and Martínez, 2006; see Discussion in Ferran et al., 2007). The downregulated *Pax6* expression at the level of the alar domain caudal to the zona limitans involves the thalamus proper in p2 and the PcP and JcP domains in p1; so that *Pax6* expression persists at the habenula, in p2, as well as at dorsal parts of PcP and JcP, and at the entire CoP, in p1, that is, the sites farthest from the zona limitans intrathalamica. This *Pax6* pattern possibly stabilizes *Pax3* and *Pax7* in the same mixed area, with a corresponding distribution of genes such as *Dbx1*, *Ebf1* (*Pax6* transiently upstream of them before it becomes downregulated at PcP). The *Meis1* pattern thus importantly establishes the neuromeric limit at this dorsal level between the thalamic habenular region and the pretectum. The overlapping distribution of pretectal *Meis1* and *Pax3* was also observed for the mouse, and comparable *Pax3* data exist also for *Xenopus* (Toresson et al., 2000; Ferran et al., 2008; Morona et al., 2010). Interestingly, *Pax3* and *Meis* genes have been suggested to be part of a specific regulatory network involved in the development of the rhombencephalon in *Xenopus* (Elkouby et al., 2010; Gutkovich et al., 2010); further studies should address whether a similar relationship of these transcription factors exists in the patterning of the pretectal region of vertebrates.

The caudal pretectal boundary was marked by the expression domains of *Pax6* and *Tcf712* (*Tcf4*; rostral side) and *Meis2* (caudal side) during early chicken development. However, only *Pax6* has been studied extensively throughout development (Ferran et al., 2007, 2008, 2009). Here we found that *Tcf712* (*Tcf4*) is also expressed in the midbrain mantle from stage Q25/HH25 onwards, but its DMB boundary persists at the ventricular zone at the stages analyzed. The same early diencephalic pattern and added mesencephalic expression in the mantle at later stages was observed in the mouse (Ferran et al., in preparation) and *X. laevis* (Morona et al., 2010, and unpublished observations). In *Xenopus*, some authors have misinterpreted early diencephalic *Tcf712* expression as “rostral mesencephalon” (Kunz et al., 2004; Koenig et al., 2008, 2010); nonetheless, comparison with the *Pax6* expression pattern illustrated in the same studies, as well as by Schlosser and Ahrens, (2004), clearly indicates that *Tcf712* expression was only diencephalic (thalamus and pretectum) at those stages. According to the genoarchitectonic data from Schlosser and Ahrens, (2004), the distinction between p1, p2, p3, and midbrain in *Xenopus* is established at stages 24–27, whereas *Tcf712* (*Tcf4*) is clearly expressed only in the diencephalon until at least stages 33–34 (Figure 1 in Morona et al., 2010). *Meis2* expression in the alar mesencephalon, stopping rostrally at the DMB boundary, was described in chicken from stage HH9/10 onward (Ferran et al., 2007; Sánchez-Guardado et al., 2011). In the present study, we identified at Q27/HH28 some *Meis2*-positive cells in the *pe* and *su* strata from the CoP, which progressively increased in number. Those cells lying at the CoP *su* stratum will become part of the lateral and dorsal terminal nuclei of the accessory optic tract at later stages (data not shown), whereas the deep *pe* ones apparently are integrated in the pretectal periaqueductal gray. The progression of *Meis2* mRNA expression suggests that the positive cells at the CoP increased progressively in clear connection with a stream of *Meis2*-positive cells translocating from the mesencephalic TG; a radial migration of the *su* cells is inconsistent with the fact that such cells never are found within the *i* stratum. Our analysis therefore suggests that these cells may be migrating from the TG

at the rostral mesencephalon, unless, alternatively, these cells are selectively upregulating *Meis2* expression independently. The few *Meis2*-positive cells observed in the *pe* stratum at all stages analyzed also show a pattern suggesting a periventricular mesencephalic TG origin. In contrast, various studies using quail/chicken chimeras or lineage analysis after infection with a recombinant retrovirus showed no quail-derived cells migrating from mesencephalon to diencephalon (Senut and Alvarado-Mallart, 1987; Gray et al., 1988; Gray and Sanes, 1991; Martínez et al., 1992; García-López et al., 2004; García-López, 2005). Nearly all of the cited quail/chicken chimera studies involved grafts of the rostral prospective optic tectum, which demonstrated a rostralward tangential migration of given tectal cell types restricted within the tectum itself, but never penetrating the TG, or crossing into diencephalic domains (Senut and Alvarado-Mallart, 1987; Figure 1 in Martínez et al., 1992; García-López et al., 2004; García-López, 2005). Similarly, the lineage studies labeled clones that were restricted to the optic tectum and confirmed strictly tectal radial and tangential migration patterns (Gray et al., 1988; Gray and Sanes, 1991; Ferran, 2002), but no rostral translocation into the diencephalon. Due either to the small size of the TG primordium, or to the traditional tendency to disregard it, no experiment so far has properly tested the fate of TG derivatives. However, alar mesencephalic quail tissue transplanted heterotopically into the chicken diencephalic alar plate produced a massive superficial dispersion of quail cells, which invaded selectively all primary retinorecipient nuclei of the chick diencephalon; it was not determined whether prospective TG tissue was included in these grafts (Martínez and Alvarado-Mallart, 1989). Notably, most of the *Meis2*-positive cells apparently entering the CoP from the TG occupy the superficial stratum. Further studies will be needed to test whether the TG is indeed a source of *Meis2*-positive cells for the pretectum or the observed pattern is a consequence of independent gene expression recruitment by pretectal cells.

At the level of the PcP domain, we similarly found *Gbx2* expression in the *pe* layer that might be related either to migration of thalamic cells (note the signal extending caudalward from the thalamus in Figure 3P'), or to gene expression recruitment. Among available lineage tracing and fate-mapping studies relevant to the TPB, the reports of Figdor and Stern (1993) and García-López et al. (2004) did not identify thalamic clones or derivatives crossing the p1–p2 boundary, whereas Larsen et al. (2001) did observe cell movement between thalamus and pretectum, concluding that this boundary is not clonally restricted. More studies are needed to resolve this controversy. *Pax6* expression starts to appear at the dorsal PcP *di* stratum at HH27, earlier than previously observed by us (shown at HH30/32 in Ferran et al., 2009; see asterisk in their Figures 4J,S). These cells will become part of the core of the medial pretectal nucleus (Figure 10F in Ferran et al., 2009).

The dorsocaudal nucleus (DCa), originally described by Rendahl (1924), and also known as the “medial or dorsomedial spiriform nucleus” (Edinger and Wallenberg, 1899; Kuhlenbeck, 1939; see Table 3 in Ferran et al., 2009) was found to be derived from the PcP domain on the basis of a characteristic genoarchitectonic code (expression of *Bhlhb4*, *Ebf1*, and *Nbea*; Figures 11–14 in Ferran et al., 2009). In our earlier study, we concluded that DCa cells migrate through the JcP domain at stage HH31 (Figures 14A,B in Ferran et al., 2009). In our present analysis, we compared in chicken



and quail markers for PcP (*Meis1*, *Ebf1*, *Bhlhb4*) and JcP derivatives (*Six3*, *Lim1*, *Tal2*, *Gata3*), and corroborated this migration, finding in addition that the first cells that move into the JcP domain constitute a pioneering migratory group departing from the PcP *di* stratum at around Q27/HH27.

At the level of JcP, its characteristic marker *Six3* was previously shown to become secondarily downregulated at a middle locus of the intermediate mantle stratum starting at HH27, while *Lim1* and *Tal2* continue to be expressed in this cluster of cells that also begins to express selectively *FoxP1* and matures as the lateral spiriform nucleus (Ferran et al., 2007, 2009). Here we confirmed the whole pattern for these genes in both species, and added *Gata3* expression to the molecular code characteristic of the cells constituting the prospective lateral spiriform nucleus.

The intermediate mantle layer of CoP was previously described by us to be segregated into *di*, *mi*, and *oi* layers at least from HH30 onward (Figure 4 in Ferran et al., 2009). Here, we provide evidence (combining *Pax3*, *Pax7*, *Gata3*, *Lim1*, and *Tal2* expression patterns) that this segregation of the intermediate mantle stratum actually begins at Q26/HH27 (Figures 5 and 7).

#### MICROEVOLUTION OF CODING SEQUENCES AND PRETECTAL GENOARCHITECTURE IN THE PHASIANIDAE FAMILY

The two species analyzed here belong to the well known avian order Galliformes; however, the phylogenetic relationships within this order are not well resolved yet, primarily due to the low variability in anatomical, and osteological traits, as well as to inconclusive molecular studies (Kimball et al., 1999; Crowe et al., 2006; Krieger et al., 2007; Mayr, 2008; Tavares and Baker, 2008; Kan et al., 2010a,b; Shen et al., 2010). Based on the combined evidence from different phylogenetic analyses, Crowe et al. (2006) put forward a tentative revised classification of the Galliformes. In this classification, the order Galliformes comprises five families (Megapodiidae, Cracidae, Numididae, Odontophoridae, and Phasianidae), and the family Phasianidae encompasses seven subfamilies (Arborophilinae, Coturnicinae, Pavoninae, Gallininae, Meleagridinae, Tetraoninae, and Phasianinae). The Coturnicinae and Gallininae subfamilies include the *C. japonica* and *G. gallus* species, respectively (Figure 1A). The order Galliformes apparently started to evolve 90 million years ago, whereas the split between the families Phasianidae and Odontophoridae occurred ~60 million years ago (Figures 1A,A'; Puelles and Medina, 2002). Analysis of mitochondrial genomes suggested that the subfamilies Phasianinae, Gallininae, and Coturnicinae evolved independently from ~40 million years (Kan et al., 2010b; blue arrow, Figure 1A'). Therefore, the species analyzed here, *C. japonica* and *G. gallus*, have undergone a combined time of divergent evolution of ~80 million years. We sequenced and compared the coding sequences of 12 genes, finding very low rates of genomic variation, consistent with the analyses of mitochondrial genomes. This high level of coding sequence conservation was associated with a striking degree of genoarchitectonic Bauplan conservation during early stages of nervous system development in both species. Since genoarchitectonic codes are expected to be controlled by *cis* regulatory elements, our data also suggest a high level of conservation of the underlying genomic regulatory sequences between both species (Davidson, 2006; Carroll, 2008).

#### PRETECTAL GENOARCHITECTURE IN VERTEBRATES: DEGREE OF PHENOTYPIC CONSERVATION THROUGHOUT ONTOGENY

Our previous studies on the genoarchitecture of the chicken pretectum have provided a starting point for comparing this region at an unprecedented level of detail among different vertebrates (Ferran et al., 2007, 2009). The present results have shown that chicken and quail pretectal genoarchitecture patterns are strikingly conserved, thus likely representing an ancestral pattern within the Phasianidae. Outgroup comparisons with other species, such as the avian superorder Neognathae (ostriches; Figure 1A), and archosaurian reptiles (e.g., a crocodile) or lepidosaurians (e.g., lizards), should corroborate whether the observed pretectal genoarchitectonic profile is plesiomorphic in all birds and reptiles. Furthermore, there is already evidence that the same molecular map and a structural anteroposterior tripartition exists in the pretectum of the mouse and *Xenopus* frog (Ferran et al., 2008; Morona et al., 2010), suggesting that the referred plesiomorphy may apply as well to tetrapods in general. A tripartite pretectum has been postulated as well for lamprey larvae (Pombal and Puelles, 1999; Pombal et al., 2009), which suggest we deal here with a fundamental aspect of forebrain structure in all vertebrates. Some elements of the studied pretectal molecular code were, in fact, used in *Xenopus* larvae to identify precisely the pretectal region, and its main subdivisions (Morona et al., 2010) working within the context of field homology in the brain (Puelles and Medina, 2002). The present results also reveal that the pattern of radial segregation of molecularly characterized mantle components (pronuclei) described for the chicken (Ferran et al., 2007, 2009) are most likely representative for all members of the family Phasianidae. This is independent of the different absolute sizes of the respective populations and domains in chicken and quail, or of the heterochronic aspects of maturation commented above. Although late stages of differentiation are commonly thought to present more phenotypic variations (Davidson, 2006), the nuclear anatomy of the adult pretectum shows little variation among studied adult birds (Karten and Hodos, 1967; Zweers, 1971; Kuenzel and Masson, 1988; Puelles et al., 2007). Our results nevertheless allow making very detailed molecular comparative analyses at the level of strata segregation or differentiation of distinct neuronal subpopulations among different bird species, or between birds and other vertebrates, thus exploring any significant variations. For example, our study in *Xenopus* disclosed that at early stages PAX7 was observed in the mantle zone of the JcP domain, an expression trait that we did not find in chicken or quail (Morona et al., 2010; Ferran et al., 2007, 2009; present results).

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