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RESEARCH ARTICLE

Cell interactions, signals and transcriptional hierarchy governing placode progenitor induction

Mark Hintze, Ravindra Singh Prajapati, Monica Tambalo*, Nicolas A. D. Christophorou[‡], Maryam Anwar[§], Timothy Grocott[¶] and Andrea Streit^{**}

ABSTRACT

In vertebrates, cranial placodes contribute to all sense organs and sensory ganglia and arise from a common pool of *Six1/Eya2*+ progenitors. Here we dissect the events that specify ectodermal cells as placode progenitors using newly identified genes upstream of the *Six/Eya* complex. We show in chick that two different tissues, namely the lateral head mesoderm and the prechordal mesendoderm, gradually induce placode progenitors: cells pass through successive transcriptional states, each identified by distinct factors and controlled by different signals. Both tissues initiate a common transcriptional state but over time impart regional character, with the acquisition of anterior identity dependent on *Shh* signalling. Using a network inference approach we predict the regulatory relationships among newly identified transcription factors and verify predicted links in knockdown experiments. Based on this analysis we propose a new model for placode progenitor induction, in which the initial induction of a generic transcriptional state precedes regional divergence.

KEY WORDS: Chick embryo, Quail graft, Cell fate, Sense organs, Sensory ganglia, Transcriptional networks, Signalling, Gene regulatory network

INTRODUCTION

During early development, many fate decisions are controlled by inductive interactions, whereby an inducing cell population instructs responding cells to change their fate. Neural induction is perhaps the best-studied inductive event, and several models have been proposed to explain the induction and patterning of the nervous system by the organizer (Stern, 2001). The ‘multiple organizer’ model argues that different parts of the organizer induce distinct regions of the nervous system (Holtfreter, 1933a,b; Mangold, 1933; Saxen and Toivonen, 1962), whereas Waddington suggested a two-step model with ‘evocation’ generating a generic, non-regionalised nervous system, followed by ‘individuation’ to impart regional character (Waddington and Needham, 1936). Nieuwkoop’s ‘activation-transformation’ hypothesis suggests that initial induction generates anterior

character, which is then posteriorised (Nieuwkoop and Nigtevecht, 1954). Currently, there is support for modified versions of both Nieuwkoop’s and Mangold’s models (Stern, 2001; Takemoto et al., 2006). However, the question remains whether these models could also explain the development of adjacent structures – the sensory placodes. Placodes arise in the head ectoderm in register with the neural tube and generate components of sense organs and cranial sensory ganglia (Baker and Bronner-Fraser, 2001; Grocott et al., 2011; Schlosser, 2010; Streit, 2008). Pushing the multiple organizer model to the extreme, it has been suggested that even placodes are induced by a subset of organizer cells (Mangold, 1933; Saxen and Toivonen, 1962; Spemann, 1938). Here we ask whether placodes are induced by a bona fide organizer that induces and patterns the placodal territory, and if so, which if any of these models explains this process.

Although contributing to diverse organs and ganglia, the cranial placodes arise from a common domain, termed the pre-placodal region (PPR), which is specified at head process stages as a strip of ectoderm surrounding the anterior neural plate (Baker and Bronner-Fraser, 2001; Grocott et al., 2012; Schlosser, 2010; Streit, 2008). PPR cells initially have the same developmental potential: they can give rise to any placode and express a common set of genes (Bailey et al., 2006; Bailey and Streit, 2006; Streit, 2008), among them *Six* and *Eya* factors, which impart PPR character to ectodermal cells (Brugmann et al., 2004; Chen et al., 2009; Christophorou et al., 2009; Laclef et al., 2003; Zheng et al., 2003; Zou et al., 2004, 2006). Surprisingly, few upstream regulators have been identified to explain how their expression is activated in, and confined to, sensory progenitors. *Six1* transcription is directly activated by *Dlx* factors, but repressed by *Msx1* (Sato et al., 2010). At gastrulation stages, BMP signalling is required for the PPR ‘competence factors’ *Gata2/3*, *Tfap2a/c*, *Foxi1* and *Dlx3*, but must be inhibited later for *Six1* activation (Ahrens and Schlosser, 2005; Brugmann et al., 2004; Kwon et al., 2010; Litsiou et al., 2005; Pieper et al., 2012). Subsequently, FGF signalling together with inhibition of BMP and Wnt is necessary for PPR formation and sufficient to induce sensory progenitors in non-placodal ectoderm (Brugmann et al., 2004; Litsiou et al., 2005). These signals emanate from the adjacent neural plate and the underlying head mesoderm (Ahrens and Schlosser, 2005; Litsiou et al., 2005), although their relative contribution to PPR induction remains unclear, as does the question of whether the signalling tissues are true organizers.

Using an established induction assay, we now dissect PPR induction and propose a new model. First, we design a molecular screen to identify new potential players and then use these factors to characterise the response to each PPR-inducing tissue and to establish a genetic hierarchy upstream of the *Six/Eya* complex. Combined with a network inference approach, our analysis proposes a new multistep model for PPR induction. We show that signals from the neural plate are unable to induce a PPR. Two different mesodermal populations, namely the lateral head mesoderm and

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the prechordal mesendoderm, initially induce a similar set of transcription factors (reminiscent of an ‘evocation’), but then gradually impart anterior and posterior bias to sensory progenitors (‘individuation’). We show that PPR induction is not mediated by an organizer, but instead involves multiple signalling centres, each inducing cells with distinct regional character.

RESULTS

A molecular screen reveals the complexity of PPR induction

In chick, PPR induction is mediated by signals from the head mesoderm: when grafted next to competent epiblast this mesoderm induces a full set of bona fide PPR markers (Litsiou et al., 2005; see Fig. 2A,B). How long does this process take? In the same induction assay, quail lateral head mesoderm (IHM) requires at least 12 h of contact to induce the PPR markers *Six1* and *Eya2* in chick epiblast [Fig. 1A-B'; *Six1*: 4 h, 2/9; 8 h, 8/8; *Eya2*: 8 h, 0/7; 12 h, 3/10; *Six4* (not shown): 4 h, 2/9; 8 h, 10/12].

To identify the components of the PPR induction cascade, we performed a microarray screen comparing the transcriptomes of epiblast from four conditions (Fig. 2C): mesoderm-induced epiblast (12 h; MIE), non-induced epiblast (control; NIE) from the contralateral side of the same embryos, and the normal anterior (aPPR) and posterior (pPPR) PPR at HH5/6 (Lleras-Forero et al., 2013). One-way ANOVA revealed 3475 probes with a significant change of more than 2-fold ($P < 0.05$) in at least one cell population, representing 2868 unique transcripts including 206 known or putative transcription factors (Table S1, GSE81023). When compared with control epiblast, 1098 transcripts are mesoderm induced (>1.2 -fold), whereas 1379 genes are repressed (Table S1). As expected, the PPR-enriched factors *Eya2*, *Dach1* and *Sox3* (Rex et al., 1977; Barembaum and Bronner-Fraser, 2007; Litsiou et al., 2005) are induced by the IHM, whereas neural crest (*Snail2*), neural plate (*Sox2*) and regionally restricted genes (*Pax6*) are not (Litsiou et al., 2005) (Fig. S1). Thus, the array screen replicates known changes in gene expression in response to mesodermal signals.

To ensure that the genes identified were induced by mesoderm rather than recruited from the host PPR, the screen was designed using extraembryonic tissue. However, for any gene to be relevant to PPR formation it should at some point be expressed in the prospective PPR. To identify synexpression groups we performed hierarchical clustering on all genes that changed significantly and *in situ* hybridisation to verify their spatiotemporal expression in normal embryos. In total, we assessed the *in situ* expression of 47 known or putative transcription factors, five chromatin modifiers and four signalling pathway components.

This analysis reveals seven major clusters with distinct profiles (C1-C7; Figs S1, S2 and S3, Table S4). C1 and C2 transcripts are largely absent from placode progenitors, like *Cux1* (C2; Fig. 2D'), and repressed by the mesoderm, with C2 genes being enriched in the extraembryonic ectoderm (Fig. 2D). Cluster C5 transcripts (Fig. 2F) are strongly enriched in the aPPR, but only weakly mesoderm induced, and include aPPR-specific genes such as *Pax6* and *Sstr5* (Lleras-Forero et al., 2013), *Nfkb1* (Fig. 2F') and *Sall1* (Fig. S2). By contrast, cluster C6 transcripts (Fig. 2G) are IHM induced, enriched in the pPPR and include *Gbx2* (Fig. 2G'), *Irx2* and *Pax7* (Goriely et al., 1999; Khudyakov and Bronner-Fraser, 2009; Steventon et al., 2012) and novel transcripts such as *Znf423*, *Znf76* and *Rnf24*. Finally, cluster C4 genes are mostly induced by mesodermal signals and are present throughout the entire PPR. *In situ* hybridisation reveals that most C4 transcripts are not restricted to sensory progenitors, but expressed broadly at primitive streak stages encompassing the future neural, neural crest and placode

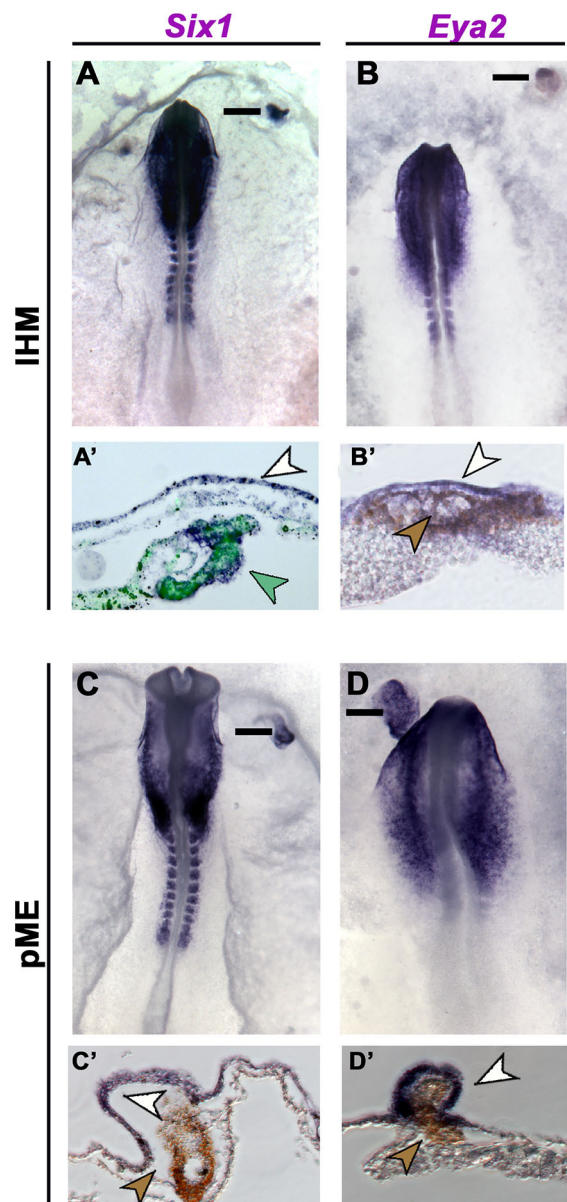


Fig. 1. Induction of placode progenitors by the lateral head mesoderm (IHM) and prechordal mesoderm (pME). (A-B') CMFDA-labelled chick (green arrowhead in A') or quail IHM (brown arrowhead in B') from HH5/6 donors induces *Six1* (A,A', white arrowhead; 8 h, 8/8) and *Eya2* (B,B', white arrowhead; 12 h, 3/10) in host extraembryonic epiblast. (C-D') pME grafts from HH5/6 quail donors (brown arrowheads) induce *Six1* (C,C', white arrowhead; 15-17 h, 5/8) and *Eya2* (D,D', white arrowhead; 15-17 h, 7/8). Bars in A-D indicate section levels in A'-D'.

territories (Fig. 2E,E', Figs S2 and S3), including *Zic1* (Khudyakov and Bronner-Fraser, 2009), *Otx2* (Bally-Cuif et al., 1995) and *Fzd8* (Paxton et al., 2010) and many new genes (Figs S2 and S3). Only a few C4 transcripts are restricted to the PPR (*Dmbx1*, *Homer2*; Fig. S3). As development proceeds, some transcripts from cluster C4 remain expressed in both neural and placode cells, whereas others become confined to either tissue (Figs S2 and S3), suggesting that *Six* and *Eya* factors are among the few bona fide PPR markers.

Thus, the screen has identified many novel transcripts expressed in placode progenitors, revealing molecular similarity between precursors for the central and peripheral nervous systems. In

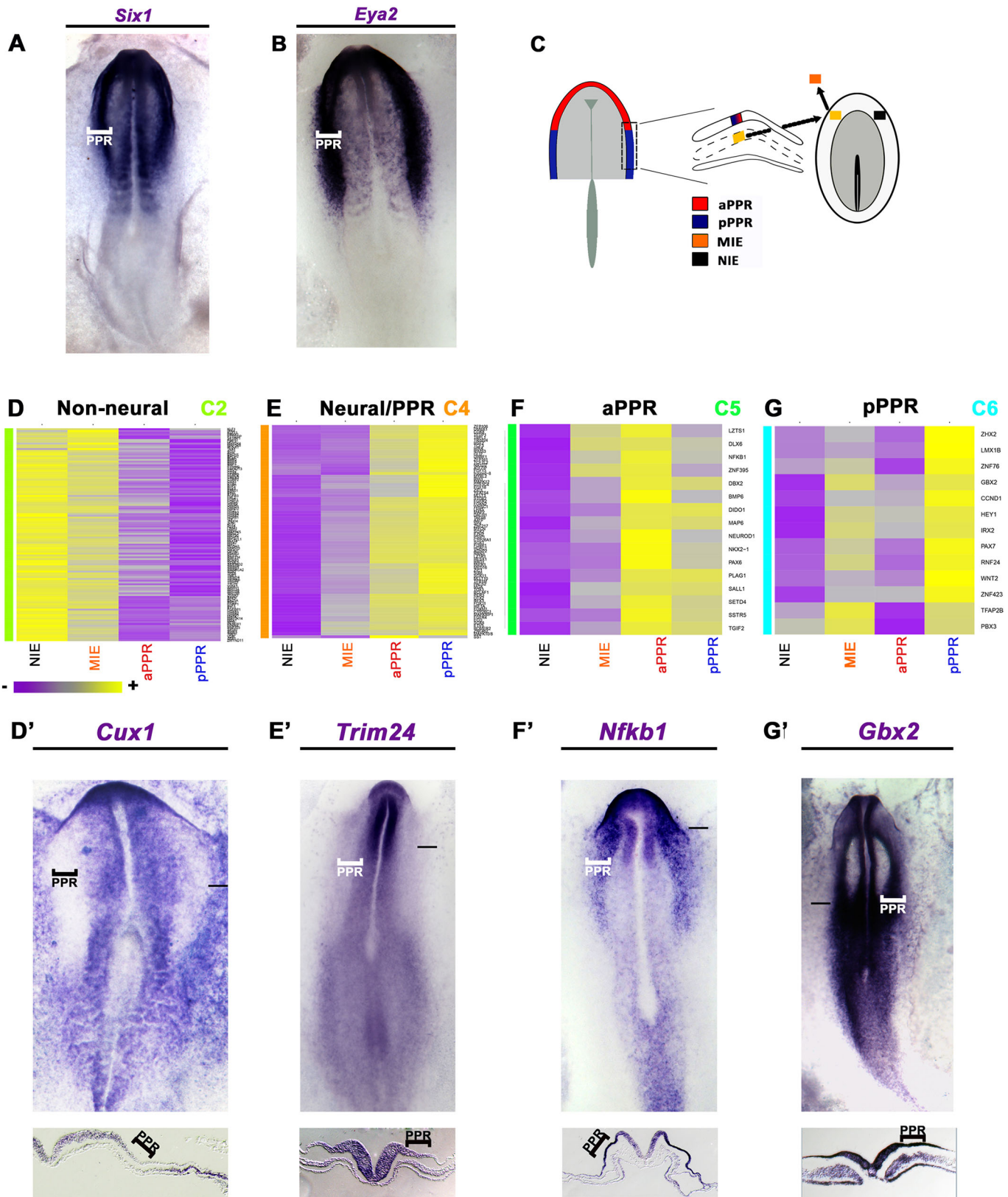


Fig. 2. New placode progenitor transcripts. (A,B) *Six1* (A) and *Eya2* (B) expression at HH7⁺8⁻. (C) aPPR and pPPR from HH6 (Lleras-Forero et al., 2013) and mesoderm-induced (MIE) and non-induced ectoderm (NIE) were analysed by microarray. (D) Cluster C2 transcripts are excluded from the PPR, such as *Cux1* (D'). (E) Cluster C4 includes aPPR- and pPPR-enriched factors, such as *Trim24* (E'). (F) Cluster C5 includes aPPR factors, such as *Nfkb1* (F'). (G) Cluster C6 includes pPPR factors, such as *Gbx2* (G'). Bars in D'-G' indicate level of sections shown beneath.

addition, ultimately the IHM appears to induce posterior placode progenitors, but does not act as an organizer that induces and patterns the PPR. This implies that other PPR-inducing tissues exist, which should impart rostral PPR identity.

Prechordal mesendoderm induces placode progenitors

Endodermal signals have been implicated in anterior placode induction in amphibians (Henry and Grainger, 1990; Jacobson, 1963a,b) and we have recently shown that the anterior prechordal mesendoderm (pME) is required for *Eya2*, *Pax6* and *Pnoc* expression in chick and zebrafish (Lleras-Forero et al., 2013). To test its PPR-inducing ability, we grafted quail HH5 pME next to chick extraembryonic epiblast (HH3^{+/4}⁻). Both *Six1* and *Eya2* are induced after 15–17 h (Fig. 1C–D'). Thus, two different mesodermal tissues, namely IHM and pME, mediate PPR induction.

Lateral and axial mesoderm provide regional bias to sensory progenitors

Our results indicate that PPR induction is not mediated by an organizer, although it remains possible that IHM first induces aPPR, which is subsequently posteriorised. Alternatively, both mesoderms might first induce generic PPR character, which is then regionalised, or that each directly induces distinct rostrocaudal identity. To distinguish these possibilities and to capture the complexity of PPR induction, we compared the response to both tissues over time assessing 126 genes simultaneously. HH5/6 IHM or pME was grafted into the extraembryonic region of HH4⁻ hosts and the underlying epiblast was collected after 3, 6 and 12 h, together with the contralateral control epiblast. Each experiment was performed in triplicate and gene expression was analysed by NanoString using a probe set containing known and new PPR transcripts, markers for different placodes, neural crest and neural plate (Table S2). Transcripts exhibiting a difference greater than 1.2-fold ($P < 0.05$) between induced and time-matched control tissues were considered as differentially expressed.

The transcriptional hierarchy in response to the IHM

IHM-derived signals initiate a dynamic response, with distinct groups of transcripts being regulated at each time point (Fig. 3A–E, Table S3). At 3 h, only a few factors are upregulated: *Cnd1*, *Etv5*, *ERNI*, *N-myc*, *Otx2*, *Sall4*, *Trim24* and *Zic3* (Fig. 3B,C',C''). Whereas some genes are maintained, others are only transiently induced: *Otx2* expression levels are negligible (Fig. S4) and *Sall4* disappears after 6 h, as do *ERNI* and *Zic3* after 12 h (Fig. 3B). At 6 h, new transcripts appear in response to the mesoderm (*Dnmt3b*, *Pdlim4*, *Irx2*, *Rybp*, *Stox2*, *Znf462*; Fig. 3B,D',D''), Table S3), whereas PPR genes (*Six1/4*, *Eya2*) are only upregulated after 12 h together with some of their known upstream regulators (*Gata3*, *Foxi3*, *Tfap2a*, *Dlx6*; Fig. 3B,E',E'') (Kwon et al., 2010; Pieper et al., 2012; Qiao et al., 2012; Sato et al., 2010) and genes previously not associated with PPR induction (*Aatf*, *Bcl7a*, *Zhx2*; Fig. 3B, Table S3). By contrast, other genes are downregulated by the IHM (3 h), including *Tfap2a*, *Dlx5*, *Gata2* and *Axin2* (Fig. 3B, Table S3). At 12 h, most of these remain absent and other PPR repressors (*Msx1*, *Bmp4*) (Ahrens and Schlosser, 2005; Litsiou et al., 2005; Sato et al., 2010) are reduced; however, *Tfap2a* is now induced and *Dlx5* is no longer repressed, consistent with their role as positive *Six1* regulators (Kwon et al., 2010; Qiao et al., 2012) (Fig. 3B, Table S3).

To determine the character of the IHM-induced cells we examined the normal expression patterns of all genes in each cohort, considering their dynamic changes and that, at different times of development, they may characterise different cell

populations (summarised in Fig. S2 and Table S4). This analysis reveals that 3 and 6 h induced transcripts are normally expressed early and broadly: at HH3–4 their domains encompass the presumptive neural plate and its border, including future neural crest and placodes (Fig. 3C,C'',D,D''), Figs S2 and S3, Table S4), except for *ERNI* and *Otx2*, which are already expressed in the pre-streak epiblast and then label much of the epiblast and the anterior ectoderm, respectively (Bally-Cuif et al., 1995). Most factors are maintained in the neural plate and/or PPR at head fold stages (HH6/7; Figs S2 and S3, Table S4). By contrast, most 12 h induced transcripts first appear at HH5/6 (*Six1*, *Six4*, *Eya2*) or begin broadly in the non-neural ectoderm (*Gata3*, *Dlx6*, *Homer2*, *Foxi3*) to become confined to, or upregulated in, the PPR (Fig. 3E,E'', Table S4). Most mesoderm-induced genes are present in all placode progenitors; however, *Foxi3* (Khatri et al., 2014) and *Gbx2* (Steventon et al., 2012) are restricted to future otic and epibranchial cells by the time PPR markers are expressed; *Otx2* remains present, although at extremely low levels (Fig. S4). Thus, at PPR stages the normal expression of all 12 h induced genes overlaps in the pPPR, suggesting that the induced tissue has acquired pPPR character.

Together, these data reveal that the IHM gradually induces PPR identity, with cells passing through sequential states. The first step does not generate anterior character but resembles pre-primitive streak or early streak stage epiblast, and a pPPR is established over time.

The transcriptional hierarchy in response to the pME

We observe a similar hierarchy in response to the pME. 3 h after grafting, the mesendoderm induces the same transcripts as the IHM (Fig. 4A,B,C',C''), Table S3), with most genes being maintained until at least 12 h. The mesendoderm appears to induce *Irx1*, *Gbx2* and *Sstr5*, albeit at extremely low levels (Fig. 4B, Figs S4 and S5, Table S3). This is followed by induction of 17 transcripts including PPR genes (*Six1*, *Eya2*, *Irx1*, *Homer2*; Fig. 4B,D',D'') and, finally, by many genes at 12 h including the *Six1* co-factor *Dach1* (Fig. 4B, E',E'', Table S3). The pME represses the same genes as the IHM, although the timing varies slightly (compare Fig. 3B with Fig. 4B).

In normal embryos, 3 h pME-induced genes are expressed widely at HH3/4, labelling the future neural plate and its border (Fig. 4C,C'',D, Figs S2 and S3, Table S4). 6 h induced genes overlap in the PPR (Fig. 4D,D''), although two neural transcripts (*Sox2*, *Znf423*) are also present (Table S4). The expression domains of most 12 h induced genes continue to overlap in the PPR and the presence of *Otx2*, *Pnoc*, *Sstr5* and *Six3* suggests that cells have acquired anterior character (Fig. 4E,E''). However, in addition, seven induced genes are expressed in the neural plate in normal embryos (Table S4). Thus, the pME rapidly induces aPPR, but over time generates tissue of mixed (neural/pre-placodal) anterior identity. The delayed induction of neural character excludes the possibility that an induced neural plate contributes to the accelerated induction of PPR transcripts by axial mesendoderm versus IHM. In summary, like the IHM, the pME initiates a sequence of transcriptional responses until PPR identity is established. Both tissues initially induce a small set of common transcription factors (Fig. S5), but subsequently impart anterior and posterior bias.

Neural plate signals are not sufficient to induce placode progenitors

The neural plate has been implicated in PPR induction in *Xenopus* (Ahrens and Schlosser, 2005), although this does not appear to be the case in chick (Litsiou et al., 2005). We now revisit this question

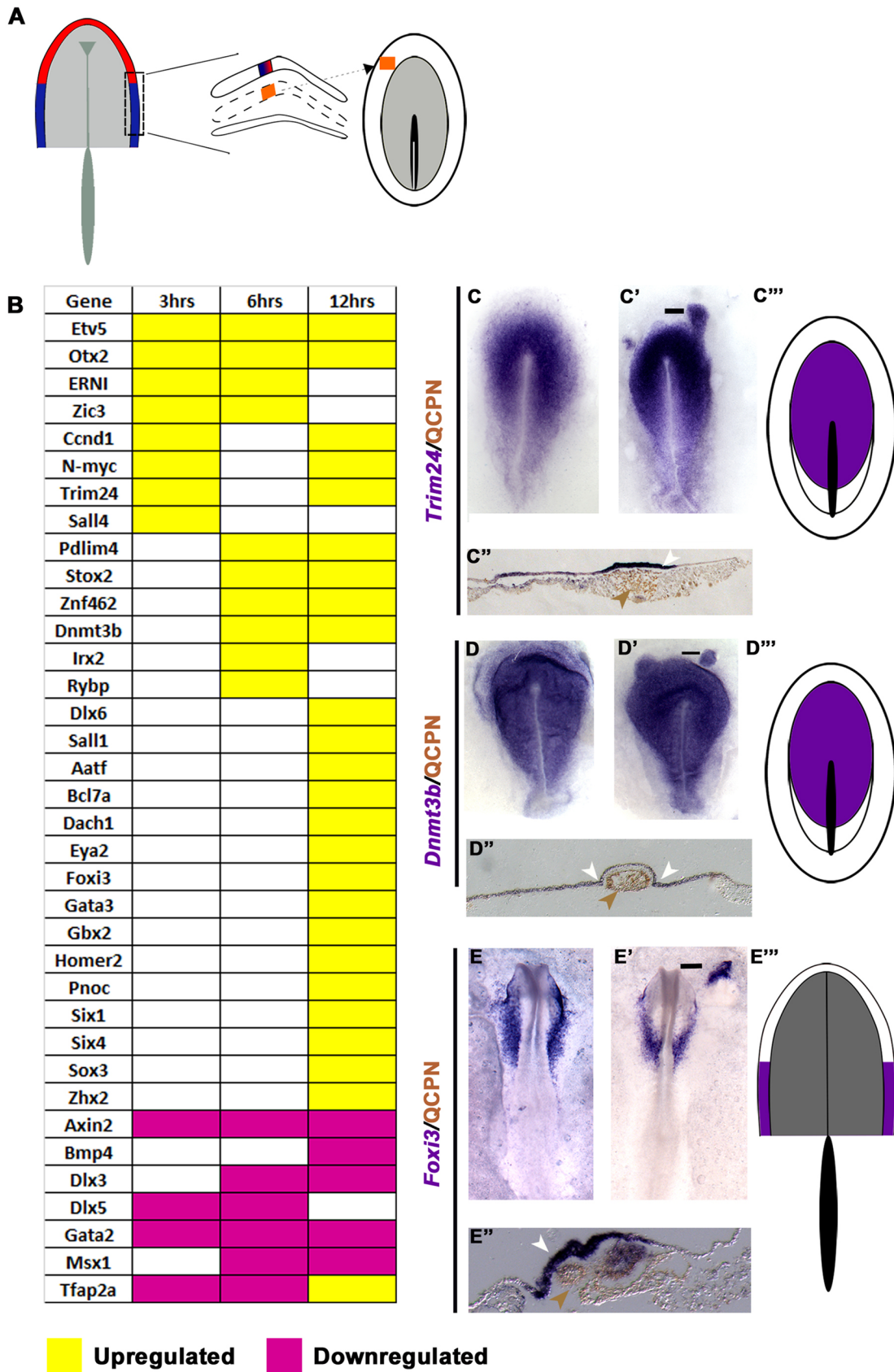


Fig. 3. Response to the IHM. (A) IHM (HH6; orange) was grafted into HH4⁻ hosts; the adjacent ectoderm was analysed after 3, 6 and 12 h. (B) Transcripts induced or repressed ($P < 0.05$; > 1.2 -fold change). (C-E'') Expression patterns of 3 h (C-C''), 6 h (D-D'') and 12 h (E-E'') induced genes. Quail IHM (brown arrowheads) induces *Trim24* (C', C'' white arrowhead), *Dnmt3b* (D', D'' white arrowheads) and *Foxi3* (E', E'' white arrowhead). Bars in C'-E' indicate the level of the sections shown in C''-E''. Area of overlap (purple) is summarised for 3 h (C''), 6 h (D'') and 12 h (E'') induced transcripts.

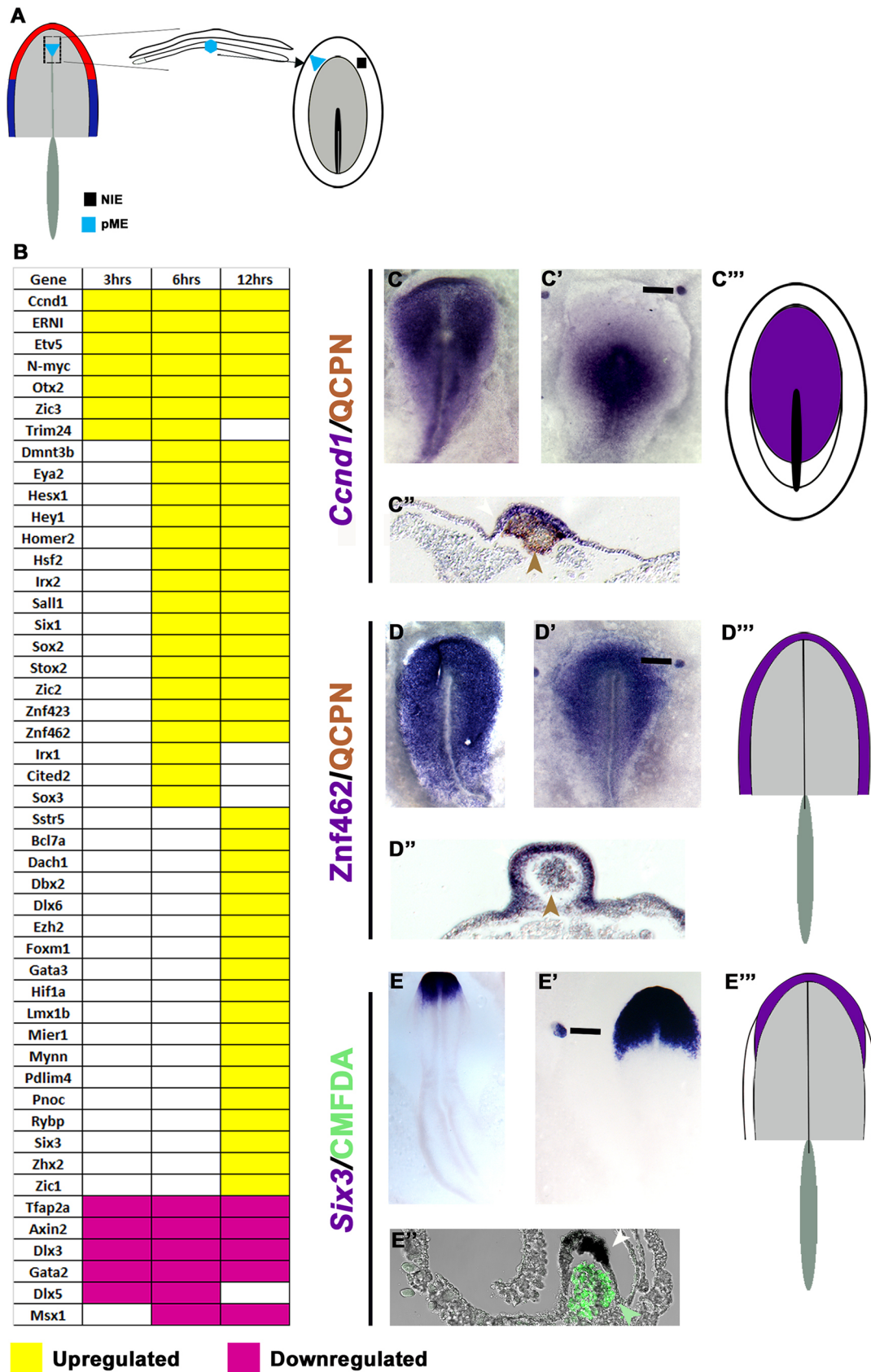


Fig. 4. Response to the pME. (A) pME (HH6) was grafted into HH4⁻ hosts; the adjacent ectoderm was analysed after 3, 6 and 12 h. (B) Transcripts induced or repressed ($P < 0.05$; > 1.2 -fold change). (C-E''') Expression patterns of 3 h (C-C'''), 6 h (D-D''') and 12 h (E-E''') induced genes. Quail pME (brown or green arrowheads) induces *Ccnd1* (C', C'' white arrowhead), *Znf462* (D', D'' white arrowhead) and *Six3* (E', E'' white arrowhead). Bars in C'-E' indicate the level of the sections shown in C''-E'''. Area of overlap (purple) is summarised for 3 h (C'''), 6 h (D''') and 12 h (E''') induced transcripts.

using our newly identified genes. The future forebrain (aNP) or hindbrain (pNP) from HH5/6 donors was grafted into the extraembryonic region of HH4⁻ hosts; epiblast exposed to neural plate signals was dissected together with non-induced epiblast from the contralateral side after 3, 6 and 12 h and processed for NanoString analysis (Fig. S6, Table S3).

After 3 h, the aNP induces many newly identified transcripts, as well as FGF mediators (*Etv4*, *Etv5*) and the neural crest marker *Pax7* (Fig. S6). Except for *Stox2* and *Mynn*, which are absent from the most central epiblast, in the embryo all other factors are expressed broadly at HH3/4 in the future neural plate and its border (Figs S2 and S3, Table S4). Most aNP-induced genes are not among those induced by mesoderm and are only transiently activated. Likewise, five 6 h induced transcripts are broadly expressed in the HH4 epiblast, while three (*Hey1*, *Foxi3*, *Six1*) are confined to the PPR. This is followed by many genes that, in the normal embryo, do not overlap in a single territory, including NP-specific transcripts (HH5/6; *Sox2*, *Znf423*) and non-neural ectoderm genes (*Gata3*, *Pax6*, *Irx1*, *Eya2*). The PPR marker *Six1* is no longer induced, while other PPR-specific genes never respond to aNP grafts (*Six4*, *Homer2*). Finally, the aNP represses known *Six1* regulators (*Dlx5*, *Tfap2a*) (Kwon et al., 2010; Qiao et al., 2012; Sato et al., 2010). Therefore, aNP-derived signals induce a mixture of neural and neural plate border cells rather than a unique territory and cannot initiate a complete PPR.

Similar results are observed in response to pNP-derived signals (Fig. S6). After 3 h, 28 genes are initiated, which are expressed widely in the normal embryo at HH3/4 (Table S4), except the neural plate border gene *Irx1*. Many transcripts continue to be present 6 and 12 h after pNP grafting, whereas 11 transcripts are only transiently induced. This is followed by neural plate (*Sox2*, *Zic2*, *Znf423*) and/or border genes (HH4^{+/5}), with *Eya2* being the only pNP-induced PPR marker. Simultaneously, non-neural markers and *Six1* regulators (*Gata2*, *Dlx3/5*, *Tfap2a*) are repressed. Thus, the pNP induces a territory of mixed identity including many neural and non-neural transcripts, with *Gbx2* and *Irx2* indicating a posterior bias. Thus, like the aNP, the pNP induces different cell populations or cells with mixed identity (neural/placodal). Initially, aNP and pNP grafts induce distinct transcripts, whereas there is significant overlap later (Fig. S6). In summary, the neural plate induces many genes, but unlike the mesoderm it cannot induce the full transcriptional profile characteristic of placode progenitors.

Integration of signalling pathways during PPR induction

FGFs together with BMP and Wnt antagonists have previously been implicated in PPR induction (Brugmann et al., 2004; Kwon et al., 2010; Litsiou et al., 2005). Having identified distinct transcriptional states as cells adopt sensory precursor identity, we can now dissect the role of each signal at different time points by combining mesoderm grafts with pathway manipulation.

FGF initiates PPR induction

FGF signalling is required for PPR induction (Litsiou et al., 2005). To test which genes are FGF induced, we grafted FGF8-coated beads into the HH4⁻ extraembryonic region and analysed FGF8-exposed and control epiblast gene expression after 3 and 6 h using NanoString. After 3 h, FGF8 induces a few genes (*Etv5*, *Trim24*, *Ccnd1*, *ERNI*, *N-myc*, *Sall4*; Fig. 5D, Fig. S7), largely overlapping with IHM-induced (6/8) and pME-induced (5/7) transcripts. *ERNI* is already known to be modulated by FGF8 (Streit et al., 2000). At 6 h, *Etv5*, *Ccnd1*, *N-myc* and *Trim24* continue to be upregulated, as are some additional transcripts (*Stox2*, *Cited2*, *Znf462*; Fig. 5D,

Fig. S7). We confirmed the induction of *Trim24* (Fig. 5A,A''; 4/4) and *N-myc* (not shown; 6/9) by *in situ* hybridisation. By contrast, FGF8 – like the mesoderm – represses *Tfap2a*, *Dlx3*, *Dlx5*, *Gata2* and *Axin2* (Fig. 5D, Fig. S7, Table S3).

To assess the requirement of FGF signalling we combined lateral or axial mesoderm grafts with control beads or beads coated with the FGF receptor antagonist SU5402 (Fig. S7, Table S3). Both tissues induce *Etv5* and *N-myc*; however, the requirement for FGF signalling differs: although FGF signalling is necessary for *Etv5* induction by the IHM (Fig. 5B,B'') it is not required for its induction by the pME, whereas the opposite is true for *N-myc* (Fig. 5D,E, Fig. S7, Table S3). *ERNI* induction by the IHM requires FGF signalling as do pME-induced genes (*Sox3*, *Homer2*) and the neural genes *Sox2*, *Zic2* and *Zic3* (Fig. 5D,E, Fig. S7, Table S3). Finally, FGF signalling is necessary for *Gata2* repression by both tissues, for *Tfap2a* suppression by the IHM, and for inhibition of *Msx1* and *Axin2* by the pME (Fig. 5D,E, Fig. S7, Table S3).

Together, these results show that FGF signalling is sufficient to induce or repress many genes that characterise the earliest response to mesoderm-derived signals (Fig. 5D,E, Fig. S7). However, only a few transcripts strictly depend on FGF, suggesting that each tissue contains other signals that can compensate for FGF loss.

BMP antagonism is required throughout PPR induction

The role of BMP signalling in PPR formation changes over time: during gastrulation it is required to induce *Six/Eya* regulators, but subsequently must be reduced to allow the emergence of sensory progenitors (Ahrens and Schlosser, 2005; Brugmann et al., 2004; Kwon et al., 2010; Litsiou et al., 2005). To test which mesoderm-response genes change after BMP pathway modulation, we reduced BMP signalling by growing primitive streak stage embryos in the BMP inhibitor dorsomorphin. After 3 and 6 h of culture, 11 transcripts are upregulated in PPR-competent epiblast when compared with stage-matched controls (Table S3). However, of these only *Trim24* is induced by both mesodermal tissues (see Fig. 3C').

To test which mesoderm-induced genes require BMP antagonism, we compared the inducing ability of mesoderm alone or mesoderm together with BMP4-coated beads after 3 and 6 h. BMP activation leads to the loss of *Trim24*, *Ccnd1*, *Sall4*, *Otx2*, *ERNI* and *Zic3* induction by the IHM (Fig. 5D, Fig. S7, Table S3); we confirmed the reduction of *Trim24* by *in situ* hybridisation (Fig. 5C,C''). Of the 23 pME-induced transcripts, ten are repressed after BMP activation, including *Trim24*, *N-myc* and the PPR genes *Six1* and *Eya2* (Fig. 5E, Fig. S7, Table S3). Likewise, the downregulation of some transcripts by the IHM is sensitive to BMP activation: *Dlx3*, *Dlx5* and *Gata2* increase when compared with control grafts (Fig. 5D, Fig. S7). By contrast, the pME continues to suppress *Dlx* genes even at elevated BMP levels, while *Msx1* and *Axin2* inhibition requires BMP antagonism (Fig. S7). Therefore, although BMP inhibition alone is sufficient to activate only a few mesoderm-induced genes, many transcripts require BMP antagonism, among them both early and late mesoderm-response genes. In addition, these findings highlight that although both mesoderm populations induce similar sets of transcription factors, each tissue provides a distinct signalling environment where other signals can compensate for the loss of a particular pathway.

Wnt antagonism is not required during early PPR induction

At the border of the neural plate increased levels of Wnt signalling promote neural crest formation at the expense of PPR, whereas Wnt antagonists have the opposite effect (Litsiou et al., 2005). To test which transcripts respond to Wnt modulation at different time

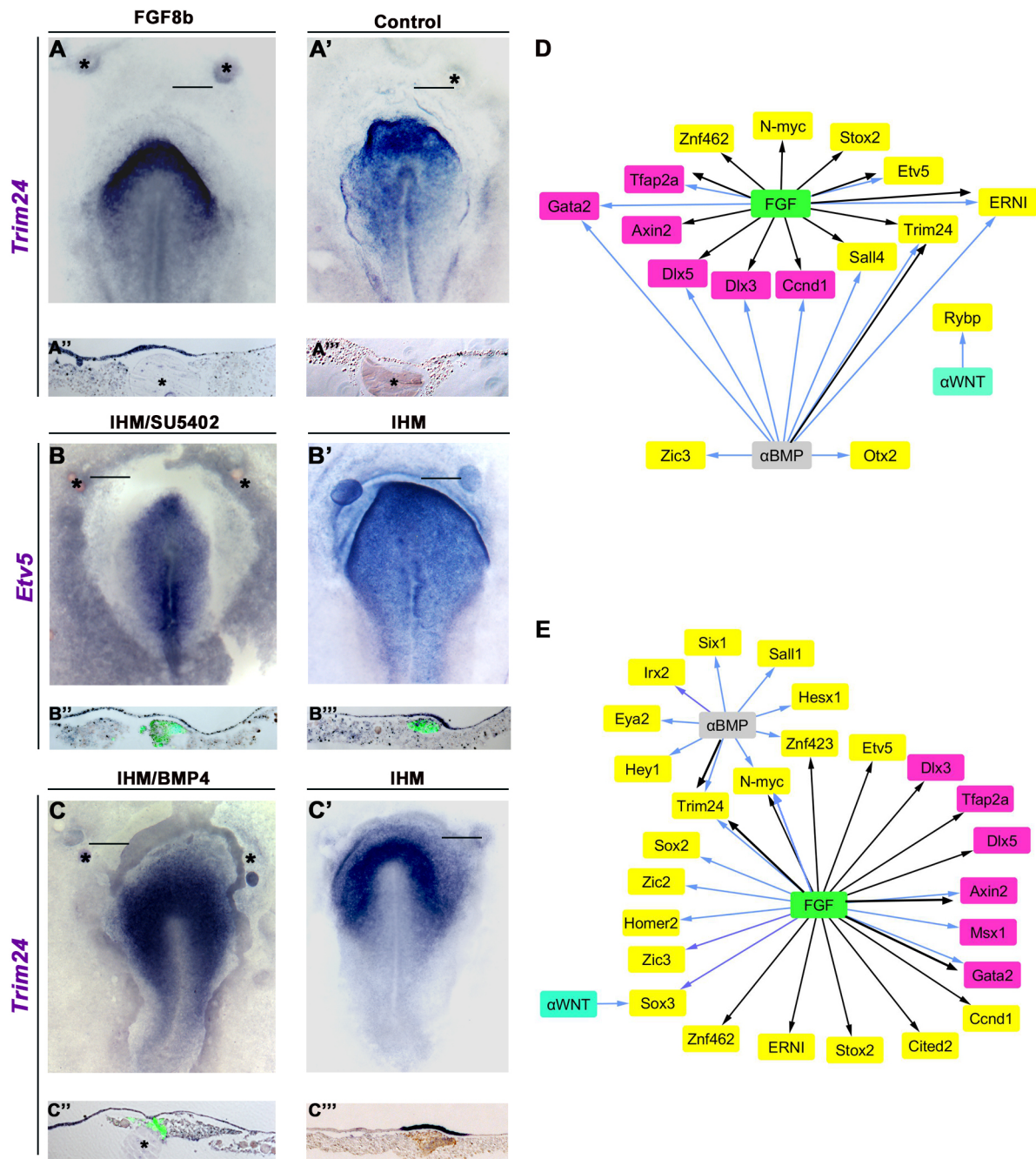


Fig. 5. Signals controlling mesoderm-response genes. (A-A'') Fgf8-coated (A,A''), but not control (A',A''), beads induce *Trim24* after 3 h (4/4). (B-B'') FGF signalling is required for the induction of *Etv5* by IHM. IHM (green in B',B'') induces *Etv5* in extraembryonic epiblast (B',B''; 5/6), and this is inhibited in the presence of SU5402-coated beads (B,B''; 4/4). (C-C'') *Trim24* is induced by IHM grafts (C', brown in C''; 3/3), and this induction is inhibited in the presence of BMP4-coated beads (C,C''; 4/4). Asterisks mark beads. Bars indicate level of sections shown beneath. (D,E) Different IHM (D) and pME (E) induced (yellow) or repressed (magenta) genes respond to modulation of FGF, BMP (α BMP, BMP antagonist) or Wnt (α WNT, Wnt antagonist). Black arrows indicate that signal is sufficient; blue arrows indicate that signal is required; see Table S3 and Fig. S7.

points, we exposed the extraembryonic epiblast to the Wnt antagonist IWR and compared gene expression with control untreated epiblast using NanoString after 3 and 6 h. Wnt inhibition results in the upregulation of several genes, including *Dlx5* and *Nfkb1* as well as the Six1 co-factor *Dach1* (Table S2). However, none of these is induced by mesoderm grafts, suggesting that antagonising Wnt signalling does not mimic the response to mesoderm signals.

To assess whether mesoderm-induced genes depend on Wnt antagonism, we combined tissue grafts with control DMSO-

coated beads or beads coated with the Gsk3 inhibitor BIO to activate canonical Wnt signalling. Only a few genes, which are normally induced after 6 h, are affected: *Rybp* is no longer induced by the IHM, while *Sox3* is no longer upregulated by the pME (Fig. 5D,E, Table S3, Fig. S7). Thus, modulation of Wnt signalling does not play a major role during the initial phase of PPR induction, but might be important for the decision between neural crest and placode progenitors (Litsiou et al., 2005; Villanueva et al., 2002).

Sonic hedgehog signalling is required for aPPR induction

The above experiments demonstrate that the axial mesendoderm is important to imbue sensory progenitor cells with anterior character. What is the nature of the anteriorising signal? Mesendoderm-derived somatostatin mediates PPR induction but does not impart regional identity (Lleras-Forero et al., 2013). Since sonic hedgehog (Shh) is prominently expressed in the pME (Dale et al., 1997), we tested whether it is required for aPPR formation. We grafted pME next to competent epiblast of HH4⁻ hosts with or without cyclopamine, an Shh inhibitor (Fig. 6A-C). After 16 h, the general PPR markers *Six1* (Fig. 6A) and *Eya2* (not shown) continue to be induced, whereas the anterior markers *Six3* and *Otx2* are reduced (Fig. 6B,C). Shh-coated beads are not sufficient to induce PPR or anterior markers in the extraembryonic epiblast. We therefore tested whether Shh can anteriorise the IHM-induced PPR. When IHM grafts are combined with Shh-coated beads, the anterior marker *Six3* is not induced (Fig. S11). Together, these data indicate that Shh alone is not sufficient to impart anterior identity or induce sensory progenitors, but plays a role in conferring rostral character to the PPR.

Network predictions support a transcriptional hierarchy during PPR induction

Our analysis suggests a gradual transition of epiblast cells towards placode progenitor fate, each step being defined by distinct transcriptional regulators and controlled by a combination of signals. Based on timecourse analysis we propose that at the top of the hierarchy a small set of transcription factors defines a 'primed ectoderm state' and controls the expression of a second tier of factors, which then directly regulate the *Six/Eya* cassette in the PPR. To test this model, we used an unbiased approach to investigate the topology of the PPR genetic network.

Using the NanoString datasets we inferred a regulatory network using the GENIE3 algorithm (Huynh-Thu et al., 2010). We created a network using all interactions with an importance measure of 0.02

or higher (Fig. S8A), which highlights important nodes (potential regulators) and weighted directional interactions. Visual inspection suggests the existence of three potential subclusters, which indeed emerge upon further dissection of the network using community clustering (Fig. S8, clusters G1-G3). *Six1*, *Six4*, *Eya2* and other PPR-specific genes cluster together with known *Six1* upstream regulators (Fig. S8C, cluster G2). In addition, G2 contains several early mesoderm-response genes including *Otx2*, *ERNI*, *Sall1*, *Sall4*, *Ccnd1*, *Trim24*, *Zic3* and the FGF mediators *Etv4* and *Etv5* (Fig. S8C). Thus, the network predictions correlate well with our proposed model.

To assess potential direct regulators of *Six1* and *Eya2*, we extracted their nearest neighbours from the network (Fig. S8A, Fig. S9). Using this information together with our induction timecourse and the spatial information from *in situ* hybridisation, we constructed a gene regulatory network in BioTapestry (Fig. 7A). Genes not induced by the mesoderm or not co-expressed with *Six1* and *Eya2* (e.g. *Hey2*, *Irx1*) were excluded from the BioTapestry network as potential direct interactors. Many first neighbours are shared by *Six1* and *Eya2*, including known upstream regulators such as *Gata3* and *Dlx6* (Kwon et al., 2010; Sato et al., 2010). In addition, the network predicts novel regulatory relationships between the *Six1/Eya2* complex and *Sall1*, *Gbx2*, *Hey1* and *Hesx1*. To corroborate these predictions, we analysed the transcription factor-binding sites of the only PPR enhancer so far identified, *Six1-14* (Sato et al., 2010), which directs *Six1* expression in the aPPR. This analysis confirms the presence of Gata, Dlx, Gbx and Sall motifs (Table S5), indicating that these factors may indeed directly regulate *Six1* (Fig. 7A, bottom, 12 h, purple genes).

By contrast, none of the early mesoderm-response genes defining the primed ectoderm state (*Etv5*, *Otx2*, *Zic3*, *N-myc*, *Trim24*; Fig. 7A, top, 3 h, blue genes) is predicted to interact with *Six1* and *Eya2* directly. We therefore analysed their first neighbours in the GENIE3 network as well as those of the predicted direct *Six1/Eya2* regulators (*Gata3*, *Dlx6*, *Gbx2*, *Foxi3*, *Sall1*; Fig. S9). This analysis

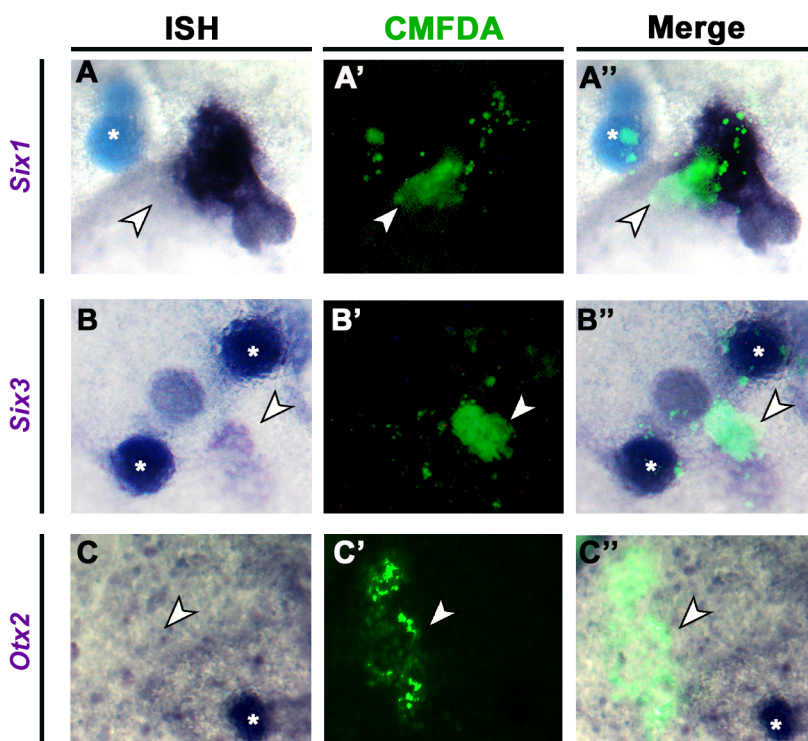


Fig. 6. Sonic hedgehog is required for aPPR induction. CMFDA-labelled (green fluorescence) pME was grafted together with cyclopamine-coated beads into the extraembryonic region of chick hosts. After 16 h, *Six1* is induced (A-A''; $n=7/9$ induced; no significant difference to controls, goodness of fit test). By contrast, *Six3* (B-B''; 4/13 induced compared with 8/11 in controls; $P=0.00166$, goodness of fit test; see Fig. 4E') and *Otx2* (C-C''; 0/4 induced compared with 3/3 in controls; $P=0.024$, two-tailed goodness of fit test; see Fig. S11) induction is lost. Arrowheads indicate grafts; asterisks indicate beads. ISH, *in situ* hybridisation.

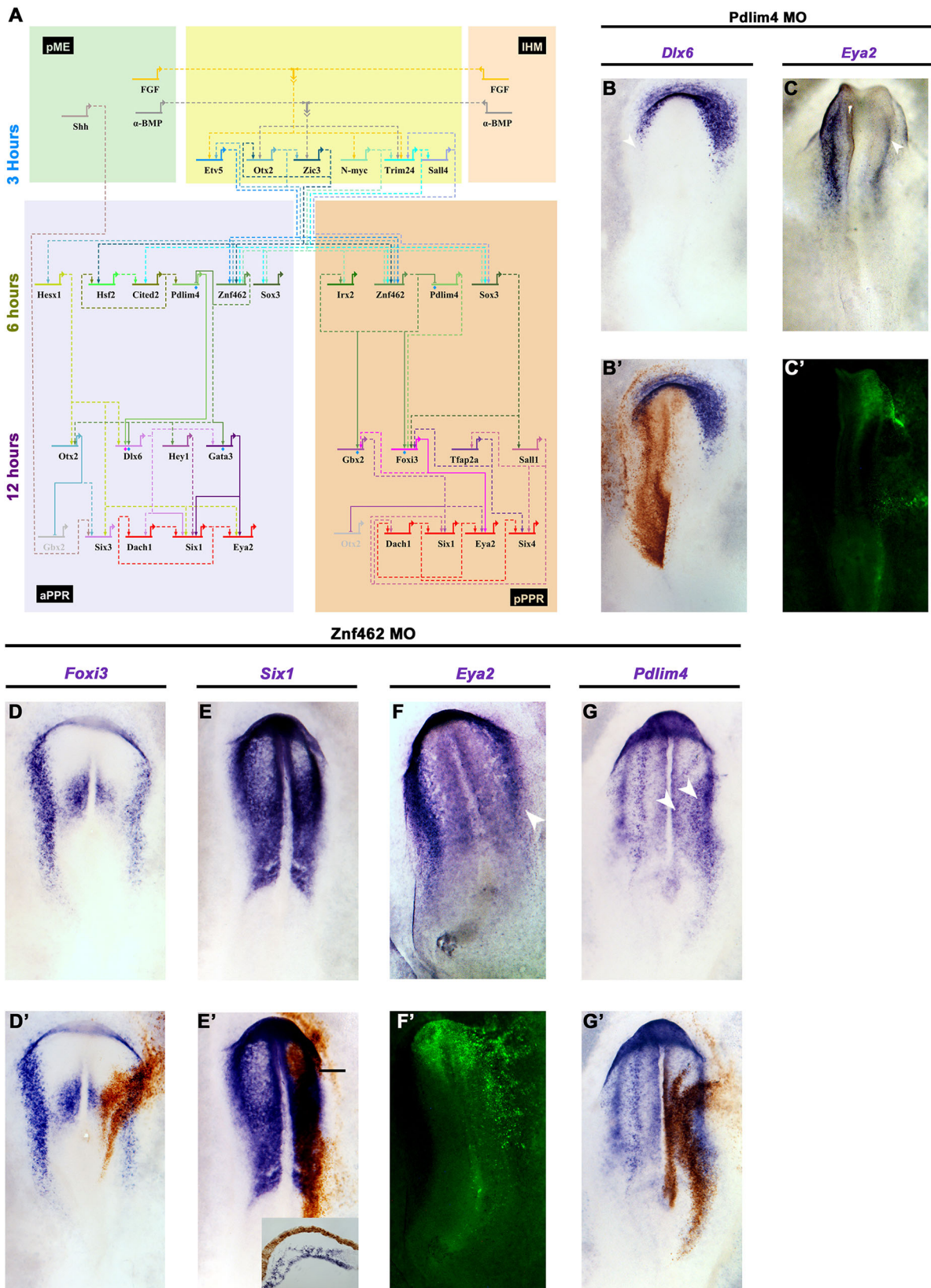


Fig. 7. A predicted gene network reveals new candidate *Six1* and *Eya2* regulators. (A) BioTapestry network that integrates time series of transcript induction and interactions predicted from GENIE3. Genes induced after 3 h, blue; 6 h, green; 12 h, purple; PPR genes, red. Dashed lines indicate predicted interactions; solid lines indicate known interactions and those experimentally verified in this study (blue diamonds for *Znf462*, pink diamonds for *Pdlim4*). (B-C') *Pdlim4* knockdown results in reduction of *Dlx6* (B,B'; 5/5) and *Eya2* (C,C'; 3/6). Arrowheads indicate changes in gene expression. (D-G') *Znf462* knockdown leads to reduction of *Foxi3* (D,D'; 4/5), *Six1* (E,E'; 4/5; inset, transverse section at level of bar) and *Eya2* (F,F'; 4/6; arrowhead indicates changes in *Eya2* expression), but to an expansion of *Pdlim4* into the neural plate (G arrowheads, G'; 3/5). Morpholinos are visualised by DAB staining (brown) or by fluorescence (green).

reveals a small set of intermediate transcripts downstream of the genes defining the primed ectoderm, among them *Hesx1*, *Znf462*, *Hsf2*, *Pdlim4*, *Irx2*, *Cited2* and *Sox3*, which are all induced 6 h after exposure to mesodermal signals (Fig. 7A, middle 6h, green genes). In turn, they are connected to the *Six1/Eya2* regulators suggesting that they provide the link between 3 and 12 h induced genes. We have termed the transcriptional state where 3 and 6 h induced genes are co-expressed the ‘PPR-primed’ state. In summary, our network inference approach supports a gradual transition of epiblast cells towards PPR fate and highlights the hierarchical nature of this process.

Confirming predicted interactions

To test whether these intermediate genes are regulators in the PPR gene network, we assessed the potential role of two transcription factors, *Znf462* and *Pdlim4*. We designed two morpholinos for each factor to knockdown their expression. Experimental or control morpholinos were electroporated at primitive streak stages, the aPPR and pPPR were dissected at HH6/7 and then analysed by NanoString. Whereas reduction of *Znf462* changes the expression of many transcripts when compared with controls (aPPR, 30; pPPR, 94), knockdown of *Pdlim4* only affects nine genes (Table S3). This analysis verifies five of six predicted links downstream of *Znf462* and one of two downstream of *Pdlim4*. To corroborate these findings we performed *in situ* hybridisation for selected genes after *Znf462* and *Pdlim4* knockdown. Whereas control morpholinos have no effect (Fig. S10), the reduction of either gene prevents *Six1* and *Eya2* expression (Fig. 7C,E,F). Furthermore, *Znf462* regulates *Foxi3* and *Pdlim4* (Fig. 7D,G): although *Znf462* is required for *Foxi3*, it seems to repress *Pdlim4*. Finally, *Pdlim4* knockdown leads to a reduction in *Dlx6* expression as predicted (Fig. 7B). Thus, these findings provide experimental support for the GENIE3-predicted gene regulatory network, and suggest that genes induced within 6 h after mesoderm grafting link early response genes and PPR competence factors.

DISCUSSION

Sensory placode progenitors are specified at head process stages in the ectoderm surrounding the anterior neural plate (Baker and Bronner-Fraser, 2001; Grocott et al., 2012; Schlosser, 2010; Streit, 2008). Initially, they are competent to give rise to all cranial placodes and share common features, but as individual placodes emerge their developmental potential becomes gradually restricted (Bailey et al., 2006; Bailey and Streit, 2006; Schlosser, 2010; Streit, 2008). Thus, induction of sensory progenitors is thought to be a common step that generates a non-regionalised placode territory, which is later subdivided along the rostrocaudal axis. Here, we provide evidence that PPR induction and regionalisation occur simultaneously, suggesting that a homogeneous PPR without anterior-posterior identity does not exist. Having identified new genes in the regulatory cascade that specifies sensory progenitors, we investigated their temporal hierarchy. This allowed us to dissect the events upstream of the PPR specifiers *Six1* and *Eya2* and revealed how signals from different tissues gradually induce placode precursors with distinct rostrocaudal identity.

A transcriptional hierarchy gradually specifies sensory placode progenitors with distinct rostrocaudal character

Three independent approaches – a PPR induction assay, gene expression during normal development, and network analysis – lead us to propose a new multistep model for PPR induction. Although we describe discrete steps as cells acquire PPR identity, it is likely

that in reality the transitions between them are fluid and represent a continuum. In the first step, signals from the IHM and the pME rapidly induce a small set of transcription factors including *ERNI*, *Etv5*, *N-myc*, *Otx2*, *Zic3* and *Trim24* (akin to ‘evocation’ in Waddington’s model). This set of transcription factors is largely identical irrespective of the signalling source, suggesting that both tissues initially promote a common transcriptional state. In the normal embryo these factors are expressed very early in development, with their domains encompassing the future neural plate, neural crest and placode territory, indicating that before acquiring PPR identity cells pass through a ‘primed ectoderm state’ that is common to progenitors for both the central and peripheral nervous systems. How this state is induced in the normal embryo remains to be elucidated; however, our results raise the possibility that FGF signalling might, at least in part, mediate this process. FGF8 largely mimics the response to mesodermal signals (see Fig. 5), is required for the early, but not the late, steps of PPR induction (Litsiou et al., 2005) and is known to initiate neural and neural crest formation (Delaune et al., 2005; Linker and Stern, 2004; Monsoro-Burq et al., 2003; Streit et al., 2000; Stuhlmiller and García-Castro, 2012; Wilson et al., 2000). It is thus possible that, like neural induction, PPR formation starts much earlier than previously thought and our induction assay rapidly recapitulates normal development.

The second phase of PPR induction is characterised by a set of new transcription factors (Fig. 7A, middle), the expression of which in the embryo starts after that of the genes defining the primed ectoderm, but is equally broad. Together, they characterise a transcriptional state that we have termed ‘PPR-primed’. Among these, the zinc-finger transcription factor *Znf462*, which is induced by both the IHM and axial mesendoderm, emerges as a potential integrator in the PPR gene network: several early response factors are predicted to provide input to *Znf462*, while *Znf462* itself regulates *Foxi3* and is predicted to control *Gbx2*, *Gata3* and *Dlx6*, which in turn control *Six1* and *Eya2*. In this phase, PPR regionalisation begins (akin to ‘individuation’ in Waddington’s model) and, under the influence of IHM and axial mesendoderm, anterior-posterior markers are upregulated. Finally, the bona fide PPR markers *Six1/4*, *Dach1* and *Eya2* are induced, as are PPR competence factors (Bhat et al., 2013; Kwon et al., 2010; Litsiou et al., 2005; Pieper et al., 2012), which might regulate *Six1* directly. Thus, as different populations of mesoderm emerge from the primitive streak, they transform primed ectoderm into PPR with regional character, without inducing definitive placode fates.

Although this hierarchical model emerges from an ectopic induction assay, epiblast cells go through the same transcriptional states as they gradually acquire PPR fate. Genes defining the primed ectoderm are present early, some before gastrulation (Grocott et al., 2012), followed by our newly identified PPR-primed factors. Where the latter overlap with PPR competence factors (*Gata*, *Dlx*, *Tfap2a*; Bhat et al., 2013; Kwon et al., 2010; Pieper et al., 2012), PPR specifiers are induced. Competence factors are initially expressed throughout the entire non-neural ectoderm at low levels (Bhat et al., 2013; Kwon et al., 2010; Pieper et al., 2012), but their expression increases in sensory progenitors as development proceeds. Our network analysis suggests that this enhanced expression might be controlled by the newly identified transcription factors, such as *Znf462*. Whether any of the competence or newly identified factors act as pioneer factors for the *Six/Eya* network remains to be elucidated.

This model for PPR induction somewhat resembles Waddington’s model for neural induction, which proposes that the organizer initially induces a tissue of generic neural character and then imparts regional

identity (Waddington and Needham, 1936). Although placode progenitor induction does not involve a bona fide organizer, but is instead mediated by different tissues, both initially induce a common transcriptional state (evocation) without rostrocaudal character, and subsequently impart regional bias (individuation). At this point PPR cells are specified as placode progenitors, but not yet committed to a specific placodal fate (reviewed by Bailey and Streit, 2006; Grocott et al., 2012; Streit, 2008). Together, our data suggest that during PPR induction cells initially pass through a primed ectoderm state, which is characterised by only a handful of genes, followed by a PPR-primed state defined by newly identified transcripts such as *Znf462*. Subsequently, mesoderm signals divert cells towards placode fate and simultaneously impart regional bias depending on their location along the rostrocaudal axis.

Tissues and signals in PPR initiation and regionalisation

Signals from the neural plate and the IHM have previously been implicated in placode progenitor induction (Ahrens and Schlosser, 2005; Litsiou et al., 2005). Here, we identify the pME as an inducer of aPPR fate. Assessing the behaviour of 126 transcripts simultaneously allows us to dissect the dynamic response to all three tissues and reveals that, although mesodermal signals initiate the full set of PPR transcripts in the induction assay, the neural plate alone is not sufficient to do so. It is likely that in normal development signals from the neural plate contribute to PPR induction. However, our findings suggest that they may play a less prominent role in initiating induction, but might be important to define regional character of the tissue induced.

FGF signalling mimics some of the earliest responses to signals from the mesoderm, although it is strictly required for only a few rapidly induced genes (Litsiou et al., 2005). Although BMP inhibition is sufficient to induce only two transcripts, namely *Trim24* and *Irx1*, it is required for the induction of many more. Together, both pathways account for the majority of IHM-induced genes and for a considerable number of transcripts initiated by the pME. In particular, our findings confirm that antagonising BMP signalling is crucial for the expression of the PPR specifiers *Six1* and *Eya* (Ahrens and Schlosser, 2005; Brugmann et al., 2004; Litsiou et al., 2005). Together, these results suggest that the cooperation of FGF activation and BMP antagonism is crucial during the early phase of PPR induction and that both contribute to establishing a PPR-primed state, but are not sufficient for PPR specification.

By contrast, Wnt signalling only appears to play a minor role during the first step of PPR specification. Wnt antagonism does not mimic any activity of the mesoderm and only very few genes depend on Wnt inhibition. This suggests that modulation of canonical Wnt signalling is important during the late phase of PPR induction, where it might mediate different processes. On the one hand, Wnt antagonists mediate the decision between neural crest and placode precursors at the border of the neural plate and protect PPR cells from Wnts emanating from surrounding tissues (Brugmann et al., 2004; Litsiou et al., 2005). On the other hand, Wnt antagonism might also be important to promote pPPR identity. Although active Wnt signalling is generally considered to posteriorise neural and non-neural ectoderm (Wilson and Houart, 2004), Wnt inhibition might be required to fine-tune pre-placodal fate. Finally, while studies in zebrafish have implicated Shh in the decision between adeno-hypophysis and lens character (Dutta et al., 2005), our results suggest an even earlier role at pre-placodal stages. Shh and somatostatin (Lleras-Forero et al., 2013) from the axial mesendoderm may cooperate to initiate *Pnoc* expression in aPPR cells, which in turn is required for the expression of aPPR transcription factors.

Conclusions

The combination of embryological timecourse experiments analysing more than 100 genes and network analysis reveals a new multistep model for PPR induction. We demonstrate that the acquisition of placode progenitor fate occurs gradually and that cells transit through different transcriptional states, each defined by distinct factors. The first step generates primed ectoderm cells, followed by a PPR-primed state, which might be shared with neural and neural crest induction. Subsequently, two different tissues, namely the pME and the IHM, gradually impart anterior and posterior PPR character, respectively. Thus, PPR induction and regionalisation occur simultaneously, suggesting that, once induced, PPR cells do not share the same transcriptional profile. However, since both tissues initially elicit an almost identical response and cells only diverge later, PPR induction is somewhat reminiscent of Waddington's evocation-individuation model for neural induction.

MATERIALS AND METHODS

Embryo techniques and *in situ* hybridisation

Fertile hens' (Winter Farm) and quails' (Potter Farm) eggs were incubated at 38°C to obtain embryos of appropriate Hamburger and Hamilton (HH) stages (Hamburger and Hamilton, 1951). Chick embryos at stage 3⁺/4⁻ were cultured according to New (1955) as modified by Stern and Ireland (1981). To isolate quail and chick tissues for grafting, embryos were collected in Tyrode's saline. IHM and pME were dissected from HH5/6 donors using fine steel needles and small amounts of dispase (1 mg/ml). Neural plate from different rostrocaudal levels was obtained from HH6 donors after removal of the endoderm and mesoderm. For fluorescently labelled grafts, tissue was collected as above and incubated for 45 min at 38°C in DMEM containing 10 μM 5-chloromethylfluorescein diacetate (CMFDA; Thermofisher). Tissues were kept on ice in Tyrode's saline until grafted into the inner margin of the extraembryonic area opaca of HH4⁻ hosts.

To inhibit Wnt or BMP signalling, whole embryos (HH4⁻) were cultured in modified New culture (Stern and Ireland, 1981) with albumen containing 30 μM IWR-1 (Sigma Aldrich) or 20 μM dorsomorphin (Tocris Bioscience), respectively. Heparin beads were coated in 50 μg/ml FGF8 (R&D Systems) in PBS containing 0.1% BSA on ice for 1 h, washed briefly in Tyrode's saline and grafted into the inner third of the area opaca of HH4⁻ stage chick embryos. Heparin beads were coated with 100 μg/ml Shh (R&D Systems) in PBS containing 0.1% BSA for 1 h on ice, washed in Tyrode's saline and grafted into the area opaca alone or in combination with IHM. To modulate different signalling pathways, AG1X2 beads (Sigma Aldrich) were coated with 1 μg/ml BMP4 (R&D Systems) in PBS containing 0.1% BSA for 1 h on ice, with DMSO (control), 25 μM SU5402 (Tocris Bioscience) or 2.5 μM BIO (Sigma Aldrich) in DMSO at room temperature for 2 h. Affi-Gel blue beads (Bio-Rad) were coated with 1 μM cyclopamine (Sigma Aldrich) in DMSO for 2 h. Beads were then washed in PBS and grafted together with IHM or pME. SU5402 is yellow, and beads retain their colour for at least 6 h after grafting, but have released all inhibitor after 15 h and appear white. To ensure that BMP4 beads are effective over the 6 h timecourse, we grafted beads next to the neural tube and assessed *Sox2* expression (Fig. S11). The effectiveness of IWR and dorsomorphin was assessed after growing embryos from HH5/6 in the presence of these drugs and assessing heart and somite formation (Fig. S11). As positive controls for Shh and cyclopamine we grafted beads next to the node at HH4 and assayed for *Pitx2* expression after 12 h of culture (Fig. S11).

For whole-mount *in situ* hybridisation, embryos were harvested in PBS, fixed in 4% paraformaldehyde and processed for hybridisation using DiG-labelled antisense mRNA as previously described (Streit and Stern, 2001). EST clones are listed in Table S6. To reveal quail tissue, QCPN antibody was used (1:5; AB 531886, Developmental Studies Hybridoma Bank).

Electroporation and morpholino knockdown experiments

Two independent fluorescein-labelled morpholinos were designed by GeneTools targeting *Pdlim4* (MO1, 3'-CGACACCACGTGCACCATACC-5'; MO2, 3'-CATCCACTTAAAGCGGCTCCGAGGC-5') and *Znf462*

(MO1, 3'-AGACACACAGATCCTTACCCTCTCT-5'; MO2, 3'-TGCAGC ACCTCCATGGTTCAAGGAT-5'). These are splice-blocking morpholinos; blastN does not reveal any other potential targets and their effectiveness was assessed by RT-PCR. Control morpholino was 3'-CCTCTTACCTCAGTT-ACAATTTATA-5'. For electroporation, each morpholino (1 µg/µl) was mixed with plasmid DNA (0.5 µg/µl) as a carrier and 0.01% Fast Green, and injected between the vitelline membrane and the epiblast. Primitive streak stage embryos (HH3^{+/4}-) were prepared for New culture; morpholinos were transferred into epiblast cells using four 5 mV pulses for 50 ms each, with an interval of 500 ms. Targeted cells are visualised by fluorescence or by immunostaining with anti-fluorescein antibodies. For NanoString analysis, electroporated aPPR and pPPR was dissected from HH6/7 after electroporation.

Microarray analysis

To identify differentially regulated transcripts, IHM was grafted into the area opaca of HH4⁻ chick hosts. After 12 h, mesoderm grafts were removed using small amounts of dispase and the underlying area opaca epiblast was collected together with control epiblast from the contralateral side. Tissue collection was repeated on three independent occasions, with 35-40 explants collected for each sample. Using 5 ng total RNA, labelled cell extracts were prepared and hybridised to Affymetrix Chick GeneChip according to Chambers and Lumsden (2008). Probe level values were derived from the raw data using the MAS5 algorithm (version 1.2, Affymetrix). Data were analysed using the GeneSpring package (version 7.3.1, Agilent Technologies). The suitability of the datasets for further analysis and the relationship between and within the biological replicates was determined using principal components analysis and hierarchical clustering. Differential expression between the conditions under investigation was determined by a stepwise process. Samples were first normalised to the 50th percentile across the whole expression dataset and then each gene was normalised to the median of its own expression across each cell type. Prior to statistical analysis, genes classed as not expressed (absent in biological replicates) or not varying in expression above a 2-fold threshold in any of the cell types were removed from the analysis. From the remaining set of genes, those with expression levels that differ significantly ($P \leq 0.05$) between each cell type were determined by one-way ANOVA. Microarray data generated in this study were deposited in Gene Expression Omnibus (GSE81023). Transcripts expressed in the aPPR and pPPR have been identified previously (Lleras-Forero et al., 2013; GEO accession GSE48116). For hierarchical clustering and heatmap generation, the R statistical packages Hclust and Heatmap.2 were used.

NanoString nCounter

Area opaca exposed to signals from different tissues and non-induced area opaca explants were collected at 3, 6 and 12 h after grafting. Three independent replicates of 5-10 explants (5000-10,000 cells) per condition were collected on ice in RNase-free PBS, which was then replaced by lysis buffer (Ambion) and the lysed tissues were quickly spun (13,000 rpm in an Eppendorf 5415R) before being snap frozen on dry ice. Samples were stored at -80°C until required. RNA lysates were hybridised at 65°C overnight, eluted according to the nCounter manual and counted by the nCounter digital analyser. Counts were normalised to the positive controls present in each hybridisation mix. Subsequently, the negative control probe values were used to create a background threshold level; transcripts with expression values below the threshold were removed from further analysis. Counts were then normalised to the total amount of mRNA counted in each sample. Differential expression of transcripts between different conditions was determined using an unpaired two-tailed Student's *t*-test comparing the average of three biological replicates ($P < 0.05$, >1.2-fold change).

GENIE3-inferred network analysis

NanoString data from tissue grafting timecourses and from signalling experiments that did not use DMSO (i.e. IHM/BMP4 for 3 and 6 h; FGF8 for 3 and 6 h) were used to generate a predicted gene regulatory network. The mean expression values for each gene under each condition were analysed by the GENIE3 algorithm (Huynh-Thu et al., 2010). In brief, this algorithm decomposes *n* genes into *n* different regression problems. For each regression problem, the expression profile of one gene (target gene) is

predicted from the expression profiles of all other genes (input genes) using tree-based ensemble methods. Within this, the importance of a single input gene in explaining the profile of the target gene is assessed and an importance measure is generated. This importance measure is then used to predict the regulatory links and their direction within the network. Following analysis, interactions above an importance measure of 0.02 were isolated, based on the strength of their importance measure, and the network was viewed using Cytoscape (www.cytoscape.org/). For subsequent analysis, genes of interest were highlighted with their first neighbours (putative regulators and targets) and small networks were created. To identify more closely related genes, community clustering was performed using the GLay plugin in Cytoscape (Su et al., 2010).

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Competing interests

The authors declare no competing or financial interests.

Author contributions

Conceptualization: M.H., A.S.; Formal analysis: M.H., A.S.; Investigation: M.H., R.S.P., M.T., N.A.D.C., M.A., T.G.; Writing - original draft: M.H., A.S.; Writing - review & editing: A.S.; Visualization: M.H., R.S.P., M.T.; Supervision: A.S.; Project administration: A.S.; Funding acquisition: A.S.

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Data availability

Microarray data generated in this study have been deposited in Gene Expression Omnibus under accession number GSE81023.

Supplementary information

Supplementary information available online at <http://dev.biologists.org/lookup/doi/10.1242/dev.147942.supplemental>

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