Endothelin receptor Aa regulates proliferation and differentiation of Erb-dependant pigment progenitors in zebrafish

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Skin pigment patterns are important, being under strong selection for multiple roles including camouflage and UV protection. Pigment cells underlying these patterns form from adult pigment stem cells (APSCs). In zebrafish, APSCs derive from embryonic neural crest cells, but sit dormant until activated to produce pigment cells during metamorphosis. The APSCs are set-aside in an ErbB signaling dependent manner, but the mechanism maintaining quiescence until metamorphosis remains unknown. Mutants for a pigment pattern gene, parade, exhibit ectopic pigment cells localised to the ventral trunk, but also supernumerary cells restricted to the Ventral Stripe. Contrary to expectations, these melanocytes and iridophores are discrete cells, but closely apposed. We show that parade encodes Endothelin receptor Aa, expressed in the blood vessels, most prominently in the medial blood vessels, consistent with the ventral trunk phenotype. We provide evidence that neuronal fates are not affected in parade mutants, arguing against transdifferentiation of sympathetic neurons to pigment cells. We show that inhibition of BMP signaling prevents specification of sympathetic neurons, indicating conservation of this molecular mechanism with chick and mouse. However, inhibition of sympathetic neuron differentiation does not enhance the parade phenotype. Instead, we pinpoint ventral trunk-restricted proliferation of neural crest cells as an early feature of the parade phenotype. Importantly, using a chemical genetic screen for rescue of the ectopic pigment cell phenotype of parade mutants (whilst leaving the embryonic pattern untouched), we identify ErbB inhibitors as a key hit. The time-window of sensitivity to these inhibitors mirrors precisely the window defined previously as crucial for the setting aside of APSCs in the embryo, strongly implicating adult pigment stem cells as the source of the ectopic pigment cells. We propose that a novel population of APSCs exists in association with medial blood vessels, and that their quiescence is dependent upon Endothelin-dependent factors expressed by the blood vessels.
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21st December 2018

Dear Editor,

Please find attached a revised version of the manuscript "Endothelin receptor Aa regulates proliferation and differentiation of Erb-dependant pigment progenitors in zebrafish", which we ask you to consider for publication in PLoS Genetics.

We have addressed all the issues raised by the reviewers, and we do hope that you will consider it worthy of publication in PLoS Genetics.

Sincerely

Robert

Professor Robert Kelsh
Response to reviewers PGENETICS-D-18-00944

We thank the referees for their careful assessment of Camargo-Sosa et al, and for their helpful suggestions which we have incorporated into our revised document. Our point-by-point response follows:

Reviewer #1: PGENETICS-D-18-00944

...additional proof of these existence of [APSCs at blood vessels] in wildtype animals or abbreviate their interpretations...... Can lineage mapping or markers be used to point out the APSCs in wildtype animals and their ectopic progeny in pde animals? Previous studies also suggest that APSCs reside with the DRG rather than sympathetic ganglia (Dooley et al. 2013).

The referee correctly understands the novel hypothesis to which our data points, and we agree that direct proof of the presence of APSCs in this location in WTs will be the next step. However, our initial studies with inducible Cre induction of neural crest clones suggests that we can identify labelled cells in the vicinity of those trunk blood vessels in only c 1% of induced embryos, so that the direct lineage test will require a considerable investment. Furthermore, the lead author’s PhD studentship conditions mean that she absolutely had to submit her thesis by the end of Sept this year, so that it has not been realistic for her to perform the lineage study, and as the referee suggests, we feel that providing this proof is out of the scope of this paper. However, following the suggestion also of Referee 2 we have shown that the transgenic marker used by Dooley et al (2013) does reveal cells in the vicinity of the dorsal aorta, consistent with our model; we have added an extra figure (S6 Fig) and a description in the Results of this new data.

Additionally, although the authors suggest that these ectopic melanocytes are not a product of disrupted migration or transdifferentiation, they did not address whether they could be a result of simple overproliferation of ventral chromatophores, some of which might home to the blood vessels. Certainly, their observations that there are more Sox10-cre cells that express PH3 (Fig 5F) in parade mutants could support a hyperproliferation mechanism.

We have considered this model, and rejected it because 1) pH3+ melanocytes are very unusual (Only 1 in 20 in WTs, and none in 20 pde mutants); and 2) it does not explain why the ErbB inhibition should rescue the phenotype. We have now added a note in the Discussion commenting on this.

Minor comments:

1. It may be that not all the readership will be zebrafish researchers, and so there are a couple place in the main text where it would be helpful to explain a couple of details that are model-specific. Specifically, can you just briefly explain: line 169, what are the characteristics of iridophores and melanophores; line 203, what is the WIK background; line 220, what was the timing of the morpholino injections

Thank you for suggesting these clarifications, which we have now made.

2. line 175: You may want to mention that the presence of the double membrane is suggestive of individual cells rather than a single cell with dual characteristics.

Thank you for suggesting this clarification, which we have now made.
3. line 247: The sentence about splicing confused me at first until I realized that the sequencing that was performed was on cDNA, and the image in 3B is of predicted mRNA transcripts. It may be helpful to remind readers at the start of the paragraph what you are assessing.

Now added.

4. line 308: Does the pde mutant show overproliferation of all neural crest within the ventral region of the medial pathway? 55% increase in proliferating cells seems like a lot in comparison to the handfuls of ectopic chromatophores that are produced in the pde mutants.

This is an interesting point, but on further consideration perhaps not as surprising as it initially seems. As shown in Fig 4, we see no evidence of increases in peripheral neuron numbers, but note that we see not only ectopic pigment cells, but also supernumerary ones in the Ventral Stripe. We also note the very low baseline (WT) level of proliferation, and suggest that the process of regulation (e.g. of glial cell numbers) may also contribute to the balance of proliferation and extra differentiated cells that we observe.

5. line 323: is this the same line mentioned in line 310? Just a different way to assess the data? This could be better explained.

Yes. We have now modified the sentence to clarify this point.

6. All graphs: The red is hard to distinguish in grayscale or B/W print. Especially for those graphs where you do not directly label the red data points (e.g. 5G) it would be easier to see the pde mutant measurements if the data points were another shape.

We have now modified the graphs as suggested in order to improve their clarity.

7. Fig 2D: Can you better explain the differences of what you are measuring here? What are each of the dots representing?

Each dot represents the number of melanocytes, iridophores or the sum of both in the ectopic position in the ventral medial pathway (VMP) from an individual pde mutant larva at 4 dpf.

8. Fig 4: The order of the subfigures should match the order of the description in the text. Perhaps you can move the DRG data to be presented first so that you can keep the immunolabeling of the sympathetic neurons with the BMP inhibitor data.

As suggested, we have exchanged panels G-J and A-F in order to reflect the order in which this data is discussed in the text.

9. Fig 5 legend, what is the...LYN-EGFP?

This label indicates that the GFP is fused with a fragment of LYN tyrosine kinase to target it to the plasma membrane. This is now indicated in the figure legend text.

Reviewer #2:
1. Evidence that ectopic pigment cells are derived from APSC located in previously uncharacterized niche within ventral peripheral nervous system needs to be strengthened.

The authors show that inhibition of ErbB signaling rescues the parade phenotype suggesting that ectopic pigment cells are derived from Erbb3-dependent adult pigment cell progenitors. This result leads the authors to conclude that a previously unidentified pigment cell progenitor population resides in peripheral nerves (likely associated with sympathetic ganglia) located near the dorsal aorta and that these cells precociously differentiate when ednraa signaling is lost. While previous studies (Dooley et al., 2013; Singh et al., 2016) also suggest the possibility of ventrally localized adult pigment cell progenitors the conclusions drawn in this paper require more direct evidence of a distinct niche closely associated with the dorsal aorta. Further support could be provided in several ways: 1. Identification of mitfa+ cells in association with sympathetic ganglia or along peripheral nerves near the dorsal aorta in WT fish (similar to what has been shown by (Budi et al., 2011; Dooley et al., 2013), but specifically focused on region near dorsal aorta); 2. Activation of mitfa expression (presence of mitfa+ cells) within this niche in response to larval pattern ablation and migration of mitfa+ cells from ventral peripheral nerves to surface observed during time lapse imaging in WT; 3. Similar type of time-lapse analysis showing that cells from this location migrate along ventral nerves and contribute to the adult pattern.

Whilst we agree that more direct evidence of stem cells in the ventral trunk would be desirable, this largely required the use of tools that we did not have available and so was outside the scope of the present study. Meanwhile, we have established the Tg(mitfa:GFP) line used by Dooley et al as the only described marker of the APSC. In line with the referee’s suggestion, we have now assessed the expression of this transgene in WT larvae, focusing on the vicinity of the dorsal aorta in the trunk. At 8 dpf we found an average of 2.3 mitfa:gfp+ cells per segment. We have added this observation to the end of the Results and included a further figure (S6 Figure).

In addition, it is somewhat curious that no adult pattern defect was observed in parade mutants and thus some explanation of potential reasons why would be appropriate in the discussion (ex. stem cells are not exhausted during precocious differentiation in parade, or DRG stem cells are able to compensate for loss).

We were also somewhat surprised by this, but consider both the explanations proffered by the referee, as well as the suggestion that these APSCs may contribute primarily to non-integumentary (internal) pigment cells, as strong hypotheses to explain the observations. We have now added a comment to this effect in the Discussion on p. 12:

‘It is somewhat surprising that homozygous ednraa mutants show no disruption of the adult pigment pattern, but we propose several hypotheses, none mutually exclusive, to explain this observation: 1) although as we have shown APSCs are precociously activated in ednraa mutants, APSC renewal may be unaffected so that the stem cells are not exhausted; 2) ventral APSCs may primarily populate internal, rather than skin, pigment cell populations; 3) APSCs localised in DRGs may compensate for the effects on the more ventral APSCs.’

2. Relationship between mutations in parade alleles and ednraa structure/function is not clear.
From the schematic shown in Figure 3, it is very difficult to assess the severity of the three parade alleles. In Figure 3, a schematic showing the various protein domains (extracellular domains, intracellular domains, transmembrane domains) and their how the parade mutations alter the structure of the ednraa protein would be an improvement over the current diagram (Examples: Lang et al., 2009; Lopes et al., 2008). An attempt at explaining the consequences of the parade mutations on ednraa structure is provided in the discussion, but it is very confusing. Similarly, Figure S5, referenced in the discussion, is confusing and the colors and labels in the diagram do not match the descriptions provided in the figure legend. The information contained in lines 406-412 of the discussion be more appropriate for the results section with the cloning of parade, but there it still needs to be more clearly explained. From the main text and Figure 2 and Figure S5, it is not clear to me that any of these alleles should be considered a null and assessment of mutant phenotypes requires an understanding of how the mutations are likely to alter ednraa protein function.

We are grateful for the opportunity to improve the explanation of the molecular effects of the alleles, and the justification for why all are likely functional null alleles. As suggested, we have revised Fig. 3 to now include an extra panel (C) showing schematically the consequences of the splicing and point mutations at the protein level, which now complements the sequences shown in Fig. S5. We have moved the explanation in lines 406-412 from the Discussion to the Results (P. 7), and have modified it in order to clarify our explanation. We apologise for the errors in the Fig legend of Figure S5; this has now been edited to correct these. We believe that the fact that all mutants truncate the protein within the series of transmembrane domains that are at the core of this G-protein coupled receptor, combined with the observation that all three mutant alleles have quantitatively similar phenotypes, indicates that all are likely nulls. We have also included a statement that definitive testing of this idea will require creation of a truncation allele upstream of all known protein domains.

3. Lack of experiment demonstrating that ednraa expression in the dorsal aorta rescue ectopic pigment cell phenotype

One of the most exciting implications of this work is the suggestion that the blood vessels provide ednraa-dependent signals required to maintain nearby adult pigment stem cells in a quiescent state. When these unknown signals are absent, the stem cells differentiate resulting in ectopic pigment cells. However, the authors fail to provide evidence that ednraa expression in the dorsal aorta or other nearby blood vessels can rescue parade phenotype which seems crucial for establishment of a direct link between ednraa expression in blood vessels and stem cell maintenance. There may be technical reasons for not including these experiments but blastula stage transplants of flia:EGFP into parade mutants would be one way to perform this critical experiment.

We agree that a test of cell autonomy to the blood vessels would be interesting, but the variability of the pigment cell phenotype in both numbers and location of the ectopic pigment cells, the inevitable mosaicism of the blood vessels in transplantation experiments, and the expectation that signals from the blood vessels are likely to be secreted, means that this experiment might well be uninformative. As explained above, Karen has been under pressure to write up her thesis, and there is no one else available, we have reluctantly decided that we cannot attempt this risky experiment within the timeframe required. We trust that the editor will understand the cost-benefit calculation we have been forced to make here.
I also have a number of more minor concerns listed below:

1. Scale bars are missing in all Figures.

We apologise for this oversight, and have now added them to each figure.

2. Figure 3 I-J: In the version of the manuscript I received, filia:EGFP was not visible in panels I and J of Figure 3. Additionally, definitions of abbreviations DA, PVC and SC are mentioned in the figure legend but these labels do not appear in any of the figure panels, presumably because they are part of the missing 3I and 3J GFP images.

We are not quite sure what happened to the reviewer’s copy, but have checked that the version uploaded this time is complete and includes panels I and J, with the labels shown.

3. Figure 3E-H: It is difficult to evaluate ednraa expression in whole embryos. ISH images of the same region as shown in 3C and D and 3 I and J would better illustrate the spatial relationship between ectopic pigment cells and ednraa expression. It is worth showing whole embryos ISH but these images would be better in a supplemental figure that could include images of some of the other timepoints mentioned in the text but not shown.

We have now modified the panels in Fig. 3 in accordance with the reviewer’s helpful suggestions. We considered showing the whole-mounts, but in the end decided that these added little to the story of this manuscript, and are anyway available in the original description by Nair et al (2007) and so have not added a further Supplementary figure.

4. Figure 3 B: No iridophores are visible in image of ednraa morpholino injected fish. Since Figure 2 shows that parade mutants have more ectopic iridophores and supernumerary iridophores in the ventral stripe, I would expect to see iridophores in this image. If ectopic iridophores were not present it would be worth explaining in the text. If they are present, I would suggest using a different image that is illuminated to better show iridophores.

We have now labelled the figure to show both the ectopic melanocytes and iridophores in the ednraa MO injected fish. We have modified the fig. legend to make this clear.

5. Lines 227-230: dose-dependent response of translation-blocking morpholino. A figure in support of this statement would be most helpful here (supplemental?).

We apologise for a careless turn of phrase. The translation morpholino was not tested at multiple doses, but at 25 ng dose showed non-specific deformations (body curvature), so was not explored more extensively. We have revised the text to make this clear, and added reference to a PhD thesis which illustrates the result.

6. Line 267-268 is confusing. Should this start with “Ectopic pigment cells”?

Yes, we have now made this change.

7. Line 308-310: It is not clear why we would expect the timing of ectopic cells to be correlated with increased proliferation. Need better motivation for examination of proliferation.
The time of first detection of the ectopic pigment cells (no obvious increase compared to WT siblings at 24 and 30 hpf, visible increase at 35 hpf) indicates a latest time when proliferation might be detected. Furthermore, given the rapid differentiation of pigment cells from the NC, and the early nature of the pigment cell markers assessed, we predicted that the 30-35 hpf time window was a good place to look for possible proliferative changes. We have now edited this paragraph to make this clearer.

8. Lines 310-313: When was GFP expression induced in reporter? At 32 hpf or at an earlier timepoint? At 30hpf. We have now edited the sentence to clarify this point.

9. Figure 4 A-D; G-H: In addition to Hu antibody staining in Green these images appear to have unidentified red cells, giving the false impression that the reader is looking for co-localization. If the red is an artifact of staining procedure or autofluorescence of other cells, please explain in the figure legend.

We apologise for this oversight. The referee is referring to autofluorescence from fixed red blood cells. This is now mentioned in the figure legend for clarity.

10. S3 Figure: Inclusion of DIC images for C-F would be helpful. Currently is it difficult to tell which GFP+ cell are ectopic pigment cells. The association between location of the ectopic pigment cells and sympathetic ganglia is important for supporting the hypothesis that this is an APSC niche, so this image should be very clear. DIC would be good, pigment cell reporter lines would be even better.

DIC was included in the previous figure, but we acknowledge that it was perhaps too dark to be helpful. We have now revised the figure to make the DIC more obvious, including adding separate panels for the DIC, and have added further labeling to clarify key features.

11. In the introduction, line 70 and 91-94 need references.

Thank you for noting this important omission. We have now added references for these points.

12. At several points in the text (ex. line 134; line 252; line 420) authors state that ednraa is expressed only in blood vessels. While this appears to be the case in the trunk, ednraa is also expressed by cranial neural crest cells. Statements should be clarified to indicate that authors are only referring to expression in region of interest (site of ectopic pigment cells).

The referee is exactly right, and we have now added explicit reference to this at each of these mentions.

13. In Figure 1: Labels for DA and PCV are to the left of these structures and within somites. A line or narrow arrow pointing from the label to structures would be helpful.

We have edited the figure as suggested.

14. S1 Figure: It would be helpful to include a bright field image of the yolk sac stripe to demonstrate the association between melanophores and iridophores in this stripe and to indicate location of TEM
We have added a panel to the figure as suggested.
Endothelin receptor Aa regulates proliferation and differentiation of Erb-dependant pigment progenitors in zebrafish

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Abstract

Skin pigment patterns are important, being under strong selection for multiple roles including camouflage and UV protection. Pigment cells underlying these patterns form from adult pigment stem cells (APSCs). In zebrafish, APSCs derive from embryonic neural crest cells, but sit dormant until activated to produce pigment cells during metamorphosis. The APSCs are set-aside in an ErbB signaling dependent manner, but the mechanism maintaining quiescence until metamorphosis remains unknown. Mutants for a pigment pattern gene, parade, exhibit ectopic pigment cells localised to the ventral trunk, but also supernumerary cells restricted to the Ventral Stripe. Contrary to expectations, these melanocytes and iridophores are discrete cells, but closely apposed. We show that parade encodes Endothelin receptor Aa, expressed in the blood vessels, most prominently in the medial blood vessels, consistent with the ventral trunk phenotype. We provide evidence that neuronal fates are not affected in parade mutants, arguing against transdifferentiation of sympathetic neurons to pigment cells. We show that inhibition of BMP signaling prevents specification of sympathetic neurons, indicating conservation of this molecular mechanism with chick and mouse. However, inhibition of sympathetic neuron differentiation does not enhance the parade phenotype. Instead, we pinpoint ventral trunk-restricted proliferation of neural crest cells as an early feature of the parade phenotype. Importantly, using a chemical genetic screen for rescue of the ectopic pigment cell phenotype of parade mutants (whilst leaving the embryonic pattern untouched), we identify ErbB inhibitors as a key hit. The time-window of sensitivity to these inhibitors mirrors precisely the window defined previously as crucial for the setting aside of APSCs in the embryo, strongly implicating adult pigment stem cells as the source of the ectopic pigment cells. We propose that a novel population of APSCs exists in association with medial blood vessels, and that their quiescence is dependent upon Endothelin-dependent factors expressed by the blood vessels.
**Author Summary**

Pigment patterns are crucial for the many aspects of animal biology, for example, providing camouflage, enabling mate selection and protecting against UV irradiation. These patterns are generated by one or more pigment cell-types, localised in the skin, but derived from specialised stem cells (adult pigment stem cells, APSCs). In mammals, such as humans, but also in birds and fish, these APSCs derive from a transient population of multipotent progenitor cells, the neural crest. Formation of the adult pigment pattern is perhaps best studied in the zebrafish, where the adult pigment pattern is formed during a metamorphosis beginning around 21 days of development. The APSCs are set-aside in the embryo around 1 day of development, but then remain inactive until that metamorphosis, when they become activated to produce the adult pigment cells. We know something of how the cells are set-aside, but what signals maintain them in an inactive state is a mystery. Here we study a zebrafish mutant, called *parade*, which shows ectopic pigment cells in the embryo. We clone the *parade* gene, identifying it as *ednraa* encoding a component of a cell-cell communication process, which is expressed in blood vessels. By characterising the changes in the neural crest and in the pigment cells formed, and by combining this with an innovative assay identifying drugs that prevent the ectopic cells from forming, we deduce that the ectopic cells in the larva derive from precocious activation of APSCs to form pigment cells. We propose that a novel population of APSCs are associated with the blood vessels, that these are held in a quiescent state by signals coming from these vessels, and that these signals depend upon *ednraa*. Together this opens up an exciting opportunity to identify the signals maintaining APSC quiescence in zebrafish.

**Introduction**

Pattern formation is a crucial aspect of development since it creates the functional arrangements of cell-types that allow an organism to thrive. Pigment pattern formation—the generation of correctly distributed pigments or pigmented cells within the skin or elsewhere in the body—is a case in point, with pigmentation crucial for diverse aspects of an animal’s ecology, including avoidance of predators, kin recognition, mate selection, thermal regulation and UV protection [Irion, 2016 #15538]. In vertebrates, all pigment cells except those of the pigmented retinal epithelium, are derived from a transient embryonic tissue called the neural crest. Neural crest cells are multipotent, generating numerous types of neurons, glia, pigment cells and other derivatives. They are also highly migratory, moving from their origin in the dorsal neural tube to occupy diverse sites throughout the embryo. Thus, correct positioning of the different cell-types is a crucial aspect of their development.

Pigment cells in mammals consist only of melanocytes, making (and secreting) black eumelanin or yellow pheomelanin granules. In fish, amphibians and reptiles, pigment cells are much more diverse [1], allowing the generation of the varied and often beautiful pigment
patterns these groups display. The zebrafish *Danio rerio* has rapidly become a paradigmatic example for the genetic and cellular study of pigment pattern formation [2-6]. Zebrafish pigment patterns consist of three pigment cells, black melanocytes making melanin, yellow xanthophores making pteridines and carotenoids, and iridescent iridophores containing reflecting platelets [1].

Zebrafish, in common with most fish, develop two distinct pigment patterns, an early larval pigment pattern generated in the embryo by direct development of pigment cells from neural crest cells, and an adult pattern formed during metamorphosis, mostly through the de novo differentiation of pigment cells from adult pigment stem cells (APSCs, also formerly known as melanocyte stem cells; [7-9]). The adult pigment pattern consists of prominent stripes consisting of melanocytes and associated blue iridophores, alternating with pale stripes (interstripes) consisting of dense silver iridophores and xanthophores. Pigment pattern formation in adults is partially well characterised, with many genes identified that regulate the production of different pigment cell-types from the APSCs or which control their cellular interactions to create the bold horizontal stripe pattern {Irion, 2016 #15538; Kelsh, 2009 #12743; Parichy, 2015 #15005; Parichy, 2009 #9549}. A key aspect of adult pigment pattern formation that is less well-understood is the generation of the APSCs from neural crest cells. Remarkably, elegant experimental studies from multiple laboratories have established that these are set-aside from the neural crest in a narrow time-window (9-48 hours post-fertilisation (hpf); [10]), with at least some occupying a niche within the dorsal root ganglia (DRGs; [9, 11]) of the peripheral nervous system. They remain quiescent until metamorphosis begins around 20 days post-fertilisation (dpf), when they become activated. The mechanisms controlling their quiescence and their activation are largely unknown.

Pigment pattern formation in embryos is also poorly understood[3]. The embryonic pigment pattern consists principally of four longitudinal stripes of melanocytes, with iridophores arranged in a characteristic association with the melanocytes in three of these (Dorsal, Ventral and Yolk Sac Stripes), whereas the Lateral Stripe consists only of melanocytes; xanthophores then occupy the space under the epidermis between these stripes. Whereas in the adult the stripes are in the dermis, in the embryo they are associated with other structures, including the CNS, horizontal myoseptum of the body muscle blocks, the internal organs and the ventral most yolk sac. One study has investigated the detailed mechanism driving the association of melanocytes with the horizontal myoseptum [12]. In general, stripes form through migration of pigment cell precursors down migration pathways used by neural crest. These neural crest migration pathways are known as the dorsal (or dorsolateral) migration pathway, consisting of cells migrating under the epidermis and over the outer face of the somites/developing muscle blocks, and the medial migration pathway, running between the neural tube and notochord and the medial face of the somites/developing muscle blocks [13]. Pigment cell precursors of different fates use distinct migration pathways. Thus, xanthoblasts only use the lateral migration pathway, iridoblasts use only the medial migration pathway, whereas melanoblasts use both [14-17]. Note that migrating pigment cell precursors often show early signs of pigmentation i.e. they are differentiating as they migrate. During the
migration phase, early differentiating melanocytes and iridophores can be found in the ventral
trunk on the medial migration pathway, but these have disappeared by 72 hpf as those cells
migrate into the Ventral and Yolk Sac Stripes[16].
As in adult pigment pattern formation, mutants affecting embryonic pigment pattern offer an
exciting entry-point to the study of the mechanisms controlling pigment pattern formation.
In a large-scale ENU mutagenesis screen performed in 1996, we identified two zebrafish
mutant alleles, \( pde^{t262} \) and \( pde^{v212} \), that defined the \( \text{parade} (pde) \) gene [18]. These mutants
showed ectopic melanophores and iridophores in a well-defined region of the ventral side of
the posterior trunk (Fig 1B and E), in addition to a stripe pattern similar to the wild type (WT)
pigment phenotype (Fig 1A and C). The striking coincidence of melanin and reflecting
platelet distribution lead us to initially propose that the \( pde \) mutant phenotype results from
differentiation of pigment cells of mixed fate i.e. with both melanin granules and reflecting
platelets within the same cell [18]. Here we perform a comprehensive analysis of the \( pde \)
mutant phenotype. We show that the \( pde \) locus encodes a zebrafish Endothelin receptor A,
\( \text{Ednraa} \), which in the trunk is expressed only in the developing blood vessels, although in the
head it is also expressed in cranial neural crest cells and is associated with craniofacial
patterning [21]. Using chemical genetics, coupled with analysis of cell fate studies, we
 propose that APSCs occupy a niche associated with the medial blood vessels of the trunk,
likely to be a second PNS niche within the sympathetic ganglion chain, and that these become
precociously activated in \( pde/\text{ednraa} \) mutants. Thus we hypothesise that key components of
that blood-vessel niche are \( \text{Ednraa} \)-dependent factors that promote APSC quiescence.

Results

Early larval \( pde \) mutants have supernumerary iridophores and ectopic melanocytes and
iridophores

To test our initial working model, that the ectopic cells are cells of mixed
melanocyte/iridophore fate, we began by asking exactly where the ectopic cells were located,
and which pigment cell-types were involved. Whilst WT siblings at 5 dpf (Fig 1D) show no
pigment cells in the ventral trunk between the notochord and the Ventral Stripe, in \( pde^{t262} \)
mutants (from now on referred to simply as \( pde \) unless specified otherwise) ectopic
chromatophores are located in a medial position, directly beneath the dorsal aorta (DA) and
above the posterior cardinal vein (PCV; Fig 1F). Thus, their location corresponds precisely
to the ventral region of the medial neural crest migration pathway.
In zebrafish, pigment cell migration (both as unpigmented progenitors and as pigmenting
cells) is pathway specific, with cells of the melanocyte and iridophore, but not xanthophore,
lineages using the medial pathway [3]. Direct observation clearly shows cells with the
pigmentation characteristics of both melanocytes and iridophores in this region of *pde* mutants, but detection of xanthophores by their pigmentation is difficult at these stages. Thus, to assess the presence of ectopic xanthophores, we used whole-mount in situ hybridisation (WISH) using fate-specific markers. Detection of the melanocyte marker *dopachrome tautomerase* (*dct*; Fig 1J), the iridoblast marker *endothelin receptor ba* (*ednrba*; Fig 1K) and the xanthophore marker *GTP cyclohydrolase I* (*gch*; Fig 1L) at 72 hpf, provided clear confirmation that *pde* mutants display ectopic melanocytes and iridophores, but importantly revealed the complete absence of ectopic xanthophores in the ventral trunk (Fig 1L).

Initial observations of *pde* mutants indicated the intriguing possibility that some of the ectopic chromatophores displayed mixed characteristics of both iridophores and melanophores, i.e. both reflecting platelets and melanin-containing melanosomes (Fig 1E and F; [18]). This suggested that the phenotype might result, at least in part, from a failure of the normal mutual repression of alternative pigment cell fates. To test this, we used transmission electron microscopy to assess the structure of the ectopic cells. Unexpectedly, we consistently saw that melanosomes, the melanin synthesising organelles of melanocytes, and the reflecting platelets that contain the reflective crystals in iridophores, formed separate clusters, and were never intermingled (Fig 2A-2C). Furthermore, these clusters were consistently separated from each other by double membranes (Fig 2A-2C), similar to the appearance of melanocytes and iridophores in the WT Yolk Sac Stripe (S1 Fig), and consistent with them being separate cells. We conclude that the ectopic pigment cells are not of mixed fate, but instead are tightly associated individual melanocytes and iridophores; we note that this arrangement is characteristic of iridophores in their normal locations, where in each of the Dorsal, Ventral and Yolk Sac Stripes iridophores are tightly associated with melanocytes.

We hypothesised that the ectopic cells might reflect a defect in pigment cell migration through the ventral pathway, with cells that would normally contribute to the Ventral Stripe becoming stuck during migration. To test this, we quantitated the ectopic pigment cells and the cells of both the Dorsal and Ventral Stripes (Fig 2D-I). Our model predicted that the counts in the Dorsal Stripe would be unaffected, but that pigment cells in the Ventral Stripe might be reduced, perhaps in proportion to the number of ectopic cells. Quantitation of the ectopic pigment cells (melanocytes + iridophores) in the ventral medial pathway revealed a variable phenotype, with a consistently larger number of iridophores than melanocytes in the ectopic position (Fig 2D). The number of melanocytes in the Dorsal (Fig 2F) and Ventral (Fig 2G) Stripes in *pde* mutants was not statistically different to those of WT siblings. Similarly, there was no significant difference in the number of iridophores in the DS of *pde* mutants and WT siblings (Fig 2H). Interestingly, and in contradiction to the disrupted migration model, the number of iridophores in the ventral stripe of *pde* mutants shows a 58% *increase* compared to WT siblings (Fig 2I). Thus, we reject the disrupted migration hypothesis, and instead note that *pde* mutants display a regionally localised increase in
melanocytes and iridophores in the ventral trunk, with some as supernumerary cells in the Ventral Stripe, but many in an ectopic position nearby.

**pde encodes the endothelin receptor Aa gene**

To identify the mutation that causes the pde phenotype we crossed pde heterozygotes onto a WIK wild-type background, and mapped the locus of the mutation using two sets of microsatellite markers, G4 and H2 [20]. The pde mutation showed strong linkage to markers Z15424 and Z23059 on chromosome 1, ~1.5 cM and ~3.8 cM away from the mutation, respectively (Fig 3A). Using the 8th version of the zebrafish genome assembly, further analysis showed that the marker P249 in clone BX51149 was only 0.2 cM (4 recombinants in 17959 embryos) away from the mutation, placing the pde mutation about 132 kb away in clone CU462997, in which three genes were annotated: 1) mineralocorticoid receptor (now renamed nuclear receptor subfamily 3, group C, member 2, nr3c2), 2) Rho GTPase activating protein 10 (arhgap10), and 3) endothelin receptor Aa (ednraa). Of these three candidate genes, ednraa has been previously reported to be required in the development and patterning of the neural crest-derived ventral cranial cartilages. Furthermore, ednraa is expressed in the developing blood vessels of the posterior trunk [21]. This striking correlation between ednraa expression and the region with ectopic pigment cells in pde mutants made ednraa a strong candidate. Previous studies had not reported a pigment phenotype during morpholino-mediated knockdown, but the focus in that work had been on craniofacial development, reflecting another region of ednraa expression [21].

To test the possible role of ednraa in pigment development, we used morpholino-mediated knockdown by injection of 1-2-cell stage wild-type embryos using the previously validated splice-blocking ednraa-morpholino (ednraa-MO1; S1 Table and Fig 3B; [21]). As controls, we used injection of a random-sequence morpholino (cMO; standard control provided by Genetools) and the ednrab morpholino (ednrab-MO; S1 Table; [21]), neither of which resulted in ectopic pigment cells, nor any other disruption to the early larval pigment pattern (Fig 3C). In contrast, after injection of ednraa-MO1 we saw ectopic melanocytes and iridophores in the ventral medial pathway in a pde-like phenotype (Fig 3D). As a further test, we designed a new translation blocking morpholino against ednraa (ednraa-MO2, S1 Table); injection of this morpholino phenocopied the pde mutant pigment phenotype, but also was prone to non-specific deformations, likely due to off-target effects [Colanesi, 2011 #15447]. Together, these data strongly support the hypothesis that the pde mutants disrupt ednraa function.

To assess directly the link between pde mutations and disruption of ednraa function we amplified ednraa cDNAs from pde mutant alleles to examine mRNA sequence and structure from mutants. In addition to the two original alleles, pde\textsubscript{tj262} and pde\textsubscript{tv212}, we also analysed
a third mutation identified in an independent screen at the Hubrecht Institute which showed a phenotype indistinguishable from the original alleles, and which failed to complement those original alleles (S2 Fig). This cDNA sequencing showed that \( pde^{hu4140} \) has a single base transition mutation in the 3’ region of exon 5 (bp 847; AGA > TGA; Fig 3B), which is predicted to result in a premature translation stop in Ednraa. The \( pde^{j262} \) and \( pde^{v212} \) alleles each showed deletions, of 103 bp and 135 bp respectively. cDNA sequence alignment indicates a deletion of exon 7 and a couple of extra bases, predicted to cause a frame shift of translation in exon 8 in \( pde^{j262} \), while \( pde^{v212} \) has a deletion of exon 6 and a couple of extra bases which results in a frame shift of translation in exon 7 and 8 of Ednraa (Fig 3B). Given that the \( pde^{j262} \) and \( pde^{v212} \) alleles were isolated from an ENU-mutagenesis screen and hence are likely to result from induced point mutations, we propose that the mutations in \( pde^{j262} \) and \( pde^{v212} \) are likely to affect key bases involved in ednraa splicing.

The Ednraa protein is a member of the rhodopsin-like G-protein coupled receptor family (GPCR, class A), and as such is characterised by seven transmembrane domains. In silico translation and structural predictions for the \( ednraa \) alleles (Fig. 3B; S5 Fig A) indicate that the N-terminus and the early transmembrane domains are likely to be intact, but that the other transmembrane domains and the C-terminus are absent. Our three \( ednraa \) mutant alleles show indistinguishable phenotypes, and are fully recessive, consistent with the similar predicted molecular effects of the mutations, and strongly indicating that the receptor is not functional. Consequently, we propose that these alleles are all likely null mutants, although formal proof of this will require generation of an N-terminal truncation allele.

We used an \textit{in situ} hybridization time-course between 6 and 72 hpf to assess the domain of \( ednraa \) expression, and in particular, whether it was detectable in neural crest or pigment cells. Our data was fully consistent with the earlier demonstration of \( ednraa \) expression in the developing blood vessels, but we saw no evidence of neural crest expression in the trunk (Fig 3E-H). We then tested whether blood vessels morphology was affected in \( pde \) mutants, which we will refer to as \( ednraa \) mutants from now on. One possible explanation for the pigment cells might be that blood vessel morphology might be disrupted in \( ednraa \) mutants, resulting in misplaced pigment cells. However, neither \textit{in situ} hybridisation for \( ednraa \) (Fig 3E-H), nor examination of trunk blood vessel morphology using the transgenic line Tg(flia:GFP) (Fig 3I,J) showed differences between WT and \( ednraa \) mutant embryos. We conclude that gross morphology of blood vessels is not affected in the \( ednraa \) mutants, but that the supernumerary and ectopic pigment cells in the \( ednraa \) mutants result from a non-cell autonomous effect of endothelin signalling in the blood vessels.

\textbf{The \( ednraa \) phenotype does not result from neural crest cell transdifferentiation in the ventral trunk}

The location of the ectopic pigment cells on the ventral medial pathway corresponds with the
location of the nascent sympathetic ganglia, which form on the medial neural crest migration pathway in the vicinity of the dorsal aorta. We considered a transdifferentiation model in which neural crest cells fated to form sympathetic neurons switch to generating pigment cells, predicting that sympathetic neuron numbers would be reduced in ednraa mutants. However, immuno-detection of the early neuronal marker Elav1 (Hu neuronal RNA-binding protein) showed no differences in the number of sympathetic neurons between phenotypically WT embryos and their ednraa mutant siblings (Fig 4A, 4B and 4K). Furthermore, we also tested whether other neural crest-derived neurons are affected in the trunk, but neither DRG sensory neuron nor enteric neuron numbers differed between ednraa mutants and their WT siblings. Thus, at 5 dpf, trunk DRGs contained around 3 neurons per ganglion (Fig 4C, D and L), while the posterior gut contained around 125 enteric neurons (Fig 4E, F and M).

As a further test of the sympathetic neuron transdifferentiating hypothesis, we reasoned that if a key signal driving sympathetic neuron specification was reduced, this might result in enhanced numbers of ectopic pigment cells. In mammals and birds, secreted BMP signals from the dorsal aorta have been shown to induce sympathetic neurons [22-24], but it is not known if this mechanism is conserved in zebrafish. To test this, we treated zebrafish embryos with Dorsomorphin (2.5 μM), a well characterised BMP signalling inhibitor[25]. Given the well-known roles for BMP signaling in early patterning in the embryo, we chose a treatment window from 1-4 days post fertilisation (dpf). Although this treatment left the larvae looking morphologically normal, immunofluorescent detection of Elav1 showed that treated larvae had a strong reduction in the number of sympathetic neurons compared to DMSO carrier-treated controls (Fig 4G, H and N). Having shown that zebrafish sympathetic neurons were BMP-dependent, we then asked whether ectopic pigment cells in ednraa mutants were increased if sympathetic neuron specification was inhibited; using the same treatment conditions, we saw no enhancement of ectopic pigment cells in ednraa mutants treated with dorsomorphin compared to DMSO-treated controls (Fig 4I, J and O). Thus, although we provide the first evidence to our knowledge that specification of zebrafish sympathetic neurons is BMP-dependent, we discount the hypothesis that the ectopic pigment cells in ednraa mutants result from transdifferentiating sympathetic neurons.

The ednraa phenotype results from localised increased neural crest cell proliferation in the ventral trunk, in the vicinity of the medial blood vessels

In order to identify the earliest stage at which ectopic pigment cells appear in the ventral trunk of ednraa mutants, we performed in situ hybridization with the melanocyte marker dct and the iridoblast marker ltk. Comparison of gene expression between WT and ednraa mutant embryos at 24, 30 and 35 hpf showed that ectopic/supernumerary expression of both dct and ltk is detected in the ventral trunk from 35 hpf (Fig 5A-D).
We tested whether this production of extra pigment cells correlated with an increased proliferation of NC-derived cells in *ednraa* mutants. Given that ectopic and supernumerary pigment cells appear in the ventral trunk of *ednraa* mutants between 30 and 35 hpf, this indicates that enhanced proliferation might be expected to be detectable during this time-window. We used our Tg(sox10:cre) [26] combined with a Tg(hsp70:loxP-dsRed-loxP-lyn-egfp) red-green switch reporter to label all neural crest cells with membrane tagged GFP (GFP fused with the membrane LYN Tyrosine-kinase), in conjunction with expression of the proliferation marker phosphohistone H3 (pH3) by immunofluorescent labelling. GFP expression in the neural crest-derived cells was induced by brief heat shock at 30 hpf. Subsequent double labelling of GFP and pH3 at 32 hpf showed a 55% overall increase in NC-derived proliferating cells in *ednraa* mutants compared with WT siblings, albeit over a very low basal rate (Fig 5E, F and G). Furthermore, over-proliferation of neural crest cells was detectable only in the ventral region of the medial pathway (Fig 5G). Our results show that ectopic pigment cells in *ednraa* mutants result from overproliferation of NC-derived cells in a localised region of the medial pathway in the vicinity of the dorsal aorta, coinciding with the region of *ednraa* expression.

During the proliferation assays, we noted that the ventral proliferation of neural crest-derived cells was often clustered at the ventral end of the migrating streams (arrowheads, Fig 5F). Furthermore, this association was true also of the ectopic pigment cells themselves, as shown by imaging of all NC-derived cells using the same transgenic line (Tg(-4725sox10:cre)ba74; Tg(hsp:loxP-dsRed-loxP-EGFP))(S3 Fig). This imaging confirmed the widespread ventral migration of neural crest-derived cells in *ednraa* mutants, comparable to that in WT siblings, at 35 hpf (S3 Fig A,B), reinforcing the conclusion from our WISH studies (Fig 1 and 5). At 5 dpf, this imaging readily showed the ectopic pigment cells, but clearly revealed that their location is ventral to the notochord, in close proximity to the dorsal aorta and associated with the sympathetic ganglial chain, and also near the ventral projections of the spinal nerves (S3 Fig C-F).

The formation of ectopic pigment cells in *ednraa* mutants requires ErbB signalling.

In order to identify possible mechanisms involved in the formation of the *ednraa* phenotype, we performed a small molecule screen of 1396 compounds using three different libraries: 1) the Sigma LOPAC library, which contains 1280 small organic ligands including marketed drugs and pharmacologically relevant structures; 2) the Screen-WellTM Kinase Inhibitor Library that comprises 80 known kinase inhibitors; and 3) the Screen-WellTM Phosphatase Inhibitor Library, containing 33 phosphatase inhibitors of well-characterised activity. Screening was performed using the methodology of our previous screen for pigmentation modifiers [27]; we took advantage of the adult viability of the *ednraa* mutants, to perform the screen on *ednraa* mutant embryos. Thus *ednraa* mutant embryos were treated from 4 hpf
to 96 hpf at a standard concentration of 10 μM and rescue or enhancement of ednraa mutant phenotype was systematically assessed at 4 dpf. After rescreening, we identified 23 compounds able to rescue the ednraa mutant phenotype, as well as 3 that enhance the phenotype (S2 Table). Of these hits, four (Tyrphostin AG 1478, U0126, PD 98059 and PD325901) target the MAPK/ERK pathway: Tyrphostin AG-1478 is an inhibitor of the Epidermal Growth Factor Receptor (EGFR)[28] while U0126, PD98059 and PD325901 are highly selective inhibitors of MEK1/2 signalling [29-31]. Our previous work has shown that MEK inhibitors interfere with production of regenerative melanocytes, whilst not affecting direct developing pigment cells forming the early larval pattern in zebrafish [32, 33]. Treatment of ednraa mutants with each of two of the latter compounds shows a clear dose-response in the degree of rescue of the ednraa phenotype (S4 Fig).

The identification of Tyrphostin AG-1478 in our screen is especially notable, because it and the more specific EGFR inhibitor, PD158780, have been shown to affect APSC biology, but not embryonic pigment cells (except a small population contributing to the Lateral Stripe), in zebrafish [10]. Similarly, mutants for the epidermal growth factor receptor (EGFR)-like tyrosine kinase erbb3b (picasso), despite developing a normal embryonic/larval pigment pattern, fail to develop a normal adult pigment pattern and are unable to regenerate melanocytes after embryonic melanocyte ablation due to a lack of APSCs [10]. Thus inhibition of Erb signalling by either AG-1478 or PD158780 selectively affects the biology of NC-derived APSCs that give rise to adult and regenerative melanocytes. The rescue of the ednraa mutant by treatment with the Erb inhibitors suggested the exciting hypothesis that the ectopic cells in the ednraa mutants might result, not from embryonic pigment cells, but by precocious differentiation of APSCs. Interestingly, this role for ErbB signaling shows a tightly constrained time window, since inhibition of ErbB signalling with AG-1478 and PD158780 during window 9-48 hpf of embryonic development is sufficient for these effects [10]. Moreover, after melanocyte ablation throughout 24-72 hpf, melanocyte regeneration normally occurs by 5 dpf, but this is prevented when embryos are treated with AG-1478 during a 9-30 hpf window [7].

Thus, our hypothesis makes the testable prediction that rescue of the ectopic pigment cells in ednraa mutants would share the very specific temporal window known to regulate APSCs development. We treated ednraa mutant embryos with a range of concentrations (0.1 – 2.0 μ M) of PD158780 from 12-48 hpf (Fig 6A-D). Quantification of the number of ectopic pigment cells showed that ednraa mutant embryos treated with PD158780 have a significant dose dependant-reduction in the number of ectopic cells compared to DMSO treated embryos (Fig 6M). Moreover, shorter treatment with PD158780 revealed that the ednraa phenotype is effectively rescued when treatment is restricted to a 19-30 hpf window (Fig 6E-H and 7M,), showing a striking match to the critical period for establishment of APSCs, while treatment after this window (24-30 hpf) does not rescue the ednraa phenotype (Fig 6I-M). This data
strongly supports the hypothesis that the source of ectopic pigment cells in ednraa mutants is likely to be APSCs and not embryonic pigment cells.

To date there is only one marker that has been proposed to label APSCs: Dooley and colleagues identified cells expressing \textit{Tg(mitfa:GFP)} in the DRGs as APSCs as early as 12 dpf (Dooley et al., 2013). Thus we assessed the expression of this transgene in WT larvae, focusing on the vicinity of the dorsal aorta. At 8 dpf, we indeed found a significant number of \textit{GFP$^+$} cells throughout this region in the posterior trunk and anterior tail (S6 Fig), consistent with our proposal that APSCs may reside in this ventral position in addition to the DRG location documented before.

**Discussion**

In this study we show that \textit{ednraa} encodes one of two zebrafish Ednra orthologues, Ednraa. In mammals the EdnrA receptor binds selectively to Edn1 and Edn2, mediates vasoconstriction, and is overexpressed in many cancers[34]). In contrast, loss of function in mouse results in homeotic transformation of the lower jaw towards an upper jaw morphology, and in humans underlies Auriculocondylar syndrome (ACS [MIM 602483 and 614669]) [35-38]. These studies did not identify pigmentation phenotypes, although an \textit{Ednra lacZ} knockin mouse strain shows prominent expression in the hair follicles [35]. In contrast, \textit{Ednrb} and \textit{Edn3} mutants, as well as mutations in the Edn-processing enzyme Ece1, lack neural crest-derived melanocytes [39-41].

Analysis of the zebrafish genome identifies eleven components of endothelin signalling system: Four ligands, Edn1, Edn2, Edn3a and Edn3b; three Endothelin Converting Enzymes, Ece1, Ece2a, Ece2b that activate the ligands; and four receptors, Ednraa, Ednrab, Ednrba and Ednrbb [42]. In adult zebrafish, \textit{ednrba} and \textit{ece2b} loss-of-function mutants have all been shown to display reduced iridophores and broken stripes, indicating their coordinated role in iridophore development and pigment patterning, but no effect on embryonic pigment pattern [43, 44]. In contrast, \textit{edn1} mutants and \textit{ednraa} and \textit{ednrab} morphants revealed disruption of the lower jaw, similar to the mammalian role in dorsoventral patterning, but did not examine pigmentation [21].

Here we identify a novel function for Ednraa signaling in pigment cell development. Our experimental studies assess in turn a series of hypotheses regarding the embryonic basis of the ectopic pigment cell phenotype, initially exploring disrupted biology of embryonic (direct developing) pigment cells, before coming to the conclusion that the phenotype must result from disruption of APSC biology. The \textit{ednraa} mutant phenotype is restricted to supernumerary and ectopic pigment cells in the ventral trunk and anterior tail; this spatial
Localization had been perplexing, but identification of the gene as *ednraa*, which is consistently expressed strongly in developing blood vessels (but not in the trunk NC) from well before the onset of detectable ectopic cells, helps to explain the restricted phenotype of *ednraa* mutants. We find no evidence that neural crest migration is disrupted, since not only is blood vessel morphology normal, but neural crest cell migration and patterning is normal except for the ectopic pigment cells themselves, and pigment cells are abundant in the Ventral and Yolk Sac Stripes. Similarly, we disprove the ‘mixed-fate’ hypothesis by showing using TEM that the ectopic cells are indistinguishable from those melanocytes and iridophores in the Yolk Sac Stripe where these two cell-types are consistently found in tight apposition. Instead, we conclude that the close association of the two cell-types reflects the natural tendency for iridophores to adhere to melanocytes (as seen in all locations in the early larval pattern). Finally, we find no evidence that the ectopic cells derive from transfating of sympathetic neurons, since neurons are unaffected in *ednraa* mutants and even when sympathetic neuron specification is inhibited using a BMP inhibitor, ectopic pigment cell number is unchanged.

However, these studies were unable to address the possible role of glial or progenitor cells in the ventral trunk. We reasoned that the supernumerary and ectopic cells were likely to be associated with increased proliferation of a subset of neural crest cells. Strikingly, we found that enhanced proliferation of neural crest did characterise the *ednraa* mutants, but that this proliferation was specifically associated with neural crest cells in ventral regions i.e. near the dorsal aorta, at around the time (35 hpf) that ectopic pigment cells begin to be identifiable. This is before the sympathetic ganglia show detectable differentiation and suggested that a subset of NC-derived cells undergoes ectopic or precocious proliferation. A key insight into the identity of these cells came from the observation that both ErbB inhibitors rescued the homozygous phenotype. Inhibition of Erb signaling *within a defined embryonic time window* by either AG-1478 or PD158780 selectively affects the biology of NC-derived APSCs that give rise to adult and regenerative melanocytes [7, 10]. We show that *ednraa* mutants are rescued by ErbB inhibitors in a dose- and time-dependent manner, and with a time-window overlapping that known to regulate APSC development. We conclude that the ectopic pigment cells in *ednraa* mutants result from precocious differentiation of pigment cells from APSCs, cells that would normally be quiescent until metamorphosis. An alternative model, that the ectopic cells result from simple overproliferation of embryonic melanoblasts and iridoblasts in the ventral region would be consistent with the localised enhanced proliferation, but in our pH3 labeling studies, labelled melanocytes are rare (1 in 20 WT embryos, and 0 in 20 *pde* mutants). Furthermore, this model would not explain the observations that ErbB inhibition rescues the *pde* mutant phenotype.

APSCs give rise to the numerous melanocyte and iridophores of the adult skin [11]. Our model would be consistent with the lack of ectopic xanthophores, since the evidence to date is that the majority of adult xanthophores derive from embryonic xanthophores and not from
APSCs, although a small contribution from the stem cells is indicated by clonal analyses [11, 45]. Finally, the localised increase in neural crest cell proliferation that we document is also consistent with the activation of otherwise quiescent stem cells. APSCs in zebrafish have been closely-linked to the developing peripheral nervous system. One intriguing aspect of our data is that this proliferation is exclusively localised to the ventral trunk, consistent with the idea that the blood vessels form a key aspect of the stem cell niche, but surprising in that to date APSCs have been exclusively associated with the DRGs [9, 46]. We note that homozygous ednraa mutants are adult viable, but have no visible skin pigment pattern defect. Our data are consistent with a second source of APSCs, associated with the peripheral nervous system in the ventral trunk, likely the sympathetic ganglia. We propose a model in which these APSCs reside in a novel niche in close proximity to the medial blood vessels, which provide signals holding them in a quiescent state (Fig. 7). In the ednraa mutants, these signals are reduced or absent, such that the APSCs become precociously activated. They then undergo proliferation and differentiate as melanocytes and iridophores. Many of these move into the Ventral Stripe location, but the excess cannot be accommodated in this stripe and remain ectopically located in the ventral trunk near the sympathetic ganglia. It is somewhat surprising that homozygous ednraa mutants show no disruption of the adult pigment pattern, but we propose several hypotheses, none mutually exclusive, to explain this observation: 1) although as we have shown APSCs are precociously activated in ednraa mutants, APSC renewal may be unaffected so that the stem cells are not exhausted; 2) ventral APSCs may primarily populate internal, rather than skin, pigment cell populations; 3) APSCs localised in DRGs may compensate for the effects on the more ventral APSCs. A key future test of this model will require direct evidence for the presence of a quiescent set of APSCs in the vicinity of the medial blood vessels.

Although novel in the context of APSCs, blood vessels form an important part of adult stem cell niches in other contexts, especially that of adult Neural Stem Cells (NSCs) [47]. Indeed in the case of NSCs in the subependymal zone of mouse, blood vessel-mediated signals play a role in maintaining quiescence[47], although other sources for these quiescence signals have also been identified in the neurepithelial component of this niche [48-51]. In these studies of NSCs, notch signaling has been shown to be important in maintenance of quiescence, so it will be fascinating to compare the molecular signals derived from the vasculature that regulate APSC quiescence in the zebrafish. Our discovery of a second niche for APSCs and identification of the key role for blood vessels in controlling their behaviour provides an entry point for uncovering an important but currently understudied aspect of zebrafish pigment pattern formation.

Our work may also have implications for human disease. For example, Phakomatosis pigmentovascularis (PPV) is a rare mosaic disorder defined as the simultaneous occurrence of a widespread vascular nevus and an extensive pigmentary nevus, and associated with activating mutations of Gα subunits of heterotrimeric G proteins [52]. It is currently
unknown if there is a common progenitor that gains an oncogenic mutation and leads to both large nevi and vascular proliferations, or if changes in one cell type impact upon another. Our study provides evidence that melanocytes can expand/differentiate in association with a modified blood vessel niche, making it conceivable that, for instance, oncogenic activation in blood vessels might result in non-cell autonomous activation of melanocyte stem cells in the niche to generate a nevus.

Materials and Methods

Fish husbandry

All fish strains were maintained in accordance with local and national guidelines on animal welfare. This study was performed with the approval of the University of Bath ethics committee and in full accordance with the Animals (Scientific Procedures) Act 1986, under Home Office Project Licenses 30/2937 and P87C67227. Animal experiments were approved by the Animal Experimentation Committee (DEC) of the Royal Netherlands Academy of Arts and Sciences. AB, pde\textsuperscript{j262}, pde\textsuperscript{v212} and pde\textsuperscript{hu4140}, Tg(\textit{flia}:GFP) and Tg(-4725\textit{sox10:cre})ba74; Tg(hsp:loxp-dsRed-loxp-EGFP).

Genetic mapping

Mapping panels

Two specific sets of microsatellite markers, the G4 and H2 panels (Geisler et al., 2007) were used for bulk segregant analysis of \textit{parade}, placing the mutation on linkage group 1. We subsequently used a consolidated meiotic map of the zebrafish genome, ZMAP, which is available on ZFIN (http://zfin.org/cgi-bin/webdriver?MIval=aa-crossview.apg&OID=ZDB-REFCROSS-010114-1; Sprague et al., 2006; Geisler et al., 2007).

Reference zebrafish line

The standard mapping wild-type line WIK was crossed with mutant \textit{parade}\textsuperscript{j262} carriers (in AB wild-type background). This founder generation F0 was incrossed to produce heterozygous F1. Eight pairs of F1 fish were incrossed to produce the F2 generation. In total 1796 F2 embryos at 5 dpf were sorted into two groups, those that display the mutant phenotype and those with a normal wild- type pigment phenotype. These were transferred into 96-well plates where the extraction of genomic DNA was performed and stored. Additionally, genomic extracts of 10 homozygous \textit{parade}\textsuperscript{j262} and 10 wild-type sibling embryos of each of the 8 parent pairs were pooled for identification of new SSLP markers.

Mapping procedure

The \textit{pde} locus was mapped to linkage group 1 (LN 1) by bulked segregant analysis using pooled genomic DNA of 48 wild-type siblings and 48 \textit{parade}\textsuperscript{j262} mutant embryos of the F2 generation obtained from F1 Pair 2. For fine mapping we designed new mapping primers using the Primer3 online software tool (http://frodo.wi.mit.edu/primer3/, default settings plus...
1 primer pair per 300–400 bp, (Rozen and Skaletsky, 2000)). Primer pairs were designed to generate PCR products of 300-400 bp, which if generating polymorphic PCR amplicons, were then tested on the 1796 single F2 embryos to determine the frequency of recombination.

PCR protocol

2 μl of individual or pooled DNA were added to 13 μl of PCR mix (1.5 μl 10 x buffer for KOD Hot Start DNA Polymerase; 1.5 μl dNTP’s with 0.2 mM for each nucleotide; 1 μl primer mix with 10 mM forward and 10 mM reverse primer; 0.3 μl DMSO; 0.6 μl 25 mM MgSO4; 8.3 μl MiliQ water; 0.3 μl KOD polymerase (Novagen)). PCR’s were run in a GStorm thermal cycler (Gene Technologies Ltd) (program: 2 min 94° C; then 35 x (30 sec 94° C, 30 sec 60° C, 30 sec 72° C); then 5 min 72° C; then stored at 10° C). The PCR products were then analysed by electrophoresis.

Morpholino injection

Custom morpholinos were purchased from Gene Tools LLC (Philomath, USA) and resuspended in autoclaved MilliQ water to a stock dilution of 20 μg/μl. Stock solutions were stored at -20° C to prevent evaporation and heated at 70° C for 5 min prior to dilution to eliminate precipitates. Wild-type embryos were injected into the yolk at the 1-cell stage with 5 ng/ml dilutions. Phenotypes were observed under the microscope at 3 dpf and 4 dpf. Volumes varied between 5 and 10 nl per embryo. Sequences of morpholino oligos can be found in S1 Table.

Whole mount in situ hybridization

Our protocol for whole-mount in situ hybridization is based on (Thisse et al., 1993). All solutions were prepared with DEPC-treated autoclaved MiliQ water or PBS. All probes for in situ hybridization were synthesized using the Dig RNA Labeling Kit (Boehringer). Depending on the orientation of the gene and choice of plasmid, we have chosen either T3, T4 or SP6 RNA polymerases for creating the antisense RNA probe.

Briefly, embryos between 5 and 120 hpf were euthanized with a lethal overdose of tricaine and fixed overnight (4 % PFA in PBS). Embryos older than 18 hpf were dechorionated with forceps before fixation, younger embryos were dechorionated with flamed forceps after fixation. All following steps were carried out in 1.5 ml microfuge tubes with 1 ml liquid volumes; up to 40 embryos were processed per tube. The fixative was rinsed out with PBTween (DEPC-treated PBS plus 0.1 % Tween). Embryos were then gradually dehydrated on ice with 25, 50, 75 and 100 % methanol. For the ISH procedure, embryos were rehydrated on ice using 5 min washes with 75, 50, 25 % methanol followed by 3 x 5 min washes with PBTween at room temperature. To improve permeability, embryos between 30 – 48 hpf were treated for 10 min with 0.01 mg/ml proteinase K/PBTween, embryos; older than 50 hpf were treated for up to 20 min. The proteinase K was removed by washing samples for 3 x 5 min with PBTween followed by a brief re-fixation step with 4% PFA for 20 min at room
temperature. The fixative was then washed out with 3 x 5 min PBTween. To prepare the samples for optimal binding conditions, embryos were pre-hybridized with hybridization mix (formamide, 20 x SSC, heparine, tRNA, Tween20, citric acid; stored at -20° C) in a water bath at 65° C for 4 h. The hybridization of the RNA probe (diluted 1:400) was done overnight at 65° C in 200 μl hybridization mix. The samples were washed with decreasing concentrations of hybridization mix (100 % hyb mix; 75 % hyb mix/25 % 2x SSCT; 50 % hyb mix/50 % 2x SSCT; 25 % hyb mix/75 % 2x SSCT; 100 % 2x SSCT), each for 10 min, followed by 2 x 30 min washing with 0.2x SSCT. All steps were carried out in a water bath at 65° C. Samples were then infiltrated for 3 x 5 min with MABT (from 5 x MAB stock solution plus 0.1 % Tween) at room temperature. Samples were blocked for 3 h in MABTween with 5 % sheep serum at room temperature under gentle shaking. Antibody binding reaction was performed over night at 4° C using 200 μl of 1:5000 diluted antidioxigenin alkaline phosphatase conjugated antiserum in blocking solution. Samples were then washed for 6 x 15 min in MABT, then prepared for signal detection reaction by infiltrating for 3 x 5 min with NBT/BCIP buffer (100 mM Tris HCl pH 9.5, 50 mM MgCl2, 100 mM NaCl, 0.1 % Tween). The embryos were transferred into 9-well glass dishes to observe the staining reaction under the dissecting microscope. 300 μl of BMPurple substrate solution were added per well and the development of the reaction regularly monitored. Signal development was stopped by washing with PBTween.

Chemical screening

Compounds used

The compounds used in this investigation originated from the 1280 compounds from the LOPAC library (Sigma LO1280), 80 compounds from a kinase inhibitor library (Biomol 2832) and 33 compounds from a phosphatase inhibitor library (Biomol 2834A). The compound libraries were stored at -80C and plates thawed at room temperature prior to use. For screening potential effects on developing zebrafish embryos compounds were diluted to 10μM in E3 embryo medium.

For drug treatments, six embryos were placed in each well of 24 well plates. Chemical treatments were prepared in 10μM doses in 1ml E3 medium. To ensure that compounds did not precipitate, the plates containing small molecule treatments were placed on The Belly Dancer (Sorval) for a minimum of 20 minutes. When embryos reached 8hpf E3 medium was removed and 1ml of E3 containing 10μM of compound was added to each well. Wells containing E3 only and 10μM DMSO were used in each chemical treatment as controls. Plates containing treated embryos were incubated at 28C until the embryos reached 5dpf. Detailed observations were recorded daily and any dead embryos removed to avoid contamination of the medium. At day 5 embryos were anaesthetised with a small dose of tricaine to enable easy manipulation for accurate detailing of pigment phenotype. Pigment cell phenotypes and rescue of the parade mutant phenotype were recorded as well as additional phenotypes such as developmental abnormalities (e.g. shortening of the tail or cardiac edema).
Re-screening of important chemical ‘hits’

Once molecules of specific interest to pigment cell development were identified the treatment was repeated on embryos at different developmental time points and across a concentration gradient in order to identify the developmental time point of greatest phenotypic significance and the optimal small molecule concentration.

Whole mount immunostaining

Antibody staining was largely performed as described in [19]. Prior to primary antibody incubation, embryos were permeabilized with 1 ml of 5 μg/ml of proteinase K/distilled water during 30 minutes at 37 °C, then rinsed with 5% of goat serum/distilled water at RT for 5 minutes and washed 3x1 hour with distilled water at RT. The antibodies used were Hu (1:500,) and Alexa Fluor 488 goat-mouse IgG (1:750, Molecular Probes, Cat. # A11001).

Image acquisition and processing

For in vivo microscopy, embryos were mounted at the Leica Fluo III or Zeiss Axiovision dissection microscope in 0.6 % agarose or 3 % methyl cellulose. For anaesthesia, 0.02 % Tricaine was added freshly to the mounting medium. The embryos were transferred to a microscope slide or to a glass-bottomed Petri dish. For immunofluorescence, embryos were visualized with a Nikon Eclipse E800 or Zeiss M2 Imager compound microscope under bright field, incident light or epifluorescence illumination, mainly using a 10x, 20x and 63x objective for magnification. Images were taken with a DS-U1 (Nikon) or Orca and color 546 camera (Zeiss). Confocal fluorescence imaging was performed on an inverted Zeiss LSM 510 Meta or LSM 880 confocal microscope with 20x and 40x objectives. Incident light was provided by installing an additional antler lamp (Leica) around the microscope to allow visualisation of iridophores. NIS-Elements D2.30, Zen blue, Fiji and/or Adobe Photoshop 7 were used to adjust white balance and exposure and to assemble the images to figures.

Acknowledgments

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References


**Figure legends**

**Figure 1** *pde* mutants display ectopic melanocytes and iridophores, but not xanthophores, in the ventral medial pathway. (A,B) Overview of early larval WT (A) and *pde* (B) pigment phenotype at 5 dpf. (C-F) Anatomical location of ectopic pigment cells in *pde*. Magnification of lateral views (white boxes in A and B) and cross sections of posterior trunks show no pigment cells on the medial migration pathway in the ventral trunk of WT larvae (C and D). Ectopic pigment cells are located in the ventral trunk of *pde* mutants (E; red arrowhead), under the dorsal aorta (DA) and above the posterior cardinal vein (PCV) as shown by cross sections (F, white arrowhead). (G-L) Whole mount in situ hybridization of 3
dpf WT (G-H) and pde mutants (J-L) embryos for dct (G and J), ednra (H and K), and gch (I and L). Ectopic dct (J; white arrowhead) and ednra (K; white arrowhead) expression is seen in the ventral trunk of pde mutant larvae. Neural tube (NT), notochord (NC) and ventral stripe (VS). Scale bar = 500 μm (A and B) and 100 μm (C and F) and 50 μm (G-L).

Figure 2 pde mutants show supernumerary melanocytes and iridophores in the Ventral Stripe and nearby medial migration pathway, but not the Dorsal Stripe. (A-C) Transmission electron photomicrographs of ectopic pigment cells in pde mutants. Magnifications of yellow (B) and blue (C) boxes show melanosomes (m) and reflecting platelets (p) separated by a double membrane (C; white arrowheads). (D) Dot-plot of quantitation of ectopic melanocytes (M), iridophores (I) and overall number of ectopic pigment cells (sum of melanocytes and iridophores; M+I) from individual pde mutant larva at 4 dpf reveals a variable phenotype, with a consistently larger number of iridophores than melanocytes in the ectopic position (iridophores mean + s.e.=17.26±0.99; melanocytes mean + s.e.=2.44±0.34, n=27). (E) Regions where number of pigment cells were counted, Dorsal Stripe (DS; orange line), Ventral Stripe (V; green line), posterior Ventral Stripe (PVS; pink line) and ventral medial pathway (VMP, red box). (F-I) Quantitation of number of melanocytes in dorsal (F; p<0.05, two-tailed t-test, WT mean + s.e.= 78.8±2.5, n=20 and pde m + s.e.=72.8 ± 2.8, n=20) and posterior ventral stripes (G; wild-type: mean + s.e. 55.0 ± 2.5 n=20; pde; 56.6 ± 1.8, n=17) show no significant (ns) difference between WT and pde mutants. Iridophore quantitation in the DS (H; p>0.05, two tailed t-test, WT mean + s.e. = 22.9 ± 0.9, n=29; pde mean + s.e. = 20.4 ± 1.1, n=22) is not different between WT and pde mutants while the ventral stripe has a 58% increased number of iridophores compared to WT embryos (I;***, p<0.0001, two-tailed t-test; WT mean + s.e. = 25.5 ± 0.6, n=49; pde mean + s.e. = 42.8 ± 1.0, n=43). Scale bar = 500 μm (E).

Figure 3 pde mutations affect ednraa, but not blood vessel formation and patterning. (A) pde map position on chromosome 1. (B) Schematic of predicted mRNA structure based upon sequencing of cDNA from pde^{ed262}, pde^{tv212} and pde^{hu4140} mutants. cDNA of pde^{ed262} mutants lack exon 7, pde^{tv212} lack exon 6 and pde^{hu4140} have a transition mutation (AGA847TGA) that causes a premature translation stop (triangle) in the 3’ region of exon 5. Predicted changes to EdnRaa protein are shown with respect to relevant extracellular domain (ECD), transmembrane domains (TD) and intracellular domains (ICD) of the multipass receptor protein. The location of the ednraa splice-blocking morpholino (MO-ednraa) is indicated (red bar). (C-D) Injection of ednraa morpholino into WT embryos phenocopies pde mutant pigment phenotype. (C) WT embryos injected with a control morpholino (MO-control) display a normal phenotype. (D) WT sibling injected with MO-ednraa display ectopic melanocytes (black arrowheads) and iridophores (white arrowheads) in the ventral medial pathway. (E-H) Whole mount in situ hybridization of ednraa at 24 hpf and 36 hpf is restricted to the developing blood vessels, and is indistinguishable between pde mutants (F...
and H) and their WT siblings. (I-J) Imaging of blood vessels in the posterior trunk using the transgenic reporter flia:GFP shows no difference in blood vessel morphology between WT siblings (I) and pde mutants (J). DA, Dorsal Aorta; PCV, Posterior Cardinal Vein; Se, Segmental Vessels. Scale bar = 100 μm (C,D, E and J) and 25μm (E-H).

**Figure 4** Neural crest-derived peripheral neurons are not reduced in pde mutants, but a role for BMP signaling in sympathetic neuron specification is conserved in zebrafish. Immunodetection of the neuronal marker Hu in 7 dpf WT embryos (A) and pde mutant siblings (B), shows no significant difference (ns) in the number of sympathetic neurons (K; WT mean+s.e.=14.0±1.27, n=18 and pde=12.78±0.76, n=18, n=18; p>0.05, two-tailed t-test). (C-F) Immunodetection of the neuronal marker Hu in 5 dpf WT embryos (C and E) and pde mutant siblings (D and F) shows no significant (ns) difference in the number of sensory neurons per dorsal root ganglion (DRG; C and D; WT=3.26±0.11, n=15 and pde =3.08±0.14, n=12; p>0.05, two-tailed t-test) nor in enteric neurons in the posterior gut (E, F and M; WT =132.3±7.14, n=8 and pde =119.4±8.49, n=8; p>0.05, two-tailed t-test). (G-J) Chemical inhibition of BMP signalling with dorsomorphin (iBMP 2.5 μM; 1-4 dpf treatment). Treatment of WT embryos shows a 63.25% reduction in the number of sympathetic neurons in comparison with 1% DMSO-treated controls (G, H and N; DMSO =13.28±0.91, n=18 and dorsomorphin=4.88±0.47, n=18; p<0.0001, two-tailed t-test). Quantification of the number of ectopic pigment cells in pde mutants treated with DMSO (I) or dorsomorphin (iBMP 2.5 μM; 1-4 dpf treatment; J) shows no significant (ns) difference (O; DMSO mean+s.e.=19.54±1.12, n=13 and pde =20.25+1.30, n=12; p>0.05, two-tailed t-test). Weak red fluorescence in the fluorescent images result from autofluorescence of red blood cells. Scale bar = 100 μm (A-H same scale bar and I and J same scale bar).

**Figure 5** Ectopic pigment cells in pde mutants are detectable by 35 hpf, and generated by localised increased proliferation of neural crest-derived cells. (A-D) Whole mount in situ hybridization of 35 hpf WT (A and C) and pde mutant (B and D) embryos shows ectopic expression in pde mutants (white arrowheads in D) of melanocyte marker dct (A and B) and the iridophore marker ltk (C and D). (E-G) Immunodetection of the proliferation marker phosphohistone 3 (PH3) in neural crest derived cells (labelled with membrane –tethered GFP due to Tg(-4725sox10:cre)ba74; Tg(hsp:loxp-dsRed-loxp-LYN-EGFP)) of 32 hpf WT (E; white arrowhead) and pde mutant (F, white arrowheads) sibling embryos. Quantification of double positive GFP+ PH3+ cells in medial migratory pathway, shows a significant increase in pde mutants compared to WT siblings (Total; WT mean+s.e.=3.9±0.42, n=20 and pde=6.05±0.41, n=20; p<0.0009, two-tailed t-test). Subdividing this quantification of GFP+ PH3 cells in the medial migratory pathway into those dorsal and ventral to the notochord shows that this increase is not detected on the dorsal medial pathway (dorsal double positive cells, WT=2.35±0.35, n=20 and pde =2.85±0.35, n=20; p<0.3188, two-tailed t-test), but is
significantly increased on the ventral medial pathway WT = 1.55 ± 0.30, n = 20 and pde = 3.2 ± 0.32, n = 20; p < 0.0007, two-tailed t-test). Scale bar = 30 μm (A-D) 15 μm (E-F).

Figure 6 Chemical inhibition of Erb signalling rescues the pde phenotype. (A-H)
Treatment of pde embryos with increasing concentrations of Erb inhibitor PD158780 (iErb; 0.5-2.0 μM) or DMSO carrier control from 12-48 hpf (A-D), 19-30 hpf (E-H) and 24-30 hpf (I-L) hpf. Quantification of the number of ectopic pigment cells in the ventral medial pathway showed a decrease in the number of ectopic cells when embryos were treated from 12-48 hpf or just from 18-30 hpf, but not in a later 24-30 hpf time-window (M). Scale bar = 200 μm (A-L).

Figure 7 Model for the role of EdnrA signaling in pigment cell development. A second source of APSCs is held in a quiescent state by Ednraa-dependent factors from the blood vessels. Figure shows a model integrating our observations with current knowledge. 1) Dorsal root ganglia associated APSCs (APSC) are maintained in a quiescent state by local factors (red); 2) we propose a second source of APSCs in the vicinity of the medial blood vessels. Ednraa/pde activity in the blood vessels results in signals that hold this novel population in a quiescent state. In the pde mutants, these factors (red) are lost locally from the blood vessels and the APSCs become precociously activated, generating melanocytes and iridophores in their vicinity.

S1 Table Morpholino sequences. A random sequence morpholino (cMO) provided by Gen Tools and a splice-blocking morpholino against ednrab (ednrab-MO) were used as control. ednraa was targeted with either a splice-blocking morpholino (ednraa-MO1) or a translation-blocking ednraa morpholino (ednraa-MO2). Scale bar = 100 μm (A-C).

S2 Table Screening of small molecules in pde mutants. Name and known targets of small molecules that enhance (E) or rescue (R) the pde phenotype.

S1 Figure Melanocytes and iridophores in the WT yolk sac stripe are consistently separated from each other by double membranes. Transmission electron photomicrographs of melanocytes and iridophores in the WT yolk sac stripe ectopic pigment cells in pde mutants. A and B show two examples of melanosomes (m) and reflecting platelets (p) separated by a double membrane (white arrowheads). Bright-field image of ventral view (C) and close up of the area in the red box (D) of WT fish, shows yolk sac stripe. Continuous layer of iridophores is indicated by white arrowhead in D, closely associated black melanocytes forming contiguous layer immediately dorsal to iridophores is indicated by red arrowhead. Scale bars = 500 μm (C) and 100 μm (D).
S2 Figure Complementation assay of the pde alleles. Overview of early larval pigment phenotype at 5 dpf of pde$^{j262/j262}$ (A), pde$^{j262/v212}$ (B) and pde$^{j262/hu4140}$ (C). All three allele combinations show ectopic melanocyte and iridophores in the ventral medial pathway of the posterior trunk.

S3 Figure Migration of neural crest cells through the medial migratory pathway. Labelling of neural crest derivatives with GFP using the transgenic line Tg(-4725sox10:cre)ba74; Tg(hsp:loxp-dsRed-loxp-LYN-EGFP) shows no difference between 35 hpf WT fish (A) and pde mutants (B), neural crest cells migrate ventrally in a intersegmental arrangement (white line in A and B). 5 dpf pde mutant larvae show ectopic pigment cells (white arrow in D) associated with the spinal nerve projections (arrowheads in D) that emerge from the dorsal root ganglia (DRG). Ectopic pigment cells (white arrows) are also associated with the sympathetic ganglion (SyG) chain that forms perpendicular to the spinal nerve projections (white arrowhead in E and F) and ventral to the notochord (No). Guided by DIC image, dorsal edge of the dorsal aorta (DA) is highlighted with a dashed white line in C-F. Neural tube (NT). DAPI labels nuclei (blue). Scale bar = 25 μm (A and B), 50 μm (C and D) and 15 μm (E-F).

S4 Figure Inhibition of MEK rescues the pde phenotype. Treatment with increasing concentrations of the MEK inhibitors U0126 (2.5-7.6 μM) and PD 325901 (0.25 -0.75μM), from 6 – 96 hpf, shows increasing rescue of the ectopic pigment cells. Scale bar = 100 μm (A-G).

S5 Figure In-silico translation and structural prediction for the pde alleles. Scheme shows 2D structure of the human ET$\alpha$ receptor, with identical amino acids of the zebrafish EdnrAa receptor shown in black for the WT allele (A), pde$^{j262}$ (B), pde$^{v212}$ (C) and pde$^{hu4140}$ (D). In the mutant alleles, changed coding regions are indicated as follows: Amino acids changed due to a shift in the reading frame are shown in dark grey and amino acids that are absent since lie after new stop codon (red arrowheads) are shown in light grey.

S6 Figure Expression of the Tg(mitfa:gfp) line in the ventral trunk of WT larvae. (A) Scheme shows 8 dpf fish, with the red box indicating the area where mitfa:gfp positive cells in the ventral trunk were found. (B) GFP$^+$ cells are readily found in the vicinity of the dorsal aorta throughout the posterior trunk and anterior tail at 8 dpf; superimposed DIC image shows these cells are not melanised. (C) Quantitation of GFP$^+$ cells from a random posterior trunk segment in each of 5 fish, given as mean±s.d.=2.3±0.44 (n=5).
Ednraa/Pde → Factor X

WT

NT

No

BV

1)

2)

3)

pde

Blood vessel (BV) or DRG expressing quiescence Factors X

Quiescent aPSC

Activated aPSC

Iridophore

Melanocyte
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Supporting Information
S4 Fig.tif