Response to comment on ‘Palovarotene reduces heterotopic ossification in juvenile FOP mice but exhibits pronounced skeletal toxicity’

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Abstract We respond to concerns expressed by Pacifici and Shore (2019) about a recent paper (Lees-Shepard and Goldhamer, 2018a) in which we reported that the drug palovarotene can have severe side effects in a mouse model of fibrodysplasia ossificans progressiva. DOI: https://doi.org/10.7554/eLife.43928.001

Introduction

We recently published a study that demonstrated the toxic effects of palovarotene on the skeletons of juvenile mice (Lees-Shepard et al., 2018a). Pacifici and Shore have raised concerns about this study (Pacifici and Shore, 2019): while they do not challenge the veracity of the results or the rigor with which the experiments were performed, they question the relevance of our study to the ongoing efficacy and safety study of palovarotene for the treatment of fibrodysplasia ossificans progressiva (FOP) being conducted by Clementia Pharmaceuticals (https://clinicaltrials.gov/ct2/show/NCT03312634). Here we respond to their concerns and provide context for the experimental design of our study (which was supported by a sponsored research agreement with Clementia Pharmaceuticals).

Discussion

Route of palovarotene administration

Pacifici and Shore strongly imply that the goal of our study was to inform the development of acceptable safety parameters for the use of palovarotene in juvenile FOP patients, and commented that the method we used for drug delivery (intraperitoneal injection: IP) is never used in patients. It will be clear to readers, however, that the comment in the introduction of our paper concerning an "acceptable safety profile" was a general comment relevant to all potential therapies for FOP because of the prevalence of heterotopic ossification (HO) in juvenile patients. A comprehensive analysis of palovarotene toxicity was neither the stated goal nor the intent of our publication.

IP administration of palovarotene was considered necessary for our study to avoid the complication of oral dosing of animals with jaw HO, which is highly penetrant in our model, and to eliminate the real risk of inducing HO in throat and jaw muscles by daily manipulation and probable irritation. No published studies have directly compared palovarotene $C_{\text{max}}$ values for IP and oral dosing, and Pacifici and Shore provide no evidence that cell and tissue levels of palovarotene are affected by route of administration. Further, regardless of possible differences in $C_{\text{max}}$, it is not at all clear that $C_{\text{max}}$ is the most relevant pharmacokinetic parameter.
FOP mouse model

Pacifici and Shore also criticize our use of a Pdgfrα-Cre driver to conditionally recombine the FOP allele, Acvr1\textsuperscript{tnR206H}, stating that our mouse model "does not mimic FOP patients". While it is true that Pdgfrα-Cre;Acvr1\textsuperscript{tnR206H/+} mice differ from FOP in humans in that the mutation in this mouse model is restricted to cells that express, or have expressed, Pdgfrα, this restriction was an essential feature of our study. The study had two primary objectives: i) to examine the effect of palovarotene on HO driven by fibro/adipogenic progenitors; ii) to assess the efficacy of palovarotene in inhibiting body-wide spontaneous HO. As Pdgfrα is the best known single marker for fibro/adipogenic progenitors (reviewed in Lees-Shepard and Goldhamer, 2018), and because Pdgfrα-Cre;Acvr1\textsuperscript{tnR206H/+} mice recapitulate the major pathogenetic manifestations of FOP, including progressive HO at all major anatomical sites affected in FOP patients (Lees-Shepard \textit{et al.}, 2018a; Lees-Shepard \textit{et al.}, 2018b), the Pdgfrα-Cre driver was ideally suited for our studies. We also note that no additional cell types need to be invoked to explain the full repertoire of HO in FOP (Lees-Shepard \textit{et al.}, 2018b).

In 2016, Pacifici, Shore and colleagues used juvenile Prrx-Cre;Acvr1\textsuperscript{R206H} mice to test the safety and efficacy of palovarotene (Chakkalakal \textit{et al.}, 2016). Prrx-Cre is broadly expressed in limb mesenchyme during development, and it is reasonable to assume that the large majority of cells in the limbs of these mice had undergone Cre-recombination at the Acvr1\textsuperscript{R206H} locus. However, this similarity in genetic make-up does not necessarily mean that Prrx-Cre;Acvr1\textsuperscript{R206H} mice are a better model of FOP in humans, as Pacifici and Shore contend. Indeed, while FOP patients exhibit only minor developmental abnormalities (most notably, great toe malformations), mice carrying the Acvr1\textsuperscript{R206H} mutation in all cells exhibit severe skeletal developmental defects and die perinatally (Chakkalakal \textit{et al.}, 2012; Kaplan \textit{et al.}, 2012). Although the reasons for this dramatic phenotypic difference between mice and humans are unknown, it is clear that genetic status of the Acvr1 gene is only one of potentially many model parameters to consider, depending on study goals.

Effects on growth plates and joints

Our publication documented severe skeletotoxic effects of palovarotene on both wild-type and Pdgfrα-Cre;Acvr1\textsuperscript{tnR206H/+} mice, including loss of tibial growth plates (Lees-Shepard \textit{et al.}, 2018a). In contrast, Chakkalakal \textit{et al.} (2016) showed that palovarotene treatment actually improved growth plate morphology in Prrx-Cre;Acvr1\textsuperscript{R206H} mice and had only slight effects on skeletal tissues of wild-type controls. They attributed the improvement in Prrx-Cre;Acvr1\textsuperscript{R206H} mice to the offsetting actions of increased BMP signaling resulting from Acvr1\textsuperscript{R206H} expression in growth plate cartilage and the inhibitory effects of palovarotene on BMP signaling. Pacifici and Shore argue that the deleterious effects of palovarotene on growth plates of Pdgfrα-Cre;Acvr1\textsuperscript{tnR206H/+} mice resulted from the lack of Acvr1\textsuperscript{R206H} expression. At present, this hypothesis remains speculative. Pacifici and Shore assume that the growth plates in our model do not express Pdgfrα (reviewed in Schatteman \textit{et al.}, 1992), so it would not be surprising if the Acvr1\textsuperscript{tnR206H} allele was recombined in the growth plates of Pdgfrα-Cre;Acvr1\textsuperscript{tnR206H/+} mice. In fact, the entire premise of using Prrx-Cre mice in their study is that broad, antecedent expression of Cre in embryonic limb mesenchyme should result in juvenile limbs in which most or all cells carry the Acvr1\textsuperscript{R206H} mutation. Neither we nor Chakkalakal \textit{et al.} (2016) assessed Acvr1\textsuperscript{R206H} expression in growth plate cartilage, and this would certainly be a worthwhile area of investigation.

Another issue, not discussed by Pacifici and Shore, is that Chakkalakal \textit{et al.} (2016) used a distinct dosing regimen (which does not resemble that used in the FOP clinical trial) and earlier study endpoint. Specifically, Prrx-Cre;Acvr1\textsuperscript{R206H} mice were dosed for the first 15 days of life by daily administration of palovarotene to lactating mothers, followed by direct administration by oral gavage from P16 to P30, using an every-other-day dosing schedule and without adjusting for body weight. To begin to address the reasons for the markedly different effects of palovarotene reported by Chakkalakal \textit{et al.} (2016) and Lees-Shepard \textit{et al.} (2018a), we conducted an additional experiment.
in which juvenile mice were treated with palovarotene by IP administration (as in Lees-Shepard et al., 2018a), but with the dosing regimen and 30 day endpoint used by Chakkalakal et al. (2016). Notably, we did not observe loss of growth plates in either wild-type (n = 4) or Pdgfra-Cre;Acvr1TnR206H/+ mice (n = 4) (unpublished observations), consistent with the results of Chakkalakal et al. (2016). These results suggest that dosing regimen and/or study endpoint may explain the disparate study outcomes. As such, attributing the differences to the route of administration or the FOP mouse model used is, at best, premature.

**Conclusion**

The criticisms of our eLife publication by Pacifici and Shore are related to its relevance to ongoing clinical programs. However, these criticisms are based on a series of assumptions that have not been experimentally demonstrated. All mouse studies should be interpreted cautiously, and translation of individual studies to humans should not be assumed or implied. Our work should not be interpreted to mean that palovarotene will necessarily result in skeletal toxicity in children with FOP. Similarly, we urge caution when extrapolating the safety and efficacy data of Chakkalakal et al. (2016) to the ongoing clinical study of palovarotene for the treatment of FOP in children.

**Additional information**

**Competing interests**

John B Lees-Shepard: is currently a scientist at Regeneron Pharmaceuticals, who are currently conducting trials for a drug treatment for FOP. The work reported in Lees-Shepard et al. (2018a), as well as the preliminary data noted above, was conducted entirely at the University of Connecticut and was completed more than a year before he accepted his position at Regeneron. The other author declares that no competing interests exist.

**Author contributions**

David J Goldhamer, Conceptualization, Formal analysis, Supervision, Methodology, Writing—original draft, Project administration, Writing—review and editing; John B Lees-Shepard, Conceptualization, Formal analysis, Methodology, Writing—review and editing

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**Ethics**

Animal experimentation: This study was performed in accordance with the approved institutional animal care and use committee protocol, #A17-015, of the University of Connecticut.

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