



# **EVIDENCE FOR THE INVOLVEMENT OF PPARs IN THE PHOTOPROTECTIVE ACTIVITY OF BIO201 (NORBIXIN) ON RETINAL PIGMENT EPITHELIUM.**

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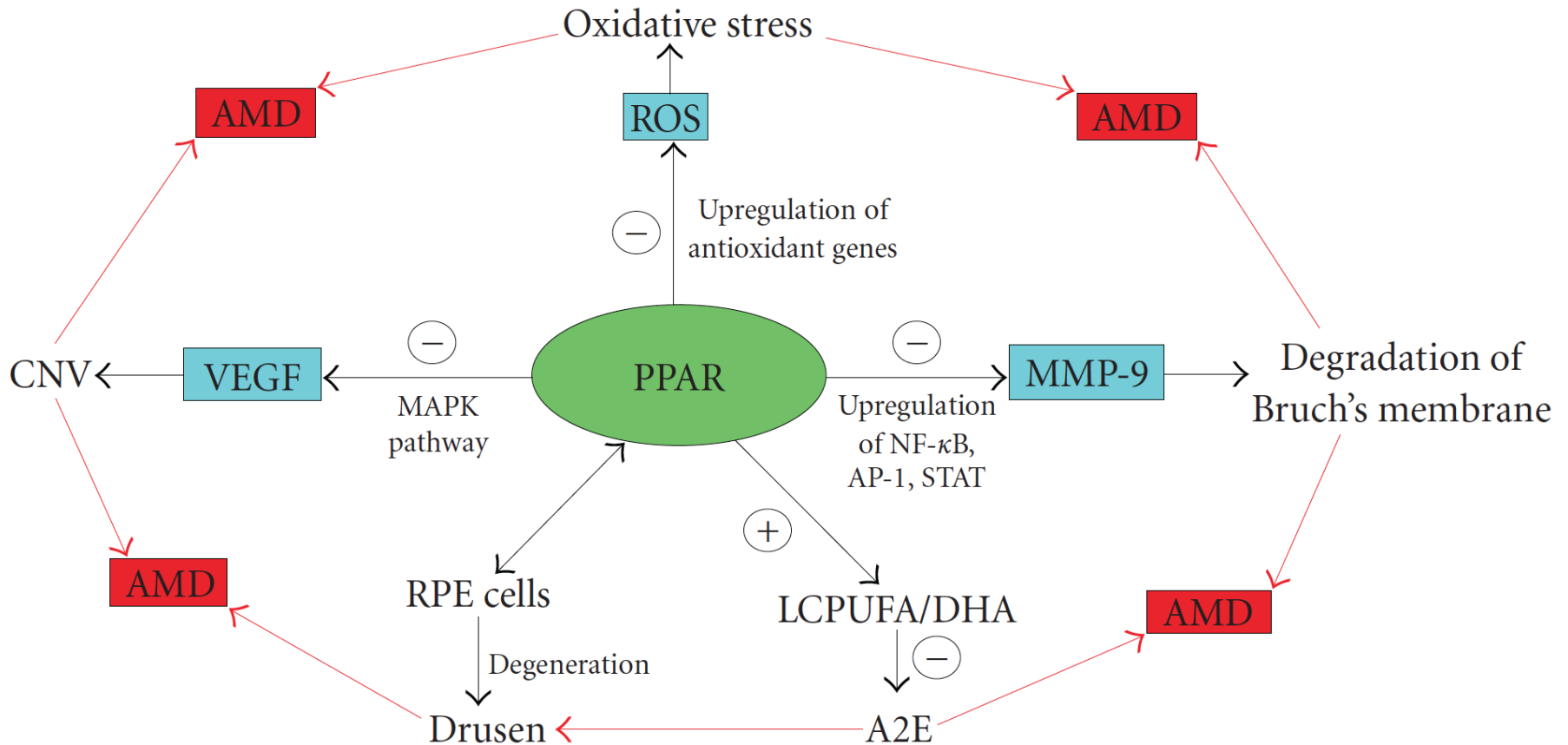
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# INTRODUCTION

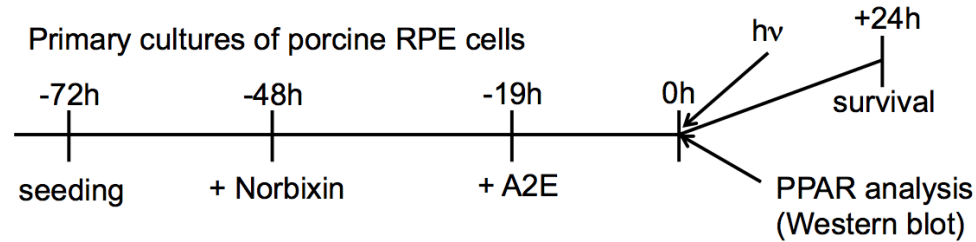
According to Herzlich (PPAR Res. 2008 Article ID 389507 ), PPARs play a key role in the protection of RPE cells against Age-Related Macular Degeneration



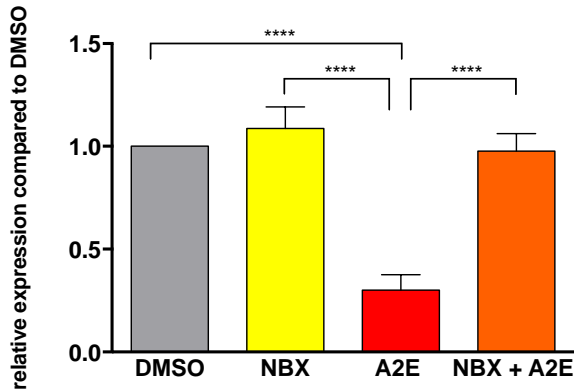
PPARs comprise three subtypes that can accommodate various types of ligands. The aim of the present work was to investigate the respective roles of PPAR $\alpha$ , PPAR $\beta/\delta$  and PPAR $\gamma$  in response to BIO201 (norbixin) on the protection against the deleterious effects of A2E.

A2E sensitizes RPE cells to blue light damage and reduces PPAR $\alpha$  protein levels. Both effects are blunted by BIO201, a preparation of the diapo-carotenoid norbixin developed by Biophytis<sup>1</sup>.

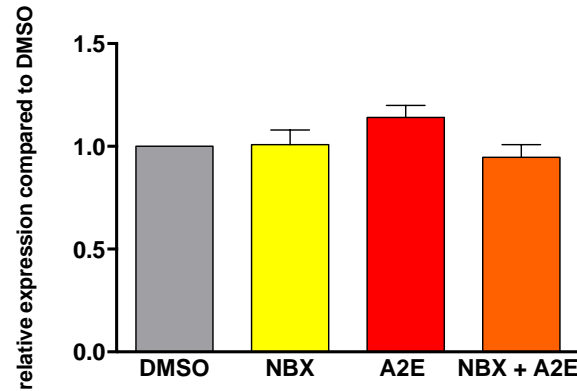
No significant effect is observed on other PPARs levels.



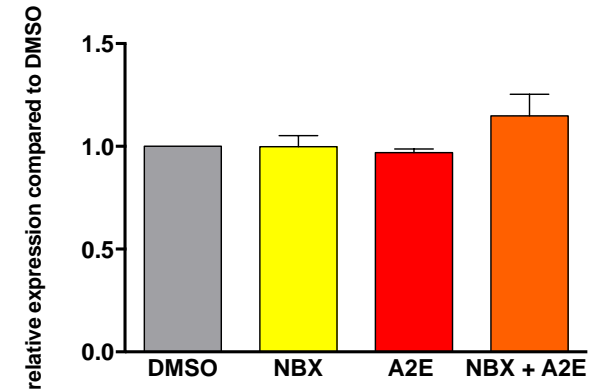
### PPAR $\alpha$



### PPAR $\beta/\delta$



### PPAR $\gamma$



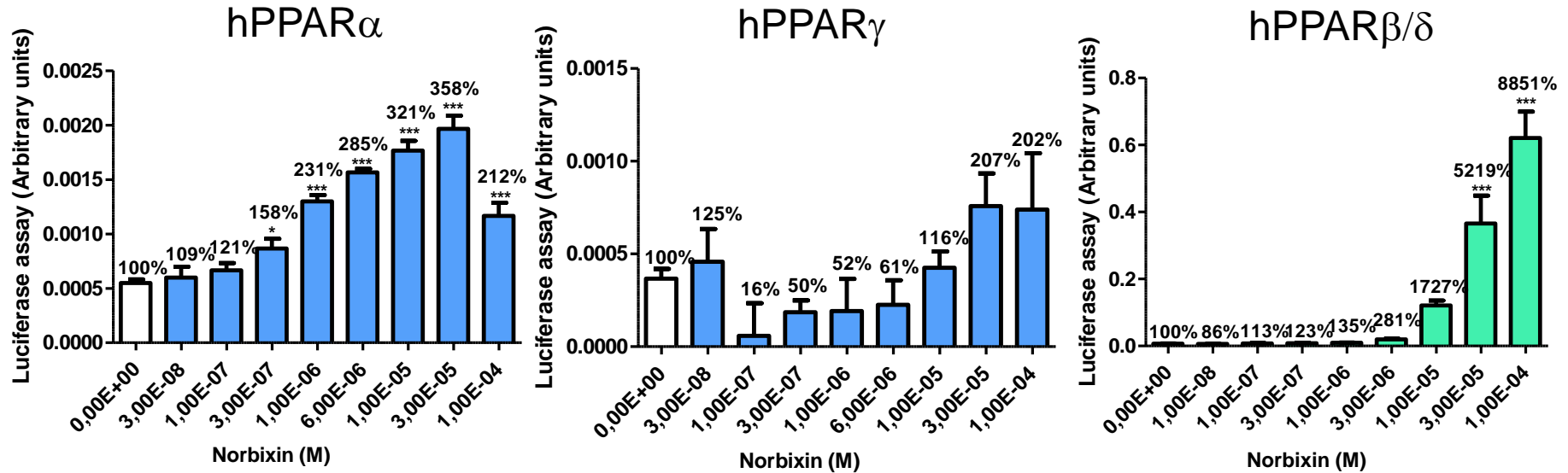
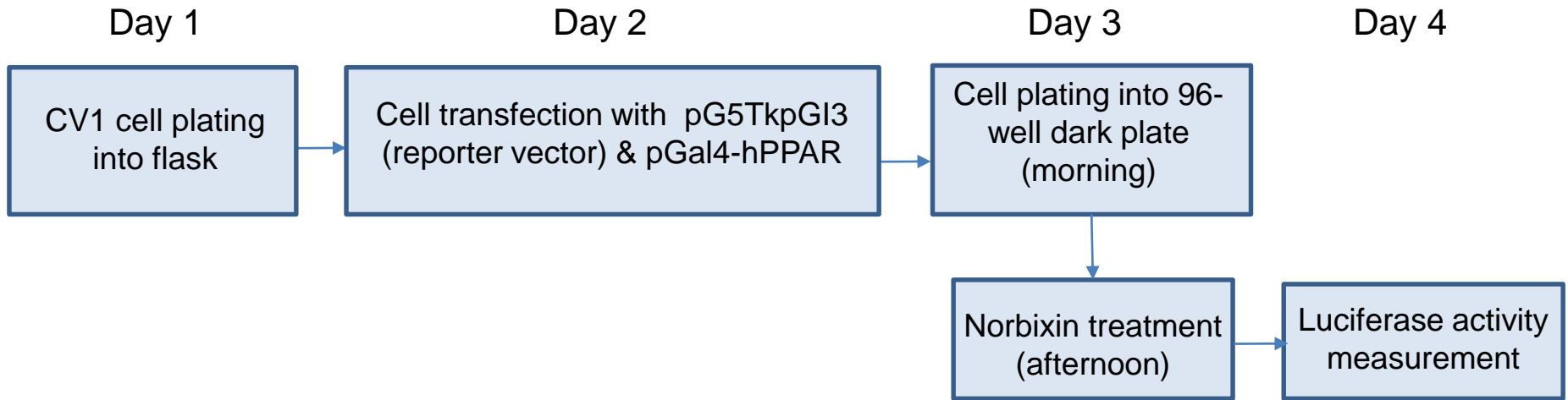
NBX: norbixin 20  $\mu$ M

A2E: 30  $\mu$ M

The effect on PPAR $\alpha$  is post-translational and is also observed with inverted protocols (where treatment with A2E precedes that with norbixin)

<sup>1</sup>(Fontaine et al., PLoS ONE 2016, DOI:10.1371/journal.pone.0167793).

# BIO201 (norbixin) activates hPPAR $\alpha$ and even more efficiently hPPAR $\beta/\delta$



Such experiments do not tell us whether these effects are direct or indirect

## **DIRECT BINDING TESTS** (competition with radioactive ligands):

BIO201 (norbixin) binds to PPAR $\alpha$  with a K<sub>i</sub> of 16.5  $\mu$ M, and to PPAR $\gamma$  with a K<sub>i</sub> of 1.15  $\mu$ M

## **FRET ACTIVITY TESTS** (induction of PPAR coactivator or corepressor recruitment):

BIO201 (norbixin) shows no agonist effect on any of the three PPARs, but has an antagonist effect on PPAR $\beta/\delta$  (K<sub>a</sub> ca. 3.2  $\mu$ M)

### **Effect of various pharmacological agents**

<b>Compound</b>	<b>Effect</b>
PPAR $\alpha$ agonist	<b>Sulindac</b> does not protect RPE cells and does not maintain PPAR $\alpha$ levels
PPAR $\alpha$ antagonist	<b>MK886</b> protects RPE cells and maintains PPAR $\alpha$ levels
PPAR $\beta/\delta$ agonist	<b>GW0742</b> protects RPE cells and maintains PPAR $\alpha$ levels
PPAR $\beta/\delta$ antagonist	<b>GSK3787</b> reduces protection by norbixin
PPAR $\gamma$ agonist	<b>Troglitazone</b> protects RPE cells and maintains PPAR $\alpha$ levels
PPAR $\gamma$ antagonist	<b>T0070907</b> reduces protection by norbixin and troglitazone

**APOPTOSIS:** BIO201 (norbixin) cytoprotective activity correlates with a significant reduction of both ROS levels (p<0.01) and apoptosis induction upon treatment with antimycin A.

# CONCLUSIONS

- These experiments have revealed the difficulty of interpreting experimental data, arising from the limited specificity of current PPAR agonists and antagonists, a potential source of multiple cross-reactions (e.g. troglitazone).
- There is some discrepancy between activation studies and binding experiments, which suggests that PPAR activation is mostly indirect.
- A2E induces PPAR $\alpha$  degradation, and all the tested compounds showing a protection of RPE cells against blue light damage in the presence of A2E are able to sustain PPAR $\alpha$  levels. Thus the presence of PPAR $\alpha$  appears required for photoprotection.
- Agonists of PPAR $\gamma$  and PPAR $\beta/\delta$  protect RPE cells, whereas their antagonists reduce the protective effect of BIO201 (norbixin). Thus, both PPARs seem involved in BIO201 action.
- Additional strategies (siRNA, shRNA, ...) are under way in order to relieve remaining ambiguities and assess definitively the role of the three PPARs in BIO201 (norbixin) protective action.
- **Norbixin** is the active principle of **BIO201**, a 97% pure 9'-cis isomer prepared from the seeds of *Bixa orellana*. BIO201 has been formulated for improving its stability and *per os* bioavailability as **Macuneos**. Biophytis will start a phase 1 clinical trial in 2017 with Macuneos as an oral treatment for dry AMD.