CHRONIC ORAL TREATMENT WITH BIO201 PRESERVES RETINAL FUNCTION IN A DRY AMD EXPERIMENTAL MODEL

Valérie FONTAINE1, Elodie MONTEIRO1, Elena BRAZHKIKOVA1, Mylène FOURNIE1, Christine BALDUCCI2, Louis GUIBOUT2, José-Alain SAHEL1, Stanislas VEILLET2, Pierre DILDA2, René LAFONT2

1 Sorbonne Université, INSERM U968, CNRS UMR 7210 - Institut de la Vision, 75012 Paris, France
2 Biophytis, Université Pierre et Marie Curie, 4 place Jussieu, 75005 Paris, France

Contact: valerie.fontaine@inserm.fr, pierre.dilda@biophytis.com

PURPOSE
We have recently demonstrated the potential of BIO201, the di-apo-carotenoid norborn, entering clinical development, on retinal pigmented epithelium (RPE) and retina photoprotection in vitro and in vivo (Fontaine et al., 2016). We also showed that BIO201 is a dual PPARα/PPARγ ligand (Fontaine et al. 2017).

Here we demonstrate that chronic BIO201 oral administration to a mouse model of dry AMD can slow down or stop the pathological course of the disease when used as preventive or curative treatment.

METHODS
Experiments were performed using the Abca4+/Rdh8- transgenic model of dry AMD. Young mice (6-week-old males; n=10) were used to test BIO201 as a preventive treatment and older mice (12-month-old males, n=7 and females, n=10) to test it as a curative treatment. Mice were fed daily with either BIO201 supplemented (0.5 g/kg) food or regular regimen for 5 or 6 months, respectively.

Full field scotopic and photopic electroretinograms (ERG) were measured at the end of the treatment. One eye was then collected for assessing A2E accumulation, and the second was used to perform histological analyses. Photoreceptor nuclei quantification was performed on eye cryosections. Oral bioavailability of BIO201 and A2E concentrations were measured in all mice by HPLC-MS/MS.

Statistical analyses were performed using either a t test or a one-way ANOVA followed by Dunnett’s test.

CONCLUSIONS
Oral administration of BIO201 (ca. 50 mg.kg1.d-1) provides a significant protection to the retina both at histological and functional levels, provided that the retinal degeneration is not too advanced when starting the treatment.

These results support the clinical development of Macuneos, a preparation based on BIO201 for dry AMD prevention and treatment.

REFERENCES
Fontaine et al. (2016) Norborn protects retinal pigmented epithelium cells and photoreceptors against A2E-mediated phototoxicity in vitro and in vivo. PLOS ONE (DOI:10.1371/journal.pone.0157793)
Fontaine et al. (2017) Involvement of peroxisome proliferator activator receptors (PPARs) in the photoprotective activity of BIO201. Ever 2017, poster #F050

RESULTS

**PREVENTIVE STUDY**

BIO201 protects the function of Abca4+/Rdh8- mouse photoreceptors

In the preventive experiment BIO201 significantly maintained the scotopic ERG A-wave at flash intensities of 1 and 10 cd.s/m². At 10 cd.s/m² A-wave of the BIO201 treated mice was still 82% of the 6W control vs 62% for the vehicle treated mice. The photopic ERG was also better in BIO201 treated mice.

**CURATIVE STUDY**

BIO201 protects the function of Abca4+/Rdh8- mouse photoreceptors

In the curative experiment 3-month-treatment of BIO201 was not highly efficient on the scotopic ERG either in females or in males but showed a slight not significant effect on photopic ERG.

Interestingly, after 5 months of treatment BIO201 completely preserved the scotopic ERG A-wave in 17-month-old females (p<0.01) but not in males. The photopic ERG was slightly higher in BIO201 treated mice (not significant).

The eyes of 12-month-old males contain more A2E and show a worsened scotopic ERG as compared to females

Males show a reduced BIO201 bioavailability as compared to females

The sex differences appear to be related to the mouse model itself rather than to the molecule. Old male mice showed an accelerated kinetics of age-related A2E accumulation in the eyes and a more deteriorated ERG at the onset of treatment. At 1 year of age, A2E level was higher in males compared to females (230.4 ± 15.5 vs 192.8 ± 6.3 pmol/eye; p<0.05) whereas ERG a-wave was lower (-101.8 ± 5.1 vs -119.5 ± 4.6 µV; p<0.05). Males also showed a reduced BIO201 bioavailability as compared to females.

CONCLUSIONS

BIO201 treatment has no impact on photoreceptor degeneration in young mice

BIO201 treatment has no impact on photoreceptor degeneration in old females

BIO201 protects residual photoreceptors in old females

BIO201 does not reduce A2E accumulation in old females

BIO201 reduces A2E accumulation

A2E accumulation was reduced by 22% (p<0.01) in BIO201 treated mice as compared to vehicle treated mice.

No difference in A2E accumulation was observed in 17-month-old mice.

The eyes of 12-month-old males contain more A2E and show a worsened scotopic ERG as compared to females

Males show a reduced BIO201 bioavailability as compared to females

The sex differences appear to be related to the mouse model itself rather than to the molecule. Old male mice showed an accelerated kinetics of age-related A2E accumulation in the eyes and a more deteriorated ERG at the onset of treatment. At 1 year of age, A2E level was higher in males compared to females (230.4 ± 15.5 vs 192.8 ± 6.3 pmol/eye; p<0.05) whereas ERG a-wave was lower (-101.8 ± 5.1 vs -119.5 ± 4.6 µV; p<0.05). Males also showed a reduced BIO201 bioavailability as compared to females.

REFERENCES
Fontaine et al. (2016) Norborn protects retinal pigmented epithelium cells and photoreceptors against A2E-mediated phototoxicity in vitro and in vivo. PLOS ONE (DOI:10.1371/journal.pone.0157793)
Fontaine et al. (2017) Involvement of peroxisome proliferator activator receptors (PPARs) in the photoprotective activity of BIO201. Ever 2017, poster #F050

DIAGNOSIS
VF, P; DM (F); SB (F); M (F); CB (D); LG (B); ARD (S); P (S); E (F); PD (S); BL (S); E (P)

Amplitude (µV)

A2E amount (pmol/eye)

Intensity (log cd.s/m²)

Intensity (log cd.s/m²)

Amplitude (µV)

Amplitude (µV)

Intensity (log cd.s/m²)

Intensity (log cd.s/m²)

Amplitude (µV)

Amplitude (µV)

-2 -1 0 1 1,5
-2 -1 0 1 1,5
-2 -1 0 1 1,5
-2 -1 0 1 1,5
-2 -1 0 1 1,5
-2 -1 0 1 1,5
-2 -1 0 1 1,5
-2 -1 0 1 1,5
-2 -1 0 1 1,5
-2 -1 0 1 1,5
-2 -1 0 1 1,5
-2 -1 0 1 1,5
-2 -1 0 1 1,5
-2 -1 0 1 1,5
-2 -1 0 1 1,5
-2 -1 0 1 1,5
-2 -1 0 1 1,5
-2 -1 0 1 1,5
-2 -1 0 1 1,5
-2 -1 0 1 1,5
-2 -1 0 1 1,5
-2 -1 0 1 1,5
-2 -1 0 1 1,5
-2 -1 0 1 1,5
-2 -1 0 1 1,5
-2 -1 0 1 1,5
-2 -1 0 1 1,5
-2 -1 0 1 1,5
-2 -1 0 1 1,5