



Efficacy and safety outcomes from the FIREFLEYE next study of children 3 years of age with retinopathy of prematurity treated with intravitreal aflibercept versus laser in the randomized FIREFLEYE study

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Disclosures

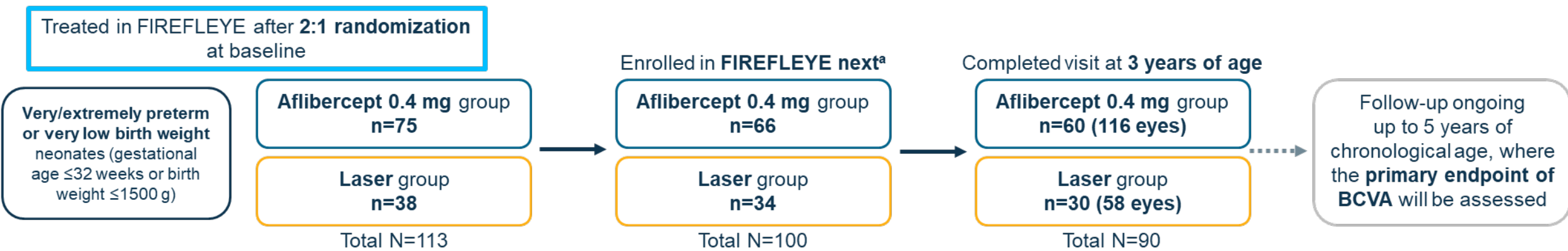
- **Domenico Lepore:** Consultant for Bayer and Novartis
 - **AS:** Speaker for Allergan, Bayer, Novartis, and Roche; attended advisory boards: Apellis, Bayer, Novartis, and Roche; research: Bayer and Novartis; contributed to clinical trials: Bayer and Novartis; board of directors: SemaThera Inc. **HN, NA,** and **CJ:** Received honoraria from Bayer. **W-CW:** Consultant for Allergan, Bayer, Novartis, and Roche. **AA, RV,** and **KC:** Employees of Regeneron Pharmaceuticals Inc. **SS, EK,** and **KB:** Employees of Bayer AG. **PI:** Former employee of Bayer AG. **FZ:** Employee of Bayer Inc. **SL:** Employee of Bayer Consumer Care AG. **TM:** Employee of Bayer U.S. LLC. **AF:** Consultant for Bayer and Novartis.
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FIREFLEYE next (NCT04015180) study design

FIREFLEYE next is the first multinational, ongoing, Phase 3b study assessing ocular and further clinical outcomes, including growth and neurodevelopmental outcomes, **through 5 years of age** following treatment of acute-phase ROP with intravitreal aflibercept 0.4 mg vs. laser photocoagulation in the **24-week, Phase 3 FIREFLEYE study**¹



Endpoints included in pre-planned interim analysis (3 years of chronological age)

- Absence of **active ROP**
- Absence of **unfavorable structural outcomes**
- **Completion of retinal vascularization**
- **Treatment need** for ROP complications
- **Refractive spherical equivalent**
- **BCVA function** (Snellen equivalent)
- Ability to **fix and follow a 5-cm toy**
- Outcomes of **growth and neurodevelopmental tests** (optional [WPPSI-IV, VABS-II])



^aThree study treatment-unrelated deaths in FIREFLEYE and the parents/carers of 10 children (aflibercept 0.4 mg, 6; laser, 4) did not consent to enrollment in FIREFLEYE next. **BCVA**, best corrected visual acuity; **n**, number; **ROP**, retinopathy of prematurity; **VABS-II**, Vineland Adaptive Behavior Scales, Second Edition; **WPPSI-IV**, Wechsler Preschool and Primary Scale of Intelligence, Fourth Edition. 1. Stahl A, et al. JAMA 2022;328:348–59. 3

Baseline characteristics

	Aflibercept 0.4 mg (n=66)	Laser (n=34)	Total (N=100)			
Male, %	54.5	50.0	53.0			
Gestational age, w and d	26w 4d ±2.1	26w 0d ±1.7	26w 3d ±1.9			
Gestational age group, %						
<24 weeks	4.5	8.8	6.0			
≥24 to <27 weeks	59.1	64.7	61.0			
≥27 weeks	36.4	26.5	33.0			
Birth weight, g	882.2±286.9	819.5±238.6	860.9±271.9			
Body weight at baseline treatment in FIREFLEYE, g	2045.8±675.8	1843.8±569.2	1977.1±645.8			
Chronological age at FIREFLEYE next entry, months	9.0±1.6	9.1±1.7	9.0±1.6			
	At FIREFLEYE study entry			At FIREFLEYE next study entry		
Detailed ROP classification by investigator, %	Aflibercept 0.4 mg (n=75)	Laser (n=38)	Total (N=113)	Aflibercept 0.4 mg (n=66)	Laser (n=34)	Total (N=100)
Absence of ROP	0	0	0	83.3	85.3	84.0
Zone I (excluding AP-ROP)	20.0	18.4	19.5	4.5	8.8	6.0
Stage 1	1.3	0	0.9	1.5	2.9	2.0
Stage 2	2.7	5.3	3.5	0	0	0
Stage 3	4.0	2.6	3.5	0	0	0
Stage 3+	12.0	10.5	11.5	0	0	0
Stage 4A	0	0	0	1.5	0	1.0
Stage 4B	0	0	0	1.5	2.9	2.0
Missing	0	0	0	0	2.9	1.0
Zone II (excluding AP-ROP)	61.3	68.4	63.7	6.1	5.9	6.0
Stage 1	0	0	0	1.5	2.9	2.0
Stage 2	0	2.6	0.9	3.0	0	2.0
Stage 2+	9.3	13.2	10.6	0	0	0
Stage 3+	52.0	52.6	52.2	0	0	0
Missing	0	0	0	1.5	2.9	2.0
Zone III (excluding AP-ROP)	0	0	0	6.1	0	4.0
Stage 1	0	0	0	4.5	0	3.0
Missing	0	0	0	1.5	0	1.0
AP-ROP	18.7	13.2	16.8	0	0	0
Zone I	16.0	10.5	14.2	0	0	0
Zone II	2.7	2.6	2.7	0	0	0



84% of children had no ROP at FIREFLEYE next study entry



Data are mean ±SD unless stated otherwise. **AP-ROP**, aggressive-posterior retinopathy of prematurity; **d**, day; **w**, week.

Unfavorable structural outcomes, ROP recurrence, and treatment for ROP complications in FIREFLEYE next

Unfavorable structural outcomes	Aflibercept 0.4 mg (n=66)		Laser (n=34)	
	At any time until 2 years of chronological age	At any time until 3 years of chronological age	At any time until 2 years of chronological age	At any time until 3 years of chronological age
Number of children, %	100.0	100.0	100.0	100.0
None	93.9	93.9	94.1	94.1
Retinal detachment	4.5	6.1	2.9	2.9
Macular dragging	1.5	1.5	2.9	2.9
Macular fold	1.5	1.5	0	0
Retrolental opacity	1.5	1.5	0	0
Any unfavorable structural outcome	6.1	6.1	5.9	5.9
Number of treated eyes, %	100.0	100.0	100.0	100.0
None	94.5	94.5	95.3	95.3
Retinal detachment	3.9	4.7	1.6	1.6
Macular dragging	1.6	1.6	3.1	3.1
Macular fold	1.6	1.6	0	0
Retrolental opacity	1.6	1.6	0	0
Any unfavorable structural outcome	5.5	5.5	4.7	4.7
Recurrence of ROP after entry into FIREFLEYE next^a	Between entry and 2 years of age	Between entry and 3 years of age	Between entry and 2 years of age	Between entry and 3 years of age
n	64	60	32	30
Recurrence, %	1.5	1.7	0	0

- **No disease reactivation occurred after 50 weeks of chronological age**
- In total, **4 patients were treated** after entry into FIREFLEYE next for ROP complications, all before 1 year of age (including 2 patients with pre-existing bilateral retinal detachment, 1 with reactivated plus disease^b, and 1 with retinal neovascularization not further specified^c)
 - 1 patient showed retinal detachment at age 3 (in the progression of macular fold reported at ages 1 and 2 years)

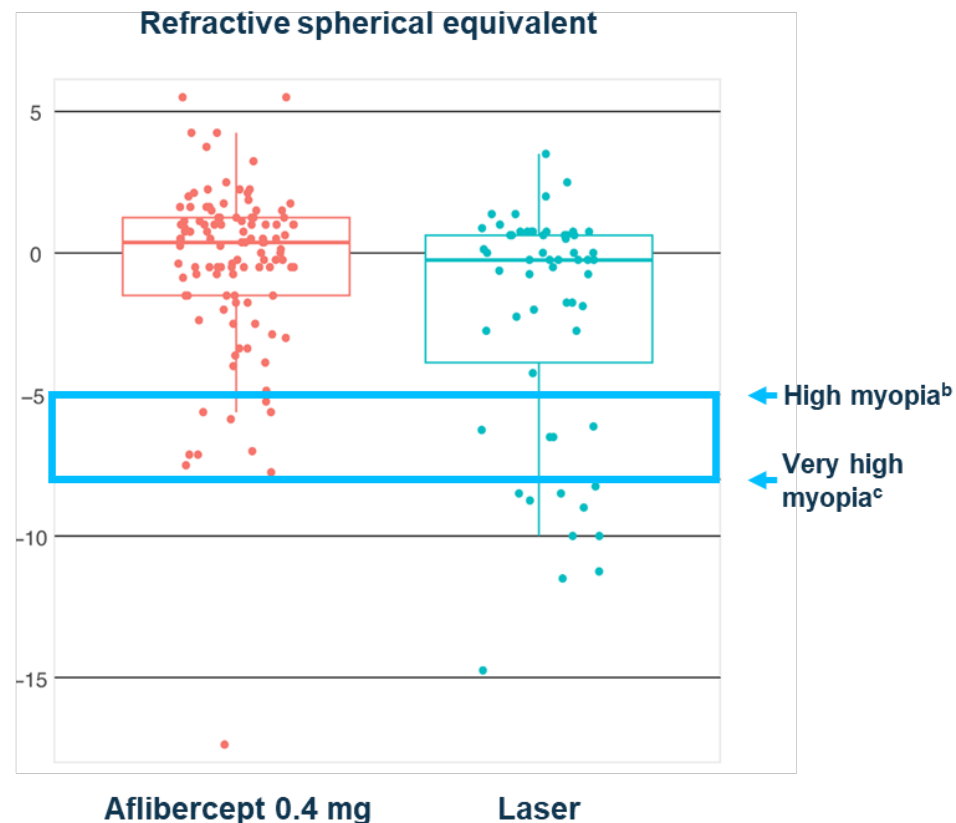
^aPost-hoc analysis. A child was considered as having ROP recurrence if: the inclusion criteria of FIREFLEYE (or worse) were reported and a previous assessment (either in FIREFLEYE or in FIREFLEYE next) of ROP not requiring treatment (according to the inclusion criteria) was available. ^bZone I, both eyes, treated at around 43 weeks of age. ^cTreated around 50 weeks of chronological age.



Ophthalmic outcomes at 3 years of age

No./Total No. (%)	Aflibercept 0.4 mg	Laser
Ocular findings, eyes		
Nystagmus	4/116 (3.4)	2/58 (3.4)
Absence of manifest strabismus	98/116 (84.5)	50/58 (86.2)
Amblyopia	1/116 (0.9)	2/58 (3.4)
Cataract	1/116 (0.9)	0
Optic nerve atrophy	2/116 (1.7)	0
Ability to fix and follow a 5-cm toy	112/116 (96.6)	57/58 (98.3)
Eyes with complete retinal vascularization^a		
At 1 year of chronological age	89/128 (69.5)	-
At 2 years of chronological age	97/121 (80.2)	-
At 3 years of chronological age	89/111 (80.2)	-
BCVA (Snellen equivalent score), patients		
≥20/200	44/45 (97.8)	23/23 (100)
≥20/40	30/45 (66.7)	11/23 (47.8)

Retinal vascularization after aflibercept treatment appeared to be complete in 80% of eyes by 2 years of age



Fix and follow a 5 cm toy

Aflibercept 0.4 mg, **97%**; laser, **98%**

BCVA ≥20/40

Aflibercept 0.4 mg, **67%**; laser, **48%**

Myopia

Mild in both groups and **less pronounced with aflibercept**

Aflibercept 0.4 mg, **-0.4 (3.1) D**;
Laser **-2.2 (4.2) D**

High myopia

Aflibercept 0.4 mg **10 eyes (9%)**;
Laser **14 eyes (24%)**

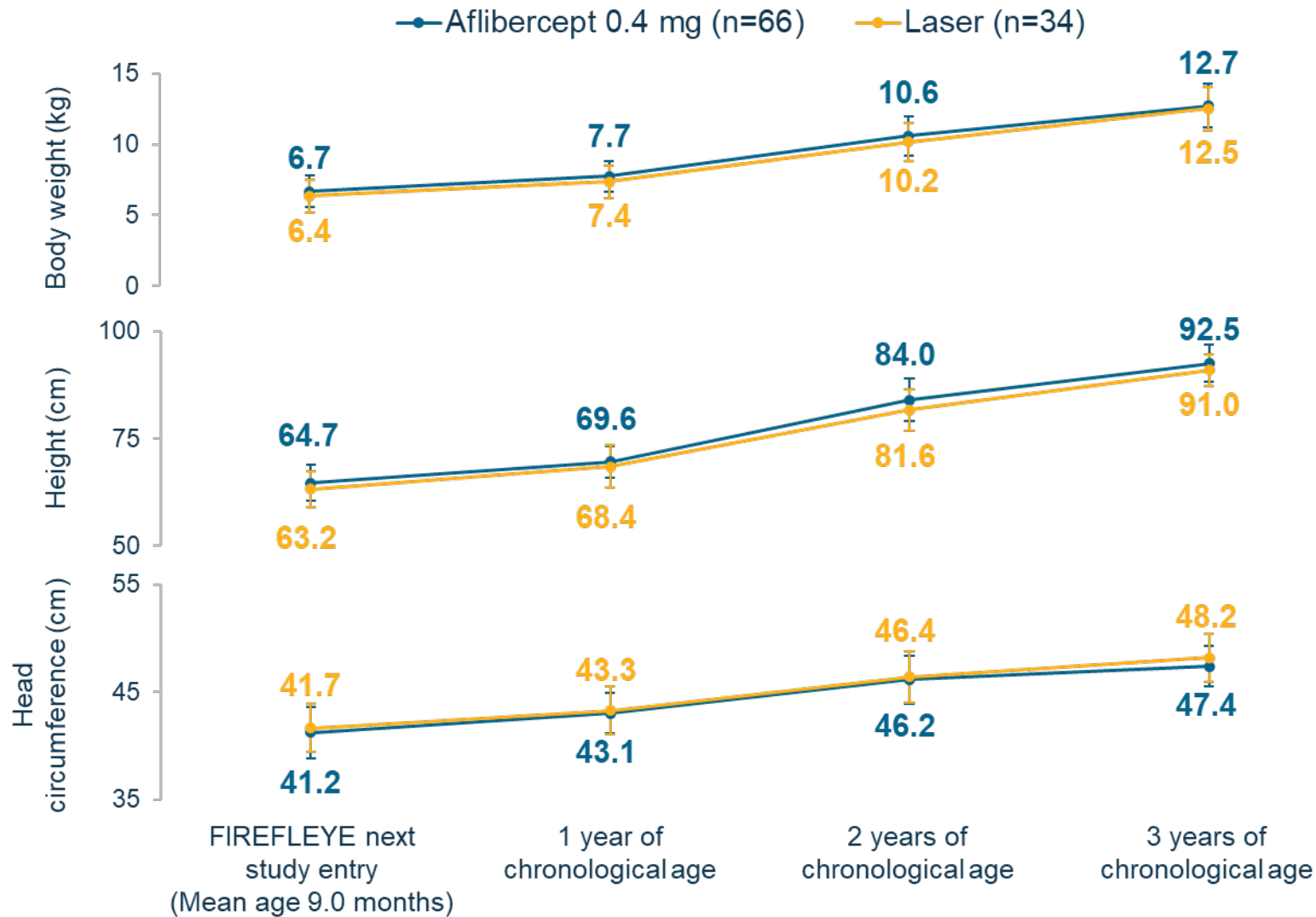
Very high myopia

Aflibercept 0.4 mg **1 eye (1%)**;
Laser **10 eyes (17%)**





Growth parameters up to 3 years of age



No clinically relevant differences in growth parameters were observed between both groups through 3 years of chronological age, and **results are in line with what is expected in this prematurely born pediatric population**





Adverse events (during FIREFLEYE next through 3 years of age)

	Aflibercept 0.4 mg (n=66)	Laser (n=34)
Any AE, n (%)	59 (89.4)	29 (85.3)
Ocular AEs	33 (50.0)	11 (32.4)
Ocular AEs in eyes formerly treated in FIREFLEYE	32 (48.5)	10 (29.4)
Systemic AEs	56 (84.8)	29 (85.3)
AEs related to aflibercept 0.4 mg	2 (3.0)	1 (2.9)
AEs related to laser treatment	3 (4.5)	6 (17.6)
Ocular AEs in eyes formerly treated in FIREFLEYE (≥10% occurrence in any group), n (%)		
Astigmatism	13 (19.7)	5 (14.7)
Myopia	9 (13.6)	5 (14.7)
Strabismus ^a	9 (13.6)	2 (5.9)
Any SAE, n (%)	21 (31.8)	14 (41.2)
Ocular SAEs in eyes formerly treated in FIREFLEYE	6 (9.1)	0
Systemic SAEs	19 (28.8)	14 (41.2)
SAEs related to aflibercept 0.4 mg	1 (1.5)	0
Death	0	0
Ocular SAEs, n (%)	6 (9.1)	0
Optic atrophy	2 (3.0)	0
Retinal detachment	2 (3.0)	0
Retinal neovascularization	2 (3.0)	0
Retinopathy of prematurity	1 (1.5)	0
Vitreous opacities	1 (1.5)	0
Retinoblastoma	1 (1.5)	0
Systemic SAEs (≥5% occurrence in any arm), n (%)		
Cerebral palsy	2 (3.0)	4 (11.8)
Bronchiolitis	2 (3.0)	2 (5.9)
Bronchospasm	0	2 (5.9)



Ocular and systemic AEs were consistent with those expected in children born preterm and who developed severe ROP, and no new safety concerns were identified

^aReported as an adverse event. No imbalance of “clinically manifest strabismus” (reported as an efficacy parameter) between groups.
AE, adverse event; SAE, serious adverse event.





Conclusions

FIREFLEYE next is the **first prospective, controlled, Phase 3b study** evaluating **long-term efficacy and safety outcomes** after treatment of **acute-phase ROP with aflibercept 0.4 mg versus laser photocoagulation** (final results through 5 years of age expected for 2026)

Through 3 years of chronological age, **efficacy outcomes were well sustained, and no ocular or systemic safety concerns, including outcomes of growth, were identified:**

Efficacy

- **Disease reactivation** after aflibercept 0.4 mg **was rare**
 - No disease reactivation occurred after 50 weeks of chronological age. One patient showed retinal detachment at age 3 years in the progression of macular fold reported at ages 1 and 2 years of chronological age
- No patient received treatment in the post-acute phase later than 50 weeks of chronological age
- **Retinal vascularization** after aflibercept 0.4 mg **appeared to be complete in 80% of eyes by 2 years** of chronological age
- **Visual function was age-appropriate**, and **myopia** was rarer and **less severe in the aflibercept 0.4 mg group** than the laser group

Safety

- **No ROP treatment-specific effects** on **growth** outcomes through 3 years of age
- No ocular or systemic safety concerns through 3 years of age were identified



Overall, aflibercept 0.4 mg injection therapy in very/extremely preterm or very low birthweight patients with acute-phase ROP (as approved^a) was **effective and generally well tolerated through 3 years of age**

^aAflibercept has been approved for treatment of ROP in Japan (September 2022),¹ the European Union (December 2022),² Switzerland,² Great Britain,² the USA (February 2023),³ and Brazil (April 2023).⁴

1. Bayer, 2023. Available at: <https://www.bayer.com/media/en-us/eylea-approved-in-japan-for-treatment-of-preterm-infants-with-retinopathy-of-prematurity/> [Accessed July 2024]; 2. Bayer AG, 2023. Available at: https://www.ema.europa.eu/en/documents/product-information/eylea-epar-product-information_en.pdf [Accessed July 2024]; 3. Regeneron Pharmaceuticals, 2023. Available at: https://www.regeneron.com/downloads/eylea_fpi.pdf [Accessed July 2024]; 4. Anvisa NHTA, 2023. Available at: <https://www.gov.br/anvisa/pt-br/assuntos/medicamentos/novos-medicamentos-e-indicacoes/eylea-aflibercepte-nova-indicacao> [Accessed July 2024].





Thank you to all FIREFLEYE next patients, parents, caregivers, and investigators

Final results through 5 years of age are expected in 2026

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