



Two-year outcomes from FIREFLEYE Next, a prospective follow-up study to evaluate long-term efficacy and safety of patients treated with intravitreal aflibercept or laser photocoagulation for ROP in the FIREFLEYE study

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Disclosures

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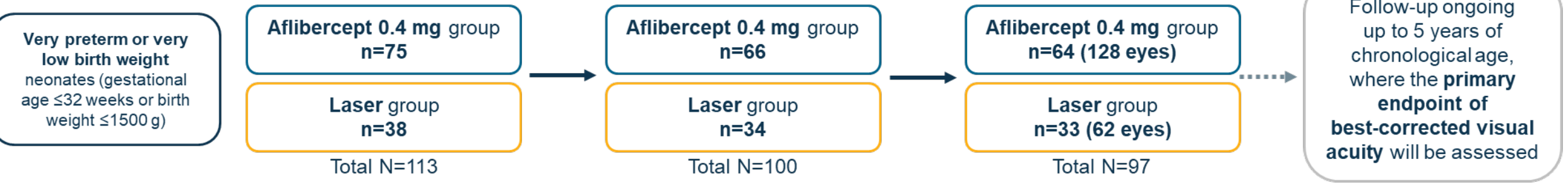
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FIREFLEYE Next (NCT04015180) study design

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

Treated in FIREFLEYE after **2:1 randomization** at baseline



Endpoints included in exploratory interim analysis (2 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**
- Ability to **fix and follow a 5-cm toy**
- Outcomes of **growth**, and **neurodevelopmental tests** (BSID-III, 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. **BSID-III**, Bayley Scales of Toddler and Infant Development, Third Edition; **ROP**, retinopathy of prematurity; **VABS-II**, Vineland Adaptive Behavior Scales, Second Edition. 1. Stahl et al. *JAMA* 2022;328(4):348–59.



Baseline characteristics

	Aflibercept 0.4 mg (n=66)	Laser (n=34)	Total (N=100)
Male, n (%)	36 (54.5)	17 (50.0)	53 (53.0)
Gestational age, w and d	26w 4d ±2.1	26w 0d ±1.7	26w 3d ±1.9
Gestational age group, n (%)			
<24 weeks	3 (4.5)	3 (8.8)	6 (6.0)
≥24 to <27 weeks	39 (59.1)	22 (64.7)	61 (61.0)
≥27 weeks	24 (36.4)	9 (26.5)	33 (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



84% of children had **no ROP** at FIREFLEYE Next study entry

Detailed ROP classification by investigator, n (%)	At FIREFLEYE study entry			At FIREFLEYE Next study entry		
	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	55 (83.3)	29 (85.3)	84 (84.0)
Zone I (excluding AP-ROP)	15 (20.0)	7 (18.4)	22 (19.5)	3 (4.5)	3 (8.8)	6 (6.0)
Stage 1	1 (1.3)	0	1 (0.9)	1 (1.5)	1 (2.9)	2 (2.0)
Stage 2	2 (2.7)	2 (5.3)	4 (3.5)	0	0	0
Stage 3	3 (4.5)	1 (2.6)	4 (3.5)	0	0	0
Stage 3+	9 (12.0)	4 (10.5)	13 (11.5)	0	0	0
Stage 4A	0	0	0	1 (1.5)	0	1 (1.0)
Stage 4B	0	0	0	1 (1.5)	1 (2.9)	2 (2.0)
Missing	0	0	0	0	1 (2.9)	1 (1.0)
Zone II (excluding AP-ROP)	46 (61.3)	26 (68.4)	72 (63.7)	4 (6.1)	2 (5.9)	6 (6.0)
Stage 1	0	0	0	1 (1.5)	1 (2.9)	2 (2.0)
Stage 2	0	1 (2.6)	1 (0.9)	2 (3.0)	0	2 (2.0)
Stage 2+	7 (9.3)	4 (13.2)	12 (10.6)	0	0	0
Stage 3+	39 (52.0)	20 (52.6)	59 (52.2)	0	0	0
Missing	0	0	0	1 (1.5)	1 (2.9)	2 (2.0)
Zone III (excluding AP-ROP)	0	0	0	4 (6.1)	0	4 (4.0)
Stage 1	0	0	0	3 (4.5)	0	3 (3.0)
Missing	0	0	0	1 (1.5)	0	1 (1.0)
AP-ROP	14 (18.7)	5 (13.2)	19 (16.8)	0	0	0
Zone I	12 (16.0)	4 (10.5)	16 (14.2)	0	0	0
Zone II	2 (2.7)	1 (2.6)	3 (2.7)	0	0	0

Data are mean ±SD unless stated otherwise. **AP-ROP**, aggressive-posterior retinopathy of prematurity; **n**, number; **SD**, standard deviation.

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 1 year of chronological age	At any time until 2 years of chronological age	At any time until 1 year of chronological age	At any time until 2 years of chronological age
Number of children, n (%)	66 (100.0)	66 (100.0)	34 (100.0)	34 (100.0)
None	62 (93.9)	62 (93.9)	32 (94.1)	32 (94.1)
Retinal detachment	3 (4.5)	3 (4.5)	1 (2.9)	1 (2.9)
Macular dragging	1 (1.5)	1 (1.5)	1 (2.9)	1 (2.9)
Macular fold	1 (1.5)	1 (1.5)	0	0
Retrolental opacity	1 (1.5)	1 (1.5)	0	0
Any unfavorable structural outcome	4 (6.1)	4 (6.1)	2 (5.9)	2 (5.9)
Number of treated eyes, n (%)	128 (100.0)	128 (100.0)	64 (100.0)	64 (100.0)
None	121 (94.5)	121 (94.5)	61 (95.3)	61 (95.3)
Retinal detachment	5 (3.9)	5 (3.9)	1 (1.6)	1 (1.6)
Macular dragging	2 (1.6)	2 (1.6)	2 (3.1)	2 (3.1)
Macular fold	2 (1.6)	2 (1.6)	0	0
Retrolental opacity	2 (1.6)	2 (1.6)	0	0
Any unfavorable structural outcome	7 (5.5)	7 (5.5)	3 (4.7)	3 (4.7)
Recurrence of ROP after entry into FIREFLEYE Next^a	Between entry and 1 year of age	Between entry and 2 years of age	Between entry and 1 year of age	Between entry and 2 years of age
n	66	64	34	32
Recurrence, n (%)	1 (1.5)	0	0	0

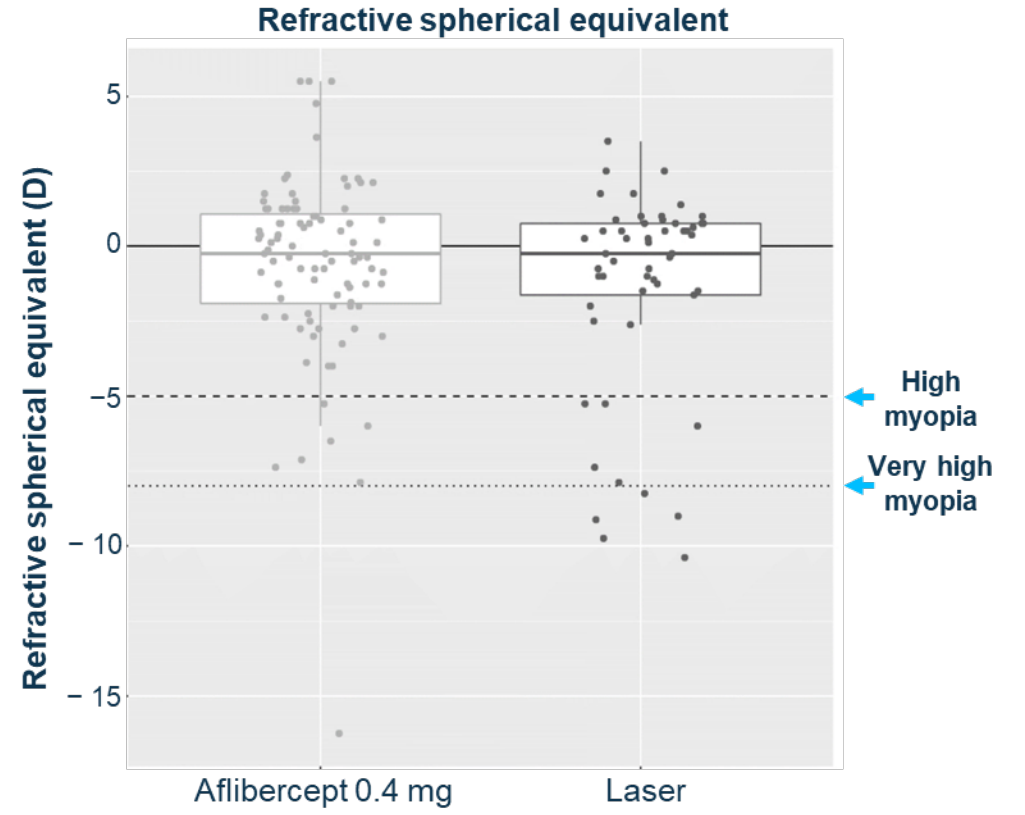
In total, **4 patients were treated** after entry into FIREFLEYE Next for ROP complications, all before 1 year of age (including 2 patients with pre-existing bilateral retinal detachment, 1 each with reactivated plus disease^b and retinal neovascularization not further specified^c)

^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 2 years of age

	Aflibercept 0.4 mg (122 eyes)	Laser (63 eyes)
Ocular findings at 2 years of chronological age, n (%)		
Nystagmus	6 (4.9)	4 (6.3)
Manifest strabismus	28 (23.0)	12 (19.0)
Amblyopia	0	0
Cataract	1 (0.8)	0
Optic nerve atrophy	2 (1.6)	0
Ability to fix and follow a 5-cm toy	118 (96.7)	62 (98.4)
Eyes with complete retinal vascularization, n (%)^a		
At 1 year of chronological age	91 (71.1)	-
At 2 years of chronological age	97 (80.2)	-



Retinal vascularization in the aflibercept 0.4 mg group continued beyond 1 year of age

Most eyes (aflibercept 0.4 mg, **97%**; laser, **98%**) were able to fix and follow a 5-cm toy

Myopia was mild in both groups and **less pronounced in the aflibercept 0.4 mg group** (aflibercept 0.4 mg, $-0.6 [3.1]$ D; laser, $-1.4 [3.4]$ D).

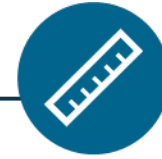
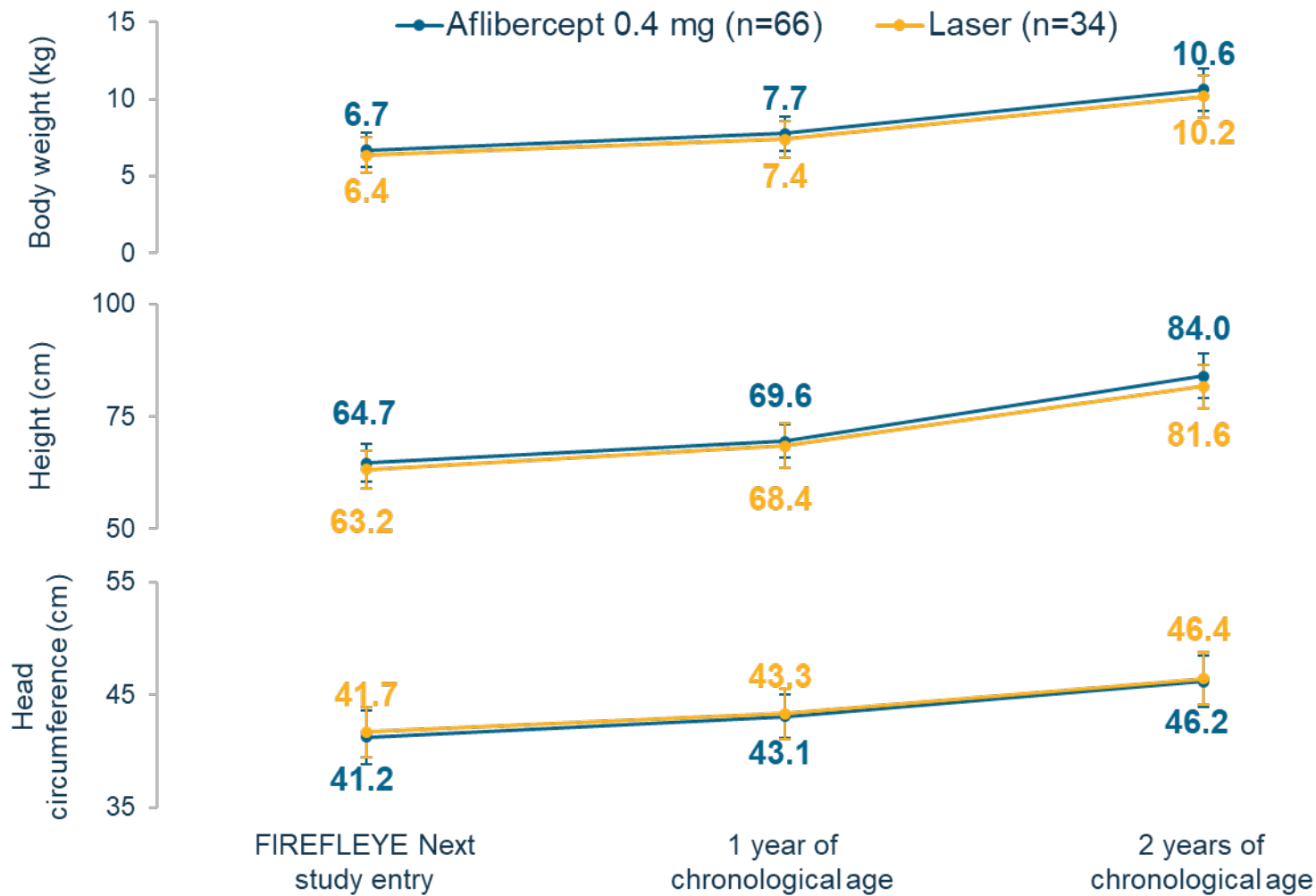
High myopia was present in **7 eyes (8%)** in the aflibercept 0.4 mg group and **10 eyes (19%)** in the laser group

Very high myopia was present in **1 eye (1%)** in the aflibercept 0.4 mg group and **5 eyes (9%)** in the laser group

^aAnalysis of complete vascularization in eyes receiving laser therapy is not reported, as laser scars prevent physiologic vascularization of the peripheral retina. Square brackets indicate standard deviation. D, diopter.



Growth parameters up to 2 years of age

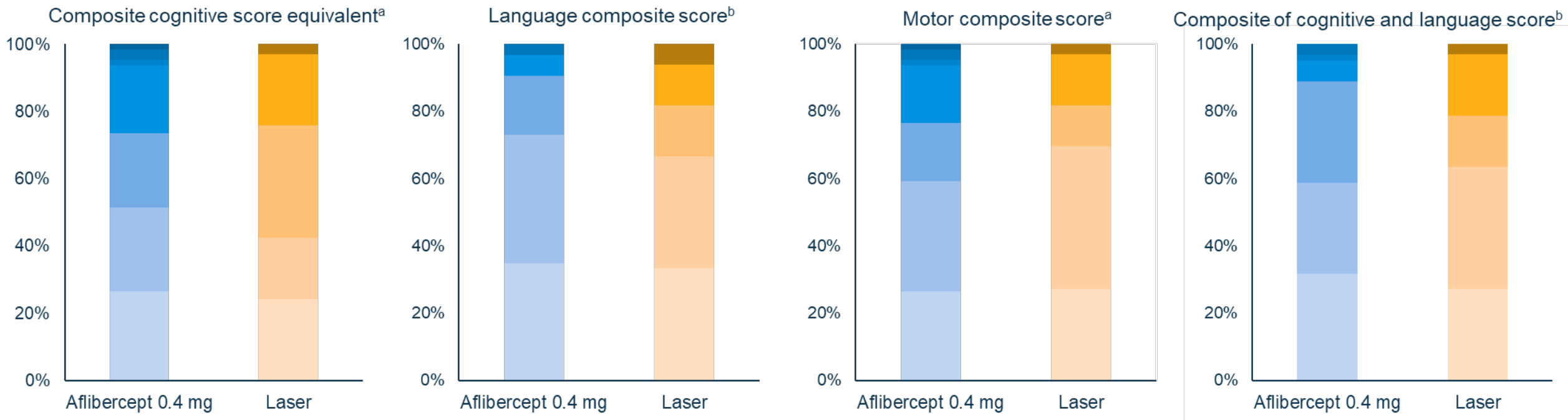


No clinically relevant differences in growth parameters were observed between both groups at 1 and 2 years of age, and results are in line with **what is expected in this prematurely born pediatric population**



Neurodevelopmental test outcomes at 2 years of age

Bayley Scales of Toddler and Infant Development, 3rd Edition (BSID-III) scores



Composite scores for BSID-III and VABS-II (not shown) at 2 years of age were comparable across both groups. Small numerical differences in both directions can likely be attributed to sample size

^aAflibercept 0.4 mg, n=64; Laser, n=34. ^bAflibercept 0.4 mg, n=63; Laser, n=33.



Adverse events (during FIREFLEYE Next through 2 years of age)

	Aflibercept 0.4 mg (n=66)	Laser (n=34)
Any AE, n (%)	53 (80.3)	27 (79.4)
Ocular AEs	30 (45.5)	9 (26.5)
Ocular AEs in eyes formerly treated in FIREFLEYE	29 (43.9)	9 (26.5)
Systemic AEs	49 (74.2)	24 (70.6)
AEs related to aflibercept 0.4 mg	1 (1.5)	1 (2.9)
AEs related to laser treatment	1 (1.5)	5 (14.7)
Ocular AEs in eyes formerly treated in FIREFLEYE (≥5% occurrence in any group), n (%)		
Astigmatism	9 (13.6)	4 (11.8)
Myopia	9 (13.6)	5 (14.7)
Strabismus ^a	9 (13.6)	2 (5.9)
Conjunctivitis	1 (1.5)	2 (5.9)
Any SAE, n (%)	17 (25.8)	10 (29.4)
Ocular SAEs in eyes formerly treated in FIREFLEYE	6 (9.1)	0
Systemic SAEs	15 (22.7)	10 (29.4)
SAEs related to aflibercept 0.4 mg	1 (1.5)	0
Death	0	0
Ocular SAEs, n (%)		
Optic atrophy	1 (1.5)	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)	2 (5.9)
Bronchospasm	0	2 (5.9)



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

^aReported as an adverse event. No imbalance of “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 vs. laser photocoagulation** (final results through 5 years of age expected for 2026)

Through 2 years of chronological age, **efficacy outcomes were well sustained**, and **no ocular or systemic safety concerns**, including outcomes of growth and neurodevelopmental testing, **were identified**:

Efficacy

- **No late retinal detachment, and disease reactivation** after aflibercept 0.4 mg **was rare**
- No patient received treatment in the post-acute phase later than 50 weeks of chronological age
- **Retinal vascularization** after aflibercept 0.4 mg **continued beyond 1 year** 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 2 years of age and **neurodevelopmental test** performance at 2 years of age
 - No ocular or systemic safety concerns through 2 years of age were identified

 Overall, aflibercept 0.4 mg injection therapy in very preterm/very low birthweight patients with acute-phase ROP (as approved^a) was **effective and generally well-tolerated through 2 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 2023]; 2. Bayer AG, 2023. Available at: https://www.ema.europa.eu/en/documents/product-information/eylea-epar-product-information_en.pdf [Accessed July 2023]; 3. Regeneron Pharmaceuticals, 2023. Available at: https://www.regeneron.com/downloads/eylea_fpi.pdf [Accessed July 2023]; 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 2023].



Thank you to all FIREFLEYE Next patients and investigators

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