

MEDICAL POLICY

POLICY TITLE	SUPRACHOROIDAL DELIVERY OF PHARMACOLOGIC AGENTS
POLICY NUMBER	MP- 4.032

Effective Date:	1/1/2024
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[POLICY RATIONALE](#)
[DISCLAIMER](#)
[POLICY HISTORY](#)

[PRODUCT VARIATIONS](#)
[DEFINITIONS](#)
[CODING INFORMATION](#)

[DESCRIPTION/BACKGROUND](#)
[BENEFIT VARIATIONS](#)
[REFERENCES](#)

I. POLICY

Suprachoroidal delivery of a pharmacologic agent is considered **investigational**, as there is insufficient evidence to support a conclusion concerning the health outcomes or benefits associated with this procedure.

Cross-references:

- MP 2.028** Eye Care
- MP 2.149** Aqueous Shunts and Stents for Glaucoma
- MP 2.159** Intravitreal Corticosteroid Implants
- MP 4.008** Photodynamic Therapy for Choroidal Neovascularization

II. PRODUCT VARIATIONS

[TOP](#)

This policy is only applicable to certain programs and products administered by Capital Blue Cross please see additional information below, and subject to benefit variations as discussed in Section VI below.

FEP PPO - Refer to FEP Medical Policy Manual. The FEP Medical Policy manual can be found at: <https://www.fepblue.org/benefit-plans/medical-policies-and-utilization-management-guidelines/medical-policies>.

III. DESCRIPTION/BACKGROUND

[Top](#)

Delivery of pharmacologic agents to the suprachoroidal space is being investigated for treatment of posterior eye segment diseases.

The structure of the eye is classified under two subheadings: 1) anterior segment, and 2) posterior segment. The anterior segment consists of the front one-third of the eye that includes; pupil, cornea, iris, ciliary body, aqueous humor, and lens; the posterior segment consists of the back two-thirds of the eye that includes vitreous humor, retina, choroid, macula, and optic nerve. Posterior segment ocular diseases (e.g., age-related macular degeneration, diabetic neuropathy) are the most prevalent causes of visual impairment. The following is a list of the various routes for ocular drug administration:

Invasive drug administration to intraocular cavities

MEDICAL POLICY

POLICY TITLE	SUPRACHOROIDAL DELIVERY OF PHARMACOLOGIC AGENTS
POLICY NUMBER	MP- 4.032

- Suprachoroidal injections
- Intravitreal surgery
- Intravitreal injections
- Intracameral surgery
- Subretinal injection
- Intracameral injections

Invasive periocular and scleral modes of drug administration

- Intrascleral surgery
- Episcleral surgery
- Periocular injections
- Subconjunctival injections
- Transscleral diffusion from controlled release systems

Noninvasive methods

- Topical administration on the eye

Systemic administration

- Intravenous infusion and injection
- Oral

Many ocular diseases are treated with either topical or systemic medications. Topical application has remained the most preferred delivery route due to ease of administration. Topical application is useful in the treatment of disorders affecting the anterior segment of the eye. Although topical and systemic routes are convenient, lack of bioavailability and failure to deliver therapeutic levels of drugs to the retina has prompted vision scientists to continue to explore alternative routes of administration.

One potential advantage of suprachoroidal injection would be the ability to minimize systemic adverse effects while delivering higher local tissue levels of drugs. This proposed benefit assumes that high local levels lead to improved outcomes. Weighed against this potential benefit is the risk of localized tissue damage from the microcannula. A microcannula system combines a drug delivery channel with a fiberoptic light source for localization of the cannula tip. This technique is being investigated for the treatment of subchoroidal neovascularization related to diseases of the retina.

Regulatory Status

The iTrack™ (iScience Interventional), which is a flexible microcannula designed to allow atraumatic cannulation of spaces in the eye for infusion and aspiration of fluids during surgery,

MEDICAL POLICY

POLICY TITLE	SUPRACHOROIDAL DELIVERY OF PHARMACOLOGIC AGENTS
POLICY NUMBER	MP- 4.032

received 510(k) marketing clearance from the U.S. Food and Drug Administration (FDA). The microcannula incorporates an optical fiber to allow transmission of light to the microcannula tip for surgical illumination and guidance. The microcannula “is indicated for fluid infusion and aspiration, as well as illumination, during surgery.”

IV. RATIONALE

[Top](#)

Controlled trials are needed to evaluate the safety and efficacy of suprachoroidal drug administration compared to the standard of care. There is insufficient peer-reviewed evidence to establish effectiveness. The evidence is insufficient to determine that the technology results in an improvement in the net health outcome.

V. DEFINITIONS

[Top](#)

ANGIOGENESIS refers to the development of blood vessels.

CHOROID is the thin, highly vascular membrane covering the posterior five sixths of the eye between the retina and the sclera.

CHOROIDAL NEOVASCULARIZATION refers to the abnormal formation of new blood vessels usually on or under the retina, usually seen in diabetic retinopathy, blockages of central retinal vision and macular degeneration.

EXUDATION refers to the pathological oozing of fluids, usually the result of inflammation.

MACULAR DEGENERATION refers to loss of pigmentation in the macular region of the retina, usually affecting persons over age fifty (50); a common disease of unknown etiology that produces central visual field loss and is the leading cause of permanent blindness in the United States.

OCULAR refers to the eye or vision.

PHOTODYNAMIC refers to the effects of light on biological, chemical, or physical systems.

VI. BENEFIT VARIATIONS

[Top](#)

The existence of this medical policy does not mean that this service is a covered benefit under the member's health benefit plan. Benefit determinations should be based in all cases on the applicable health benefit plan language. Medical policies do not constitute a description of benefits. A member's health benefit plan governs which services are covered, which are excluded, which are subject to benefit limits, and which require preauthorization. There are different benefit plan designs in each product administered by Capital Blue Cross. Members and providers should consult the member's health benefit plan for information or contact Capital Blue Cross for benefit information.

MEDICAL POLICY

POLICY TITLE	SUPRACHOROIDAL DELIVERY OF PHARMACOLOGIC AGENTS
POLICY NUMBER	MP- 4.032

VII. DISCLAIMER

[Top](#)

Capital Blue Cross's medical policies are developed to assist in administering a member's benefits, do not constitute medical advice and are subject to change. Treating providers are solely responsible for medical advice and treatment of members. Members should discuss any medical policy related to their coverage or condition with their provider and consult their benefit information to determine if the service is covered. If there is a discrepancy between this medical policy and a member's benefit information, the benefit information will govern. If a provider or a member has a question concerning the application of this medical policy to a specific member's plan of benefits, please contact Capital Blue Cross' Provider Services or Member Services. Capital Blue Cross considers the information contained in this medical policy to be proprietary and it may only be disseminated as permitted by law.

VIII. CODING INFORMATION

[Top](#)

Note: This list of codes may not be all-inclusive, and codes are subject to change at any time. The identification of a code in this section does not denote coverage as coverage is determined by the terms of member benefit information. In addition, not all covered services are eligible for separate reimbursement.

Investigational; therefore, not covered:

Procedure Codes							
67516							

IX. REFERENCES

[Top](#)

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MEDICAL POLICY

POLICY TITLE	SUPRACHOROIDAL DELIVERY OF PHARMACOLOGIC AGENTS
POLICY NUMBER	MP- 4.032

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9. Kim YC, Oh KH, Edelhauser HF, Prausnitz MR. Formulation to target delivery to the ciliary body and choroid via the suprachoroidal space of the eye using microneedles. *Eur J Pharm Biopharm.* 2015;95(Pt B):398-406 PMID 26036448
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11. Hancock SE, Wan CR, et al. Biomechanics of suprachoroidal drug delivery: From benchtop to clinical investigation in ocular therapies. *Expert Opinion on Drug Delivery* 2021;18(6):777-788 PMID 33393391
12. Archived: Blue Cross Blue Shield Association Medical Policy Reference Manual. 9.03.19, Suprachoroidal Delivery of Pharmacologic Agents December 2014.

X. POLICY HISTORY

[Top](#)

MP 4.032	CAC 10/28/12 Adopting BCBSA. <ul style="list-style-type: none"> • New policy • Extracted information regarding Suprachoroidal Delivery of Pharmacologic Agents from MP 4.008 Ocular Therapy. • No change to policy statement, remains investigational. • Codes reviewed 9/19/2012
	CAC 11/26/13 Consensus review. References updated but no changes to the policy statement. Rationale added. FEP variation revised to refer to the FEP medical policy manual.
	CAC 11/25/14 Consensus review. No change to policy statements. References and rationale updated. Coding reviewed and updated new code 11/10/14
	CAC 11/24/15 Consensus review. No change to the policy statement. Reference and rationale update. Coding updated.
	CAC 11/29/16 Consensus review. FEP variation removed due to archiving of their policy. References updated. New code 0465T added to policy with effective date of 1/1/2017. Variation reformatting.
	12/19/17 Consensus review. No change to the policy statement. References reviewed. Coding and configuration reviewed. 2/28/18 Admin coding review, no changes.
	11/1/18 Consensus review. No change to the policy statement. References reviewed.
	10/01/20 Consensus review. No changes to policy statement.
	9/14/20 Consensus review. Policy statement unchanged. References updated. Removed MP4.023 from cross references retired.
	11/19/2021 Consensus Review. Policy statement unchanged. References

MEDICAL POLICY

POLICY TITLE	SUPRACHOROIDAL DELIVERY OF PHARMACOLOGIC AGENTS
POLICY NUMBER	MP- 4.032

	updated.
	10/28/2022 Consensus Review. Policy statement unchanged. References updated.
	12/12/2023 Administrative update. Deleted 0465T and added 67516

[Top](#)

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