

Figure 6. Intravitreous treatment study, showing late-phase fluorescein angiograms at postlaser days 15, 20, and 29 for representative animals in the groups receiving placebo and VEGF Trap treatment. The placebo-treated animal shows grade 4 leakage in most of the 9 treatment areas at all 3 times. The VEGF Trap-treated animal shows grade 4 leakage at all 9 laser sites on postlaser day 15 prior to receiving a single intravitreous injection of VEGF Trap. By postlaser day 20 (5 days following VEGF Trap injection), there are no grade 4 spots. No recurrence of grade 4 leakage is evident at postlaser day 29.

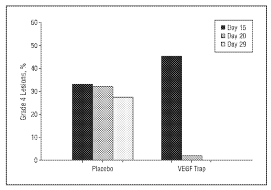


Figure 7. Percentage of grade 4 lesions at postlaser days 15, 20, and 29 for groups that received intravitreous placebo and a single treatment with 500 µg of VEGF Trap given on postlaser day 15.

In this animal model of CNV, VEGF Trap was highly effective at preventing the development of grade 4 leakage on FA regardless of dose or whether it was administered intravenously on a weekly schedule or intravitreously every 2 weeks (Table 3, Figure 4, and Figure 5). Histological assessment confirmed that choroidal new vessel formation, fibrotic changes, and retinal thickness also were markedly less in the treated eyes (Table 5).

Moreover, when a single intravitreous injection of VEGF Trap was given after grade 4 CNV had developed, leakage was stopped within 5 days in approximately 95% of previously active grade 4 lesions and within 14 days

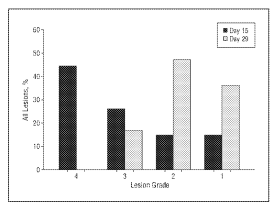


Figure 8. Percentage of all lesion grades at postlaser days 15 and 29 for a single intravitreous treatment with 600 μ g of VEGF Trap given on postlaser day 15.

following treatment in 100% of the lesions (postlaser days 20 and 29, respectively) (Table 3, Figure 6, and Figure 7). Although the effect of a single intravitreous injection of placebo was not evaluated, grade 4 lesions persisted for the duration of the study in all animals receiving multiple intravitreous or intravenous injections of placebo (Table 3). Histological examination revealed a trend toward decreased CNV and fibrosis relative to controls, which was not statistically significant. VEGF is a powerful mediator of vascular permeability in addition to new vessel formation, so VEGF Trap may have blocked VEGF-induced leakage from choroidal neovessels. Alterna-

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		Score,	Mean	
Histological Finding	CNV Pr	CNV Prevention ^a CNV Treatmen		eatment ^b
	Placebo	VEGF Trap	Placebo	VEGF Trap
Fibroplasia	1.74	1 85°	1.88	1.91
Retinal elevation	1.31	0.62°	1.62	1.43
Neovascularization	0.69	0.12°	0.90	0.58
Total	3.74	1.79 ^c	4.40	3.92

Abbreviation: CNV, choroidal neovascularization.

^aMean scores for all lesions in all eyes (n = 6 per group).

^bMean scores for lesions that were grade 4 at postlaser day 15 (prior to the single VEGF Trap injection).

^c P < .05, Mann-Whitney U test.

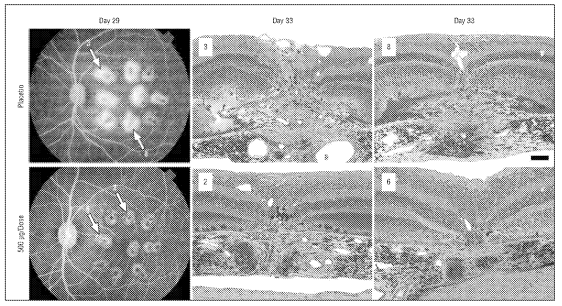


Figure 9. Intravitreous prevention study, showing late-phase fluorescein angiograms at postlaser day 29 and histological sections at postlaser day 33 (glycol methacrylate sections stained with toluicine blue; scale bar=250 µm) for 2 animals that received 3 intravitreous doses of either placebo or VEGF Trap (500 µg/eye/dose). The representative histological sections correspond to the numbered laser treatment areas in the fluorescein angiograms. The placebo-treated sections at bicker and more vascular compared with the VEGF Trap-treated eyes. Note the presence of subretinal fluid in lesion 3 on day 33.

tively, VEGF Trap may have reduced or stopped blood flow through the new vessels.

Intravitreous administration of VEGF Trap was well tolerated, with only a mild inflammatory response noted in the eyes that underwent intravitreous VEGF Trap treatment. Except for 1+ or fewer anterior chamber and vitreous cells in some eyes, no other ophthalmoscopic signs of inflammation were seen.

HUMAN TRIALS OF VEGF TRAP-EYE

VEGF Trap is now in clinical trials (for a recent review, see the article by Dixon et al⁴²). A phase 1 trial of 25 patients with exudative AMD evaluated the tolerability and efficacy of intravenous administration of VEGF Trap at 3 different dose levels. Subjects had a significant decrease in retinal thickness as determined by optical coherence tomography,⁴³ although visual acuity was not significantly improved in this small safety study. However, 1 subject experienced grade 4 hypertension and 1 subject developed grade 2 proteinuria. Hypertension and proteinuria are now well-established class effects of systemic VEGF inhibition, and both patients exhibiting these adverse events in the study by Nguyen et al⁴³ had received the highest intravenous dose of VEGF Trap (3 mg/kg).

Another phase 1 study (Clinical Evaluation of Antiangiogenesis in the Retina, CLEAR-IT 1) used intravitreous administration of VEGF Trap-Eye (aflibercept ophthalmic solution).⁴⁴ The first part of this study was a sequential cohort dose escalation (from 0.05 to 4.0 mg/ eye) in 21 patients with exudative AMD. No serious systemic or ocular toxic effects were observed. However, a marked decrease in retinal thickness⁴⁴ and improvement in visual acuity⁴⁴ were noted. VEGF Trap-Eye also

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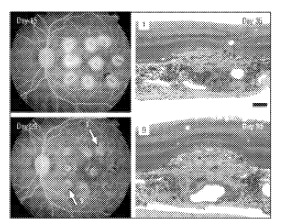


Figure 10. Intravitreous treatment study, showing late-phase fluorescein angiograms at postlaser days 15 and 29 and histological sections (corresponding to the numbered laser treatment areas in the day 29 fluorescein angiogram) obtained at necropsy on postlaser day 35 (glycol methacrylate sections stained with toluidine blue; scale bar=250 µm) from an animal that was treated with a single intravitreous 500-µg dose of VEGF Trap on day 15 following angiography. Note the marked reduction in fluorescein leakage from all of the treatment areas on the day 29 angiogram. In the histological sections, lesions are somewhat thicker and contain more patent choroidal new vessels than was observed in the VEGF Trap prevention study (Table 5 and Figure 9).

has been used in a small open-label safety study for treatment of diabetic macular edema.⁴⁵ A single dose of 4 mg was administered intravitreously to 5 patients who had undergone multiple prior treatments for diabetic macular edema. There was a median decrease in central macular thickness of 79 µm as well as some improvement in vision. A phase 2 trial in diabetic macular edema is in progress.

In a double-masked phase 2 trial (CLEAR-IT 2), VEGF Trap-Eye was evaluated in 157 patients with exudative AMD randomized to either monthly or quarterly intravitreous injections for 12 weeks at doses of 0.5 or 2 mg (monthly injections) and 0.5, 2, or 4 mg (quarterly). Following the 12-week fixed dosing period, patients continued to receive treatments on an as-needed basis at their originally assigned dosages. Reports of the 1-year results described a statistically significant improvement in vision, retinal thickness, and size of the CNV lesions,46,47 with few re-treatments required during the 40week phase of as-needed treatment. Patients initially dosed on a schedule of 2.0 mg monthly received, on average, only 1.6 additional injections during the 40-week period of as-needed treatment, and those initially dosed on a schedule of 0.5 mg monthly received, on average, 2.5 injections. While as-needed dosing following a fixed quarterly dosing regimen (with dosing at baseline and week 12) also yielded improvements in visual acuity at week 52 as compared with baseline, the results generally were not as robust as those obtained with initial fixed monthly dosing. VEGF Trap-Eye was generally well tolerated and there were no drug-related serious adverse events. The most common adverse events were those typically associated with intravitreous injections.

Two phase 3 trials of 2 years' duration are under way to further investigate the efficacy and safety of VEGF Trap-

Eye in wet AMD, VIEW 1 in the United States and Canada⁴⁸ and VIEW 2 in Europe, Japan, and Latin America.⁴⁹ For both trials, VEGF Trap-Eye is being administered intravitreously. In the first year of treatment, VEGF Trap-Eye was administered every 4 weeks at doses of either 0.5 or 2 mg. Another study arm used 3 initial monthly doses of 2 mg followed by 2-mg doses given at 8-week intervals. The active control arm comprised subjects receiving ranibizumab (0.5 mg) at 4-week intervals. The 1-year outcomes from these studies are pending publication.

CONCLUSIONS

Using an established primate model of CNV, administration of VEGF Trap in a prevention protocol markedly reduced vasoproliferative responses of the macaque retina to laser injury, substantially preventing the development of all components of CNV lesions as well as vascular leakage. When a single intravitreous VEGF Trap injection was given after grade 4 lesions had developed, there was resolution of vascular leakage. This also resulted in a trend toward lower histological scores for the neovascular components of the lesions, suggesting partial regression of newly formed vessels.

Submitted for Publication: February 17, 2010; final revision received February 21, 2011; accepted February 24, 2011.

Correspondence: T. Michael Nork, MD, MS, Comparative Ophthalmic Research Laboratories (CORL), 600 Highland Ave, F4/336, Madison, WI 53792-3220 (tmnork @wisc.edu).

Author Contributions: All of the authors had full access to all of the data in the study and take responsibility for the integrity of the data and the accuracy of the data analysis.

Financial Disclosure: Drs Cao, Zimmer, and Wiegand own stock and/or stock options in Regeneron Pharmaceuticals, Inc.

Funding/Support: This study was funded by Regeneron Pharmaceuticals, Inc.

Additional Contributions: Huihao Fan, MA, and Ajit Thakur, PhD, provided the statistical analysis.

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Patent No. 10,828,345 Petition For Post Grant Review

UNITED STATES PATENT AND TRADEMARK OFFICE

BEFORE THE PATENT TRIAL AND APPEAL BOARD

CHENGDU KANGHONG BIOTECHNOLOGY CO., LTD., Petitioner

v.

REGENERON PHARMACEUTICALS, INC., Patent Owner

Patent No. 10,828,345 Issue Date: November 10, 2020 Title: USE OF A VEGF ANTAGONIST TO TREAT ANGIOGENIC EYE DISORDERS

Case: PGR2021-00035

PETITION FOR POST-GRANT REVIEW

UNDER 35 U.S.C. §§ 321-329 AND 37 C.F.R. § 42.200 et seq.

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sf-4338006

Celltrion Exhibit 1014 Page 903

I. INTRODUCTION

Petitioner Chengdu Kanghong Biotechnology Co., Ltd. ("Petitioner" or "Kanghong") respectfully petitions for post grant review ("PGR") in accordance with 35 U.S.C. §§ 321-329 and 37 C.F.R. § 42.200 *et seq.* of all claims of U.S. Patent No. 10,828,345 (the "345 patent" (Ex. 1001)), which issued on November 10, 2020 to Regeneron Pharmaceuticals, Inc. ("Patent Owner" or "Regeneron"). As shown in this petition, all claims are invalid as anticipated, obvious, and lacking written description support.

The '345 patent is premised on the supposed "surprising discovery" that VEGF antagonists are effective with "less frequent dosing . . . compared to prior administration regimens for angiogenic eye disorders which require monthly administrations." Regeneron was not the first to have this revelation. In a PCT application ("Shams")¹ filed more than six years and published almost five years earlier, Genentech described the same discovery: "It has been discovered that the treatment effects of a VEGF antagonist, e.g., Ranibizumab, are maintained for an extended period of time, such as more than one month."²

¹ Ex. 1004.

² Ex. 1004, Page 24, Lines 27-28.

The '345 patent describes methods of treating angiogenic eye disorders with an initial dose, one or more secondary doses, and one or more tertiary doses of a specific VEGF antagonist. Shams describes the same regimen and the VEGF antagonist. The '345 patent's only independent claim limits this dosing regimen by specifying that each secondary dose is administered every 4 weeks and each tertiary dose is administered every 12 weeks. That dosing frequency is also described in Shams. The '345 patent is thus anticipated by Shams.

The '345 patent is also obvious over Regeneron's own prior art press release publicizing the claimed dosing regimens. During prosecution, Regeneron overcame its own press release (the "2009 Press Release") ³ by focusing on the 12– week tertiary dosing frequency as the supposedly inventive aspect of the claims. But the 12–week tertiary dosing frequency was known, as evidenced by Shams. Although the 2009 Press Release expressly disclosed a 12–week tertiary dose, Regeneron argued that "[m]ere mention of a prospective possibility of dosing at 12 weeks does not specifically indicate or teach towards a method where 12–week dosing would be undertaken, let alone successful."⁴ To the extent the 2009 Press release did not adequately disclose a 12–week tertiary dose, the 2009 Press Release

³ Ex. 1005.

⁴ Ex. 1002, Response to Office Action 03/16/2020, Page 4.

in combination with Shams renders the claims obvious. One of skill in the art would have been motivated to combine the Shams 12–week tertiary dosing regimen with the 2009 Press Release because Shams, in listing suitable VEGF antagonists for the regimen, specifically identifies Regeneron's VEGF Trap.⁵ Thus, the '345 patent is invalid as obvious over the 2009 Press Release in view of Shams.

Regeneron's prosecution arguments about the 2009 Press Release also support a finding that the '345 patent is invalid as lacking written description. Regeneron argued during prosecution that one of skill in the art "would not have understood a . . . dosing regimen with 4 and 12 week limits as encompassing a q12w regiment," but the '345 patent specification provides the same disclosure. Regeneron also argued during prosecution that the disclosure of a 12–week dosing "possibility" was insufficient to identify a 12–week dosing, but the '345 patent specification does not differentiate a 12–week tertiary dosing regimen from myriad other possibilities; such "undifferentiated descriptions" of a specific invention are insufficient to satisfy 35 U.S.C. § 112(a). *Novozymes A/S v. DuPont Nutrition Biosciences APS*, 723 F.3d 1336, 1346 (Fed. Cir. 2013).

⁵ Wu Decl. (Ex. 1003) at ¶ 112.

The '345 patent is eligible for PGR. The '345 patent is a "transition" application, filed after the America Invents Act ("AIA") went into effect but claiming priority to several pre-AIA applications. Under the AIA, a patent that issues from a transition application is eligible for PGR if it contains a claim that lacks written description support in a pre-AIA application. Claim 8 of the '345 patent recites "branch retinal vein occlusion," a disorder that was first mentioned in a continuation-in-part (CIP) patent application filed on July 12, 2013. Thus, the '345 patent is eligible for PGR. Further, because the '345 specification does not provide written description support for the claimed 12–week tertiary dosing frequency, none of the pre-AIA priority applications provides written description support for the '345 patent's earliest effective filing date is its actual filing date, October 12, 2018, making the patent eligible for PGR for this separate reason.

The Board should institute PGR and find all of the claims unpatentable under 35 U.S.C. §§ 102, 103, and 112.

II. GROUNDS FOR STANDING

Pursuant to 37 C.F.R. § 42.204(a), Kanghong certifies that the '345 patent is available for PGR and that Kanghong is not barred or estopped from requesting PGR on the grounds identified in this Petition. Specifically: (1) neither Kanghong nor any of its privies own the '345 patent; and (2) neither Kanghong nor any of its

privies have filed a U.S. civil action challenging the validity of any claim of the '345 patent.

Despite claiming priority to applications filed before the effective date of the AIA, the '345 patent is eligible for PGR pursuant to Section 3(n)(1) of the AIA⁶ because multiple granted claims do not find Section 112 support in any of the pre– AIA applications. As explained below, the '345 patent has an earliest effective filing date in July 2013 or October 2018, thus rendering it eligible for PGR.

III. STATEMENT OF RELIEF REQUESTED

Kanghong requests review under 35 U.S.C. § 321 of claims 1-11 of the '345 patent, and seeks a finding that claims 1-11 are unpatentable as anticipated under 35 U.S.C. § 102(a), as obvious under 35 U.S.C. § 103, and for lacking written description support under 35 U.S.C. § 112(a).

IV. THE '345 PATENT

On October 12, 2018, Regeneron filed U.S. Application No. 16/159,282 (the "282 application"), which matured into the '345 patent.⁷ The '345 patent, titled

⁶ Pub. L. No. 112-29, AIA § 3(n)(1), LEAHY-SMITH AMERICA INVENTS ACT,

PL 112-29, 125 Stat. 284, 293 (Sept. 16, 2011).

⁷ Ex. 1001, Cover Page.

"Use of a VEGF Antagonist to Treat Angiogenic Eye Disorders,"⁸ has one independent claim:⁹

A method for treating an angiogenic eye disorder in a patient, said method comprising

sequentially administering to the patient

[a] a single initial dose of a VEGF antagonist,

[b] followed by one or more secondary doses of the VEGF antagonist,

[c] followed by one or more tertiary doses of the VEGF antagonist;

[d] wherein each secondary dose is administered 4 weeks after the immediately preceding dose; and

[e] wherein each tertiary dose is administered 12 weeks after the immediately preceding dose;

[f] wherein the VEGF antagonist is a receptor-based chimeric molecule comprising an immunoglobin-like (lg) domain 2 of a first VEGF receptor which is Fltl and lg domain 3 of a second VEGF receptor which is Flk1, and a multimerizing component.

⁸ Id.

⁹ *Id.* at Col. 21:56-23:14.

Claim 1 has three sequential steps: ¹⁰ (1) administer an initial dose of a specific VEGF antagonist (step [a], narrowed by clause [f]): (2) administer one or more secondary doses, each 4 weeks after the immediately preceding dose (step [b], narrowed by clause [d]); and (3) administer one or more tertiary doses, each 12 weeks after the immediately preceding dose (step [c], narrowed by clause [e]).

The '345 patent includes ten dependent claims. These claims narrow independent claim 1 by specifying the drug administered (claim 2),¹¹ modes of administration (claims 3 and 4),¹² dose amount (claims 5-7),¹³ and the disorder(s) treated (claims 8-11).¹⁴

A. Background: VEGF Trap and AMD

Vascular endothelial growth factor (VEGF) is a promoter of angiogenesis and causes ocular disorders such as neovascular age-related macular degeneration

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- ¹² Ex. 1003 at ¶ 68 (citing Ex. 1001, Col. 22:58-62).
- ¹³ Ex. 1003 at ¶ 68 (citing Ex. 1001, Col. 22:63-23:2).
- ¹⁴ Ex. 1003 at ¶ 68 (citing Ex. 1001, Col. 23:3-13).

¹⁰ Ex. 1003 at ¶ 67.

¹¹ *Id.* at ¶ 68 (citing Ex. 1001, Col. 22:56-57).

("AMD"¹⁵).¹⁶ For many decades, VEGF antagonists have been known to inhibit VEGF and have proven to be an effective strategy in treating diseases involving pathological angiogenesis, such as AMD.¹⁷

Many VEGF antagonists have received FDA approval for AMD. For example, Ranibizumab, an anti-VEGF antibody manufactured by Genentech, received FDA approval for treatment of AMD in 2006.¹⁸ Regeneron developed Aflibercept and received FDA approval for treatment of AMD in November 2011.¹⁹

B. '345 Patent's Specification

The '345 patent's "Background" acknowledges that the prior art includes "FDA-approved treatments of angiogenic eye disorders [which] include the administration of an anti-VEGF antibody called ranibizumab (Lucentis®,

¹⁸ Ex. 1006; *see also* Ex. 1001, Col. 2:30-31 (citing "prescribing information for Lucentis® [ranibizumab], Genentech, Inc.").

¹⁹ Ex. 1001, Col. 2:51-52; see also Ex. 1007.

¹⁵ Neovascular AMD is also referred to as "Wet AMD." In this petition, "AMD" is used to refer to neovascular/wet AMD.

¹⁶ Ex. 1003 at ¶ 50.

¹⁷ *Id.* at ¶¶ 56-60.

Genentech, Inc.) on a monthly basis by intravitreal injection" but identifies a "need in the art for new administration regimens for angiogenic eye disorders, especially those which allow for less frequent dosing while maintaining a high level of efficacy."²⁰ The "Summary of Invention" describes the inventor's contribution to the art as "less frequent dosing" than, e.g., Genentech's ranibizumab:²¹

> The present inventors [sic] have surprisingly discovered that beneficial therapeutic effects can be achieved in patients suffering from angiogenic eye disorders by administering a VEGF antagonist to a patient at a frequency of once every 8 or more weeks, especially when such doses are preceded by about three doses administered to the patient at a frequency of about 2 to 4 weeks. . . . One advantage of such a dosing regimen is that, for most of the course of treatment (i.e., the tertiary doses), it allows for less frequent dosing (e.g., once every 8 weeks) compared to prior administration regimens for angiogenic eye disorders which require monthly administrations throughout the entire course of treatment. (See, e.g., prescribing information for Lucentis® [ranibizumab], Genentech, Inc.).

²⁰ Ex. 1001, Col. 1:57-67.

²¹ Id. at Col. 2:12-31 (modification "[ranibizumab]" in original).

The Detailed Description begins with a section titled "Dosing Regimens," which explains the meanings of "initial dose," "secondary dose," and "tertiary dose:"²²

The terms "initial dose," "secondary doses," and "tertiary doses," refer to the temporal sequence of administration of the VEGF antagonist. Thus, the "initial dose" is the dose which is administered at the beginning of the treatment regimen (also referred to as the "baseline dose"); the "secondary doses" are the doses which are administered after the initial dose; and the "tertiary doses" are the doses which are administered after the secondary doses.

The '345 patent has little discussion of 12-week tertiary dosing. When a 12-week tertiary dose is mentioned, the '345 patent includes it as one of 14 "or more" possible tertiary dosing frequencies: "each tertiary dose is administered at least 8 (e.g., 8, $8\frac{1}{2}$, 9, $9\frac{1}{2}$, 10, $10\frac{1}{2}$, 11, $11\frac{1}{2}$, <u>12</u>, $12\frac{1}{2}$, 13, $13\frac{1}{2}$, 14, $14\frac{1}{2}$, or more) weeks after the immediately preceding dose." ²³ When the '345 patent next mentions a 12-week dose frequency, the 12-week dose(s) is preceded by four 8-week "tertiary doses" and 12 weeks is again only one choice, among many, for the

²² *Id.* at Col. 3:42-56.

²³ Id. at Col. 3:60-62 (emphasis added).

subsequent tertiary dose.²⁴ Because the tertiary dosing in this description includes 8–week dosing followed by 12–week dosing, it does not describe the claimed 4– week secondary dosing followed by "each tertiary dose is administered 12 weeks after the immediately preceding dose."²⁵ The remainder of the '345 patent includes no discussion of a dosing regimen where "each tertiary dose is administered 12 weeks after the immediately preceding dose." The only other mention of 12 weeks is as an upper limit of a PRN²⁶ regimen,²⁷ the same dosing regimen recited in Patent Owner's 2009 Press Release and distinguished by Patent Owner during prosecution.

²⁵ This discussion in the '345 patent corresponds to Claim 5 of the parent PCT Application. That claim does not teach "sequentially administering" secondary doses and tertiary doses of the same frequency for the same reasons as the corresponding paragraph in the '345 patent.

²⁶ "PRN" is an abbreviation for "*pro re nata*" meaning "as needed." Ex. 1003 at
¶ 62.

²⁷ Ex. 1001, Col. 10:4-25.

²⁴ *Id.* at Col. 4:41-43 ("each subsequent tertiary dose is administered from 8 to 12 (e.g., 8, $8\frac{1}{2}$, 9, $9\frac{1}{2}$, 10, $10\frac{1}{2}$, 11, $11\frac{1}{2}$, **12**) weeks after the immediately preceding dose") (emphasis added).

C. Regeneron's Clinical Trials and 2009 Press Release of VEGF Treatments for Angiogenic Eye Disorders

In 2005, Regeneron began a Phase 1 study for VEGF Trap treatment of AMD with a single dose per patient.²⁸ Regeneron announced preliminary positive results in the Phase 1 study on May 1, 2006.²⁹ The Phase 1 study is described in Example 1 of the '345 patent.³⁰

In May 2007, Regeneron presented interim results of a Phase 2 VEGF Trap trial in AMD, describing dosing regimens comprising a single initial dose and one or more secondary doses at 4 or 12–week intervals for the first 12 weeks of treatment.³¹ The arms of the Phase 2 study are described in Example 2 of the '345 patent.³²

On September 14, 2009, Regeneron, through the 2009 Press Release, announced a Phase 3 trial with various initial, secondary, and tertiary doses.³³ This

²⁹ Ex. 1009.

³⁰ Ex. 1003 at ¶ 74.

³¹ Ex. 1010.

³² Ex. 1003 at ¶ 75.

³³ Ex. 1005.

²⁸ Ex. 1008.

trial is disclosed as Example 4 of the '345 patent.³⁴ According to the 2009 Press Release, after the first year of initial and 4–week secondary dosing, patients would continue to be followed and treated for another year on a flexible, criteria-based extended PRN regimen with a dose administered between four and 12 weeks.³⁵

D. '345 Patent's Prosecution History

The '345 patent claims priority to a provisional application (the first of three) filed in January 2011, more than one year after the 2009 Press Release, through a PCT application filed January 2012, a CIP application filed July 2013, and a series of continuations of the CIP.³⁶ The PCT application and three provisional applications are the only pre-AIA applications in the family.³⁷

The '282 application originally presented two independent claims, claim 21 (abandoned in prosecution) and claim 32 (issued as claim 1 of the '345 patent). In a first action, the PTO rejected all claims for obviousness type double patenting

³⁵ Id.

³⁶ Id.

³⁴ Ex. 1003 at ¶ 77.

³⁷ Ex. 1001, Cover Page. The PCT, the CIP, and the continuation applications all claim priority to the three provisional applications.

over seven Regeneron patents.³⁸ After the first action (but before Regeneron's response), a "Third Party Submission" disclosed the 2009 Press Release to the PTO.³⁹

Regeneron overcame the double patenting rejection by repeating the purported inventiveness of non-monthly dosing. In characterizing the state of the art, Regeneron told the PTO that the "standard of care for the treatment of [AMD] was to administer an antibody formulation (ranibizumab) by injection to the eye once per month."⁴⁰ Turning to the claimed invention, Regeneron said that "by administering the VEGF antagonist in accordance with a dosage regimen as claimed in independent [claim 1], it is possible to treat angiogenic eye disorders . . . on a less frequent basis than previously thought possible."⁴¹

⁴¹ Id.

³⁸ Ex. 1002, Rejection, 04/03/2019.

 ³⁹ Ex. 1002, Office Communication, 05/31/2019. Neither Petitioner nor its affiliates filed the Third-Party Submission. The record does not reflect who filed the Third-Party Submission, and Petitioner is unaware of the identity of the filer.
 ⁴⁰ Id.

In the next action, the PTO withdrew the double patenting rejection and added a rejection of (now) claims 1-11 as anticipated by the 2009 Press Release.⁴² To overcome the 2009 Press Release, Regeneron argued that claims 1-11 "require[e] tertiary dosing administered 12 weeks after the immediately preceding dose" and discussed no other limitations of the claims.⁴³ In distinguishing the disclosure of the 2009 Press Release, Patent Owner argued that "[m]ere mention of a prospective possibility of dosing at 12 weeks does not specifically indicate or teach towards a method where 12–week dosing would be undertaken, let alone successful."⁴⁴ The Examiner explicitly relied on this argument to allow claims 32-42,⁴⁵ which issued on November 10, 2020 as claims 1–11 of the '345 patent.⁴⁶

⁴³ Ex. 1002, Response, 03/16/2020, Page 4.

⁴⁴ *Id.* at 5; *see also* (Ex. 1002, Response, 01/23/2020, Pages 6-7 (The 2009 Press Release "does not disclose tertiary dosing administered 12 weeks after the immediately preceding dose. Accordingly, the press release does not anticipate the claims and the rejection should be withdrawn.")).

⁴⁵ Ex. 1002, Notice of Allowance, 04/01/20, Pages 2-3.

⁴⁶ Ex. 1001, Cover Page.

⁴² Ex. 1002, Rejection, 10/01/2019, Pages 4-5.

E. Level of Ordinary Skill in the Art

At the time of invention, a person of ordinary skill in the art of the '345 patent would have been a person with a medical doctorate, an internship and residency in ophthalmology, and a 1-year medical retina fellowship or 2-year vitreoretinal surgical fellowship.⁴⁷ A person with less education but more relevant practical experience with retinal disease treatment may also be a person of ordinary skill in the art.⁴⁸

V. CLAIM CONSTRUCTION

Pursuant to 83 Fed. Reg. 51340, a claim is construed using the standard set forth by *Phillips v. AWH Corp.*, 415 F.3d 1303 (Fed. Cir. 2005) (*en banc*). Petitioner relies on the plain language of the claims in the '992 patent to demonstrate that the claims are anticipated and/or obvious in light of the prior art. Accordingly, a formal claim construction is unnecessary. *See Hakim v. Cannon Avent Grp.*, *PLC*, 479 F.3d 1313, 1318-19 (Fed. Cir. 2007) ("When there is no dispute as to the meaning of a term that could affect the disputed issues of the litigation, 'construction' may not be necessary."); *Vivid Techs.*, *Inc.*, 200 F.3d at

⁴⁸ Id.

⁴⁷ Ex. 1003 at ¶ 82.

803 (only those terms that are in controversy need to be construed and only to the extent necessary to resolve the controversy).⁴⁹

VI. THE '345 PATENT IS ELIGIBLE FOR PGR

A patent is eligible for PGR if it "contains or contained at any time . . . a claim to a claimed invention that has an effective filing date as defined in section 100(i) of title 35, United States Code, that is on or after [March 16, 2013]." AIA §§ 3(n)(1), 6(f)(2)(A). The "effective filing date" of a patent is defined under 35 U.S.C. § 100(i)(1)(B) as "the filing date of the earliest application for which the patent . . . is entitled, as to such invention, to a right of priority under section 119, 365(a), or 365(b) or to the benefit of an earlier filing date under section 120, 121, or 365(c)." In order for a patent application to be entitled to a "right of priority" or "an earlier filing date" based upon an earlier filed application, the earlier filed application must have been disclosed "in the manner provided by section 112(a) (other than the requirement to disclose the best mode)." 35 U.S.C. § 119(e)(1); 35 U.S.C. § 120.

⁴⁹ Petitioner reserves the right to propose constructions for claim terms in this proceeding in response to arguments raised by Patent Owner in any future submission.

Accordingly, for purposes of determining PGR eligibility, a patent application may rely on the filing date of an earlier filed application only if it is described in the manner provided by 35 U.S.C. § 112(a), including written description support for the claims. If an application is not entitled to claim priority to a prior application, the effective filing date for the patent is the "actual filing date of the patent or the application for the patent containing a claim to the invention." 35 U.S.C. § 100(i)(1)(A). Here, that means that the '345 patent's earliest effective filing date is either in July 2013 or in October 2018. The '345 patent is thus eligible for PGR.

That the '345 patent is a continuation "transition" application does not affect the requirements for PGR eligibility.

A. "[T]he Angiogenic Eye Disorder Is . . . Branch Retinal Vein Occlusion" of Claim 8 Is not Supported by a Pre-AIA Application

Claim 8 includes the indication "branch retinal vein occlusion," which first appeared⁵⁰ in the '345 patent's family when Regeneron filed U.S. Application No. 13/940,370 (the "'370 application") on July 12, 2013, as a CIP of the PCT application. ⁵¹ The '345 patent, and its grandparent '370 application, thus contains at least one claim that has an effective filing date on or after March 16, 2013.

⁵⁰ Ex. 1011, at ¶ 0026 and claim 6.

⁵¹ Ex. 1001, Cover Page.

"[S]ome subject matter of a CIP application is *necessarily different* from the original subject matter." *Uni. of W. Va. Bd. of Trustees v. VanVoorhies*, 278 F.3d 1288 (Fed. Cir. 2002) (citing MPEP § 201.08 (7th ed. Rev. 1 Feb. 2000)) (emphasis added). The CIP's new subject matter is the additional eye disorders not disclosed in earlier filed patent applications, such as branch retinal vein occlusion ("BRVO").

In the CIP application, BRVO appears in a section titled "Angiogenic Eye Disorders," which consists of the paragraph below.⁵² To illustrate the subject matter added to the CIP, the paragraph below compares the pre and post 2013 paragraph, where underlined text indicates subject matter not appearing in the '370 application's parent, *italics* indicates text moved in the paragraph, and *italics* with strikethrough indicates the original position of the moved text (no text was deleted⁵³).

The methods of the present invention can be used to treat any angiogenic eye disorder. The expression "angiogenic eye disorder," as used herein, means any disease of the eye which is caused by or associated with the growth or proliferation of blood vessels or by blood vessel leakage. Non-limiting examples of angiogenic eye disorders that

⁵² Ex. 1011; *see also* Ex. 1003 at ¶ 120.

⁵³ Ex. 1003 at ¶ 123.

are treatable using the methods of the present invention include choroidal neovascularization, age-related macular degeneration (e.g., wet AMD, *diabetic retinopathies*, exudative AMD, etc.), retinal vein occlusion (RVO), diabetic macular edema (DME), central retinal vein occlusion (CRVO; e.g., macular edema following CRVO), branch retinal vein occlusion (BRVO), diabetic macular edema (DME), choroidal neovascularization (CNV; e.g., myopic CNV), iris neovascularization, neovascular glaucoma, post-surgical fibrosis in glaucoma, proliferative vitreoretinopathy (PVR), optic disc neovascularization, corneal neovascularization, retinal neovascularization, vitreal neovascularization, pannus, ptervgium, vascular retinopathies retinopathy, and diabetic retinal neovascularization.

BRVO was neither explicitly nor implicitly disclosed in the pre-AIA applications.⁵⁴ As the above paragraph demonstrates, the PCT application (i.e., the latest pre-AIA application) was completely silent about BRVO, and it was only added in the AIA CIP application. Having examined the pre-AIA indications, one of skill in the art in 2011-13 would have recognized that all of the listed disorders are different from BRVO; for example, each has different pathologies and has

⁵⁴ *Id.* at ¶ 120-24.

different standards for treatment.⁵⁵ One of skill in the art would not understand that successful treatment of one vascular disease (e.g., choroidal neovascularization, AMD, diabetic retinopathies, DME, CRVO, corneal neovascularization, or retinal neovascularization) means another (e.g., BRVO) is necessarily treated.⁵⁶

Other disclosures within the specification reinforce the inventor's failure to possess a treatment for BRVO. In the "Background," the PCT application identifies various eye disorders treatable by VEGF antagonists, but nowhere mentions BRVO.⁵⁷ Similarly, the "Treatment Population and Efficacy" section identifies various disorders treatable by the "present invention," but no mention of BRVO.⁵⁸ The examples describe various disorders studied in the clinical trials, but none of the examples in pre-AIA applications (i.e., Examples 1-6) mention BRVO.⁵⁹

⁵⁵ *Id.* at 124.

⁵⁶ *Id.* at 124-25.

⁵⁷ Ex. 1001, Col. 1:31-61 (the CIP did not modify this teaching).

⁵⁸ Id. at Col. 7:26-44.

⁵⁹ Example 7 was added with the CIP '370 application. The additional eye disorders are the only subject matter in Example 7 not supported by the PCT.

The pre-AIA applications' discussion of "central retinal vein occlusion" ("CRVO") does not constitute a written description of BRVO. One of skill in the art would not recognize a disclosed treatment of CRVO to be possession of a treatment for BRVO.⁶⁰ These are different indications, with their own standard of care in 2012–13.⁶¹ Anatomically, they are different.⁶² Further, they affect populations differently—Asians and Hispanics appeared to have an elevated risk of BRVO compared to Caucasians, whereas no similar difference was found for CRVO.⁶³ The specification and prosecution history confirm that CRVO and BRVO are different—Patent Owner claimed BRVO as a separate indication from CRVO and amended the specification to explicitly recite BRVO, confirming that one of skill in the art would recognize that BRVO and CRVO are separate indications.⁶⁴ Consistently, Regeneron conducted separate trials for BRVO and CRVO.⁶⁵

- ⁶⁰ Ex. 1003 at ¶¶ 126-29.
- ⁶¹ *Id.* at ¶ 127.
- ⁶² Id. at ¶ 126.
- ⁶³ Id.
- ⁶⁴ *Id.* at ¶ 130.
- ⁶⁵ *Id.* at ¶ 131.

Accordingly, none of the pre-AIA applications provides adequate written description support for claim 8⁶⁶ of the '345 patent. The patent is therefore eligible for PGR.

B. The Dosing Regimen of Claim 1 Is not Supported by a Pre-AIA Application

As explained in Section VIII below, the '345 patent is invalid under

35 U.S.C. § 112(a) for failing to support the claimed 12–week tertiary dosing regimen. That regimen first entered the patent family when Regeneron filed the '345 patent. For this reason, the earliest filing date of the '345 patent is its actual filing date, October 28, 2018, making the patent eligible for PGR.

VII. GROUNDS 1 & 2: THE '345 PATENT'S CLAIMS ARE ANTICIPATED AND OBVIOUS

Petitioner respectfully requests the Board cancel all claims of the '345 patent on the following prior-art grounds:

Ground 1: Claims 1-11 are anticipated by Shams under pre-AIA 35 U.S.C. § 102(b) and post-AIA 35 U.S.C. § 102(a)(1).

⁶⁶ Additionally, the '345 patent is eligible for PGR because a post-AIA application in its priority chain, the '370 application, includes a claim ('370 application claim
6) which is not supported by a pre-AIA patent application. The '370 application claim
claim 6 includes the same list of disorders as the '345 patent claim 8.

Ground 2: Claims 1-11 are rendered obvious by the 2009 Press Release in view of Shams under pre-AIA 35 U.S.C. § 103 and post-AIA 35 U.S.C. § 103.

A. Ground 1: The '345 Patent's Claims Are Anticipated by Shams 1. Shams

Shams published on May 4, 2006 and thus is prior art to the '345 patent under pre-AIA 35 U.S.C. § 102(b) and 35 U.S.C. §§ 102(a)(1) and (2). Shams discloses effective VEGF antagonist treatments with extended dosing frequencies: "It has been discovered that the treatment effects of a VEGF antagonist, e.g., Ranibizumab, are maintained for an extended period of time, such as more than one month."⁶⁷ Exemplary VEGF antagonists provided in Shams include Regeneron's VEGF trap.⁶⁸ Six years after Shams' filing and five years after its publication, the '354 patent described the same "surprising discovery:"⁶⁹

> The present inventors [sic] have surprisingly discovered that beneficial therapeutic effects can be achieved in patients suffering from angiogenic eye disorders by administering a VEGF antagonist to a patient at a frequency of once every 8 or more weeks, especially when such doses are preceded by

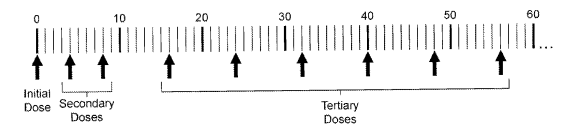
⁶⁹ Ex. 1001, Col. 2, Lines 12-31 (emphasis added) (modification "[ranibizumab]" in original).

⁶⁷ Ex. 1004, Page 21, Lines 27-28.

⁶⁸ *Id.* at Page 6, Lines 27-33.

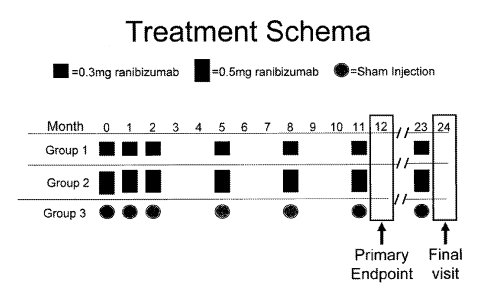
about three doses administered to the patient at a frequency of about 2 to 4 weeks. . . . One advantage of such a dosing regimen is that, for most of the course of treatment (i.e., the tertiary doses), *it allows for less frequent dosing (e.g., once every 8 weeks) compared to prior administration regimens for angiogenic eye disorders which require monthly administrations* throughout the entire course of treatment. (See, e.g., prescribing information for Lucentis® [ranibizumab], Genentech, Inc.).

Like the '345 patent, Shams describes a dosing regimen of "initial, secondary, and tertiary" doses. Shams uses different terms ("first individual doses" and "second individual doses") to describe the dosing scheme, but the difference is in name only.⁷⁰ The '345 patent Figure 1 illustrates the "initial dose," "secondary doses," and "tertiary doses:"



⁷⁰ Ex. 1003 at ¶¶ 90, 96.

Shams Figure 2 "illustrates a dosing regimen for treating, e.g., age-related macular degeneration (AMD) with a VEGF antagonist."⁷¹ The figure includes three groups: Group 1 received a 0.3 mg dose of VEGF antagonist; Group 2 received a 0.5 mg dose of VEGF antagonist; and Group 3 received a sham injection.⁷² Each group illustrates an initial dose at "month 0," two secondary doses at "month 1" and "month 2," and subsequent tertiary doses every 3 months thereafter until two years.

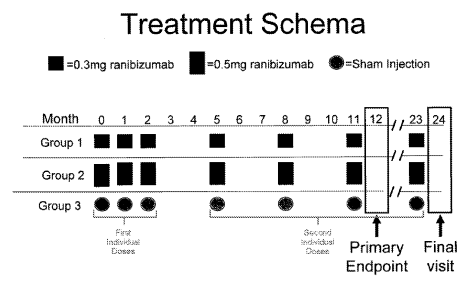




⁷¹ Ex. 1004, Page 6, Lines 8-9.

⁷² *Id.* at Page 31, Lines 8-13.

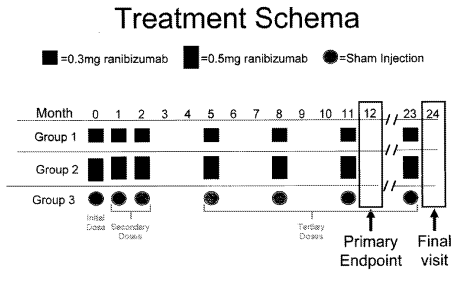
Figure 2 is annotated below to include the "first individual doses" and "second individual doses" terminology used by Shams.





Using the '345 patent's terminology, Shams' "first individual doses" encompass initial doses (month 0) and secondary doses (months 1 and 2), and the application's "second individual doses" (months 5, 8, 11, ...) are thus "tertiary doses."⁷³ The annotated Shams Figure 2 below illustrates the same regimen, but using the '345 patent's terminology.

⁷³ Ex. 1003 at ¶¶ 90-96.





In addition to the '345 patent's purported discovery of less frequent dosing and its "initial, secondary, and tertiary" dosing regimen, Shams discloses claim 1's specific dosing frequency of "each secondary dose is administered 4 weeks after the immediately preceding dose" and "each tertiary dose is administered 12 weeks after the immediately preceding dose:" "In one embodiment of the invention, the first individual doses are administered at one month intervals (e.g., about 3 individual doses). The second [individual] dose is administered less frequently, e.g., at three month intervals (e.g., about 6 individual doses)."⁷⁴ This dosing frequency is illustrated in Figure 2.

⁷⁴ Ex. 1004, Page 23, Lines 16-18.

In addition to teaching the limitations of claim 1, Shams also discloses the limitations of claim 2 (specific drug administered), claims 3 and 4 (modes of administration), claim 5-7 (dose amounts), and claims 8-11 (disorders treated).

2. Claim 1

a) Shams discloses a "method for treating an angiogenic eye disorder in a patient, said method comprising sequentially administering to the patient"

Shams' "Summary of Invention" describes:75

Methods for *treating intraocular neovascular disease* are provided. For example, methods include *administering to a mammal* a number of *first individual doses* of a VEGF antagonist, followed by *administering* to the mammal *a number of second individual doses* of the antagonist, wherein the second individual doses are administered less frequently than the first individual doses.

Shams' "intraocular neovascular disease" meets claim 1's "angiogenic eye disorder."⁷⁶ "Administering to a mammal"⁷⁷ discloses the claimed "administering

⁷⁶ Ex. 1003 at ¶ 88 (observing that the '345 patents lists, under "angiogenic

disorders," disorders that Shams lists as "intraocular neovascular diseases").

⁷⁷ Shams describes "human patients" as preferred examples of "mammals." See

⁷⁵ Id. Page 4, Line 31-Page 5, Line 2 (emphasis added).

Ex. 1004 at Page 23, Lines 30-34 ("Another aspect of the invention is the treatment

to the patient." ⁷⁸ "Sequentially administering" is taught by Shams' "administering a number of first individual doses . . . followed by administering . . . a number of second individual doses." ⁷⁹ Shams' administration meets the '345 patent's definition of "sequentially administered:" "each dose of VEGF antagonist is administered to the patient at a different point in time, e.g., on different days separated by a predetermined interval (e.g., hours, days, weeks or months)."⁸⁰

b) Shams discloses the claimed initial dose of VEGF

Shams discloses claim 1's "single initial dose of a VEGF antagonist." The '345 patent defines an "initial dose" as "the dose which is administered at the beginning of the treatment regimen (also referred to as the 'baseline dose')."⁸¹ The initial dose of the "number of first individual doses" in Shams (*see* annotated

of an intraocular neovascular disease, e.g., wet form AMD, by administering to a mammal, preferably a human patient, a number of first individual doses of a compound, e.g., a VEGF antagonist, followed by administering a number of second individual doses of the compound.")

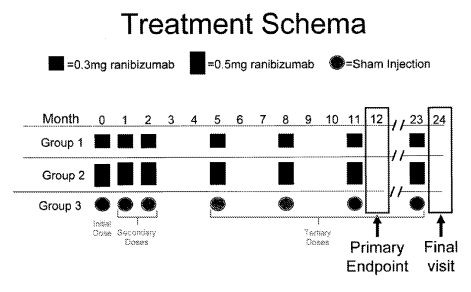
⁷⁸ Ex. 1003 at ¶ 88.

⁷⁹ *Id.* at ¶ 89.

⁸⁰ Ex. 1001, Col. 3:32-36.

⁸¹ Id. at Col. 3:44-46.

Figure 2 below) corresponds to the claimed "single initial dose of a VEGF antagonist."





Shams discloses "wherein the VEGF antagonist is a receptor-based chimeric molecule comprising an immunoglobin-like (lg) domain 2 of a first VEGF receptor which is Flt1 and lg domain 3 of a second VEGF receptor which is Flk1, and a multimerizing component." Shams describes VEGF antagonist as including "VEGF-Trap (Regeneron)."⁸² One of skill in the art would understand that "VEGF Trap (Regeneron)" includes Regeneron's fusion protein, which, in 2006, included "a receptor-based chimeric molecule comprising an immunoglobin-like (lg)

⁸² Ex. 1004, Page 6, Line 27 – Page 7, Line 6.

domain 2 of a first VEGF receptor which is Fltl and lg domain 3 of a second VEGF receptor which is Flk1, and a multimerizing component."⁸³ Consistent with its description of specific VEGF antagonists applicable to the 4+12 week dosing regimen, Shams teaches that "[a]ny compound which binds to VEGF or a VEGF receptor and reduces the severity of symptoms or conditions associated with an intraocular neovascular disease may be used in this embodiment of the invention."⁸⁴ Regeneron's VEGF Trap binds to VEGF receptors to hinder VEGF interaction and interfere with the normal biological activity of VEGF.⁸⁵

c) Shams discloses the claimed "followed by one or more secondary doses of the VEGF antagonist . . . wherein each secondary dose is administered 4 weeks after the immediately preceding dose"

The '345 patent defines "secondary dose" as "the doses which are administered after the initial dose." ⁸⁶ Shams discloses doses administered after the

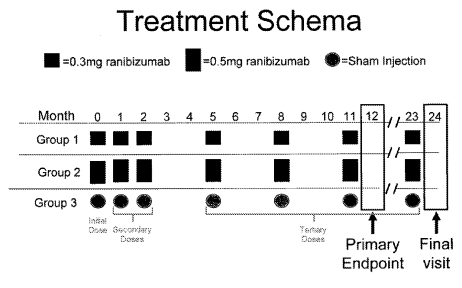
⁸⁴ Ex. 1004, Page 26, Lines 6-8; *see also id.* at Page 22, Lines 15-18 ("The term "therapeutic" in this context means that the compounds binds to the ligand, VEGF, and produce a change in the symptoms or conditions associated with the disease or condition which is being treated.")

⁸⁵ Ex. 1001, Col. 4:48-60.

⁸⁶ Id. at Col. 3:46-48.

⁸³ Ex. 1003 at ¶ 89.

initial dose; for example, in Figure 2, the initial dose is followed by a second dose and a third dose.





The second and third doses are thus "secondary doses," as defined by the '345 patent.

Further, Shams discloses that the one or more secondary doses of VEGF antagonist are "administered 4 weeks after the immediately preceding dose." For example, Shams discloses "the first individual doses are administered at one month intervals (e.g., about 3 individual doses),"⁸⁷ which are illustrated in Shams' Figure 2 (annotated above).

⁸⁷ Ex. 1004, Page 23, Lines 16-17.

One of skill in the art would understand that Shams' "one month interval" dosing discloses claim 1's "4 week" dosing frequency for at least three separate reasons.⁸⁸ <u>First</u>, the '345 patent equates monthly dosing to 4 week dosing.⁸⁹ For instance, the '345 patent's examples are based on an equivalence between monthly dosing and 4 week dosing: "For purposes of the following Examples, 'monthly' dosing is equivalent to dosing once every four weeks."⁹⁰ The patent's specific discussion of Example 4 also equates monthly and 4 week dosing: "patients receiving VEGFT 2 mg monthly (2Q4) achieved a statistically significant greater mean improvement in visual acuity at week 52 versus baseline."⁹¹ "2Q4" is understood in the art to be shorthand for dosing "every 4 weeks"⁹² and so the '345 patent's use of "monthly (2Q4)" would be understood to mean that "monthly"

⁸⁹ *Id.* at ¶ 93.

90 Ex. 1001, Col. 7:67-8:2.

⁹¹ Id. at Col. 13:47-49.

⁹² Ex. 1003 at ¶ 93; see also Ex. 1004, Page 22:31-32 (describing a sequence of "weekly, biweekly, and monthly" dosing).

⁹³ Ex. 1003 at ¶ 93.

⁸⁸ Ex. 1003 at ¶¶ 92-94.

describes a study where "patients received 6 monthly injections," and then described those injections as administered "once every four weeks from Week 0 through Week 20."⁹⁴ Thus, the '345 patent, in the paragraph describing Example 6, uses the terms interchangeably. ⁹⁵ In Example 7, Regeneron described a "dosing regimen within the scope of the present invention" as including VEGF Trap "administered by intravitreal injection once every 4 weeks (monthly).⁹⁶ Once again, the '345 patent discloses that "4 weeks" and "monthly" are to be used interchangeably.⁹⁷

Second, it is common in the art to use "one month dosing" and "4 week dosing" interchangeably.⁹⁸ Typically, surgeons and patients calendar follow-up treatments on a weekly basis (i.e., the same day (and time) of a following week), instead of returning on the same date in a future month.⁹⁹ In those cases, returning "monthly" is understood to mean returning in 4 weeks on the same day of the

- 94 Ex. 1001 at Col. 14:59-66.
- ⁹⁵ Ex. 1003 at ¶ 93.
- ⁹⁶ Ex. 1001 at Col. 15:40-41.
- ⁹⁷ Ex. 1003 at ¶ 93.

⁹⁸ *Id.* at ¶ 92.

⁹⁹ Id.

week. ¹⁰⁰ Additionally, many surgeons have practices in different locations and visit a specific office on the same days every week. ¹⁰¹ In such instances, the surgeon and patient may arrange a one-month "follow-up" but imply meeting on the same day in a future week. ¹⁰²

<u>Third</u>, Regeneron has frequently equated monthly and 4–week dosing.¹⁰³ For example, Heier (which Patent Owner cited as evidence of "unexpected results" during prosecution¹⁰⁴) equates monthly with 4 week dosing: "Patients were randomized to intravitreal aflibercept 0.5 mg monthly (0.5q4), 2 mg monthly (2q4), 2 mg every 2 months after 3 initial monthly doses (2q8), or ranibizumab 0.5 mg monthly (Rq4)."¹⁰⁵ In a different Heier publication, Regeneron stated that

- 100 Id.
- ¹⁰¹ Id.
- 102 Id.
- ¹⁰³ *Id.* at ¶ 94.
- ¹⁰⁴ Ex. 1002, Response, 06/28/2019, Page 8.

¹⁰⁵ Ex. 1012, Page 2538, "Intervention." *See also id.* at 2546 ("all treatment groups' dosing intervals were changed to a common protocol of modified quarterly dosing with their originally randomized dose and drug ("[A]ll patients were

"During the 12–week fixed dosing phase, patients in the monthly dosing groups received 0.5 or 2 mg of VEGF Trap-Eye every 4 weeks on day 0 and at weeks 4, 8, and 12 for a total of 4 doses."¹⁰⁶

d) Shams discloses the claimed "followed by one or more tertiary doses of the VEGF antagonist... each tertiary dose is administered 12 weeks after the immediately preceding dose"

Shams discloses the secondary dose(s) "followed by one or more tertiary doses of the VEGF antagonist." The '345 patent defines "tertiary dose" as "the doses which are administered after the secondary doses." ¹⁰⁷ Shams' "second individual doses" are administered after the "first individual doses" (which correspond to the claimed "secondary doses"), and, thus, the second individual doses are the claimed "tertiary doses."¹⁰⁸

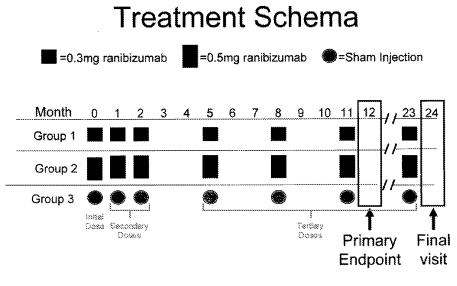
monitored monthly and received a minimum of dosing every 12 weeks with

interim as-needed monthly intravitreal injections).").

¹⁰⁶ Ex. 1013 at 1110, Legend for Figure 2.

¹⁰⁷ Ex. 1001, Col. 3:48-49.

¹⁰⁸ Ex. 1003 at ¶¶ 95-96.





Further, Shams discloses that the tertiary "doses are administered at three month intervals (e.g., about 6 individual doses)."¹⁰⁹ One of skill in the art would understand that a three month dosing frequency discloses the claimed "12 week" dosing.¹¹⁰ As explained above, one of skill in the art would equate monthly dosing with a 4–week frequency, evidenced by the '345 patent's specification and Regeneron's prosecution arguments. One of skill in the art would further understand "3 month" dosing to be equivalent to a 12–week frequency.¹¹¹

¹¹¹ Id.

¹⁰⁹ Ex. 1004, Page 5, Lines 23-24

¹¹⁰ Ex. 1003 at ¶ 97.

Surgeons frequently refer to 12–week dosing as "three month" dosing or "quarterly" dosing.¹¹² Consistently, a 2011 Regeneron publication equated "4 week" with "monthly" dosing and equated "12–week" with "quarterly" dosing when describing the results of Regeneron's Phase 2 study: "During the 12–week fixed dosing phase, <u>patients in the monthly dosing groups</u> received 0.5 or 2 mg of VEGF Trap-Eye <u>every 4 weeks</u> on day 0 and at weeks 4, 8, and 12 for a total of 4 doses; <u>those in the quarterly dosing groups</u> received 0.5, 2, or 4 mg of VEGF Trap-Eye <u>every 12 weeks</u> on day 0 and at week 12 for a total of 2 doses."¹¹³ Thus, Shams discloses "each tertiary dose is administered 12 weeks after the immediately preceding dose."

3. Dependent claims

a) Claim 2: Shams discloses the claimed drug

Shams discloses that the "VEGF antagonist" includes "VEGF Trap (Regeneron)."¹¹⁴ Shams also discloses that the VEGF antagonist treats eye

¹¹² Id.

¹¹³ Ex. 1013 at 1110, Legend for Figure 2 (emphasis added).

¹¹⁴ Ex. 1003 at ¶ 99.

disorders,¹¹⁵ which would lead one of skill in the art to understand that "VEGF Trap (Regeneron)" refers to Regeneron's VEGF Trap treatment for eye disorders.¹¹⁶ "Aflibercept" is another name for Regeneron's VEGF Trap treatment for eye disorders.¹¹⁷ Shams thus discloses "wherein the VEGF antagonist is aflibercept."

b) Claims 3 and 4: Shams discloses the claimed modes of administration

Shams discloses "wherein all doses of the VEGF antagonist are administered to the patient by intraocular administration" (claim 3), and "the intraocular administration is intravitreal administration" (claim 4). ¹¹⁸ For example, Shams discloses "[t]he therapeutic compound for treatment of an intraocular neovascular disease is typically administered by ocular, intraocular, and/or intravitreal injection."¹¹⁹

Diseases").

¹¹⁷ Id.

¹¹⁸ Id. at ¶ 98.

¹¹⁹ Ex. 1004, Page 25, Lines 15-16.

¹¹⁵ E.g., Ex. 1004 at Title ("Method for Treating Intraocular Neovascular

¹¹⁶ Ex. 1003 at ¶ 99.

c) Claims 5-7: Shams discloses the claimed dose amounts

Shams explicitly discloses "wherein all doses of the VEGF antagonist comprise from about 0.5 mg to about 2 mg of the VEGF antagonist" (claim 5) and "wherein all doses of the VEGF antagonist comprise 0.5 mg of the VEGF antagonist" (claim 6). ¹²⁰ For example, Shams discloses administering a 0.5 mg dose.¹²¹ Further, Shams discloses a range $(0.1 \text{ mg} - 20 \text{ mg})^{122}$ which covers the range of claim 5 (0.5 mg – 2.0 mg), the dose of claim 6 (0.5 mg), and the dose of claim 7 (2.0 mg).

d) Claims 8-11: Shams discloses the disorders treated

Shams discloses "wherein the angiogenic eye disorder is selected from the group consisting of: age related macular degeneration, diabetic retinopathy, diabetic macular edema, central retinal vein occlusion, branch retinal vein occlusion, and corneal neovascularization" (claim 8), "wherein the angiogenic eye disorder is age related macular degeneration" (claim 9), "wherein the angiogenic eye disorder is diabetic retinopathy" (claim 10), and "wherein the angiogenic eye

¹²⁰ Ex. 1003 at ¶ 101.

¹²¹ See, e.g., Ex. 1004 at Figure 2; Page 31, Lines 8-9.

¹²² Ex. 1004, Page 24, Lines 18-20.

disorder is diabetic macular edema" (claim 11). ¹²³ For example, Shams lists various "intraocular neovascular disease[s]" treatable by the disclosed VEGF antagonists, including "age-related macular degeneration (AMD), diabetic and other ischemia-related retinopathies, diabetic macular edema." ¹²⁴

B. Ground 2: The '345 Patent's Claims Are Rendered Obvious by the 2009 Press Release in view of Shams

1. The 2009 Press Release

The 2009 Press Release published September 14, 2009, more than one year before the '345 patent's earliest priority date, and thus is prior art to the '345 patent under 35 U.S.C. §§ 102(a)(1) and (2).¹²⁵ The 2009 Press Release teaches, among other arms, "that patients received intravitreal doses of 0.5 mg or 2g VEGF Tap-Eye [sic] at 4–week intervals in the first year, followed by continual treatment for another year on a flexible, PRN regiment, with a dose administered at least every 12 weeks."¹²⁶ The PTO cited this teaching in rejecting the '345 patent as anticipated by the 2009 Press Release. To overcome the 2009 Press Release, Regeneron argued that as-needed tertiary dosing between 4 and 12 weeks does not

- ¹²⁴ Ex. 1004, Page 21, Lines 1-6.
- ¹²⁵ Ex. 1005.

¹²⁶ Id.

¹²³ Ex. 1003 at ¶ 102.

explicitly disclose a tertiary 12–week dosing frequency component because "[a] practitioner of ordinary skill in the art would not have understood a PRN dosing regimen with 4 and 12–week limits as encompassing a [12–week dosing] regimen."¹²⁷ Regeneron also argued that the claims were not inherently disclosed in the Press Release because "[t]hough the Press Release discussed a PRN dosing regimen wherein a dose interval <u>may</u> extend out as far as 12 weeks, the dosages administered to patients were not <u>necessarily</u> this infrequent. For this reason, the Press Release was insufficient as an inherently anticipating reference."¹²⁸

An examiner interview followed on March 6, 2020.¹²⁹ Later that month, Regeneron filed a supplemental response presenting more arguments to distinguish claims 32-42 (issued claims 1-11) from the 2009 Press Release:¹³⁰

> Claims 32-42 relate to a method requiring tertiary dosing administered 12 weeks after the immediately preceding dose. There is a single appearance of the words "12 weeks" within the fourth paragraph of the Press Release. However, this paragraph is referring to a "flexible, criteria-based extended PRN regimen with a dose administered at least every 12 weeks, but not more

 130 Id.

¹²⁷ Ex. 1002, Response, 01/23/2020, Pages 6-8.

¹²⁸ *Id.* (emphasis in original).

¹²⁹ Ex. 1002, Response, 03/16/2020, Page 4.

often than every four weeks". As explained in our January 23, 2020 Response, this is not a disclosure of a regimen having 12– week tertiary dosing as specified in the claims. Mere mention of a prospective possibility of dosing at 12 weeks does not specifically indicate or teach towards a method where 12–week dosing would be undertaken, let alone successful.

The Examiner explicitly relied on Regeneron's arguments that the 2009

Press Release did not teach or suggest 12-week tertiary dosing.

This petition adds Shams to provide the 12–week tertiary dosing allegedly absent from the 2009 Press Release. Specifically, this petition relies on a different teaching in the Press Release than that relied upon by the PTO, a tertiary 8–week dosing regimen, and combines with Shams' teachings of 12-week tertiary dosing.

2. Claim 1

a) The 2009 Press Release teaches a "method for treating an angiogenic eye disorder in a patient, said method comprising sequentially administering to the patient"

The 2009 Press Release teaches a "method for treating an angiogenic eye disorder in a patient, said method comprising sequentially administering to the patient." The 2009 Press Release announced "Phase 3 clinical trials evaluating VEGF Trap-Eye in the treatment of the neovascular form of age-related macular

degeneration (wet AMD).^{*131} The 2009 Press Release explains: "In the first year of the studies, the safety and efficacy of VEGF Trap-Eye at doses of 0.5 mg and 2.0 mg administered at four week intervals and 2.0 mg at an eight-week dosing interval following one additional 2.0 mg dose at week four are being evaluated." ¹³² The 2009 Press Release also describes a "development for the treatment of [DME]" where VEGF Trap-Eye is dosed at 0.5 mg or 2.0 mg monthly, 2 mg on an as-needed basis after three monthly loading doses, or 2 mg every eight weeks after three monthly loading doses.¹³³ This disclosure in the 2009 Press Release corresponds to the '345 patent's Examples 4 and 5¹³⁴ and teaches a "method for treating an angiogenic eye disorder in a patient, said method comprising sequentially administering to the patient."¹³⁵

b) The 2009 Press Release teaches the claimed initial dose of VEGF

The 2009 Press Release teaches "a single initial dose of a VEGF antagonist." The 2009 Press Release teaches two studies with 8-week tertiary dosing, both of

- ¹³³ *Id.* at 2, Second Paragraph.
- ¹³⁴ Ex. 1003 at ¶¶ 77-78.

¹³⁵ Id. at ¶ 106.

¹³¹ Ex. 1005, Title.

¹³² Id. at 1, Fourth Paragraph.

which include an "initial dose." First, the Press Release teaches: "In the first year of the studies, the safety and efficacy of VEGF Trap-Eye at doses of 0.5 mg and 2.0 mg administered at four week intervals and 2.0 mg at an eight-week dosing interval following one additional 2.0 mg dose at week four are being evaluated." Second, the 2009 Press Release teaches: "VEGF Trap-Eye is also in Phase 2 development for the treatment of Diabetic Macular Edema (DME). VEGF Trap-Eye dosed at 0.5 mg or 2 mg monthly, 2 mg every eight weeks after three monthly loading doses." The first dose in each of these arms corresponds to the claimed "single initial dose."¹³⁶

The 2009 Press Release teaches "wherein the VEGF antagonist is a receptorbased chimeric molecule comprising an immunoglobin-like (lg) domain 2 of a first VEGF receptor which is Fltl and lg domain 3 of a second VEGF receptor which is Flk1, and a multimerizing component." The 2009 Press Release describes studies related to VEGF Trap-Eye, which one of skill in the art would understand includes "a receptor-based chimeric molecule comprising an immunoglobin-like (lg) domain 2 of a first VEGF receptor which is Fltl and lg domain 3 of a second VEGF receptor which is Flk1, and a multimerizing component." ¹³⁷

¹³⁶ *Id.* at ¶ 106.

¹³⁷ Id. at ¶ 105.

c) The 2009 Press Release teaches the claimed "one or more secondary doses"

The 2009 Press Release teaches the initial dose "followed by one or more secondary doses of the VEGF antagonist . . . wherein each secondary dose is administered 4 weeks after the immediately preceding dose."¹³⁸ In one study, the 2009 Press Release teaches an initial dose of 2.0 mg and "one additional 2.0 mg dose at week four." In other words, the 2009 Press Release teaches a secondary dose of VEGF Trap-Eye at week four, i.e., "administered 4 weeks after the immediately preceding dose." In another study, the 2009 Press Release teaches that after an initial dose of 2.0 mg, two additional "monthly loading doses" are administered. In other words, the 2009 Press Release teaches two secondary doses of VEGF Trap-Eye at weeks four and eight, i.e., "administered 4 weeks after the immediately preceding dose."

d) The 2009 Press Release in view of Shams teaches the claimed "one or more tertiary doses"

The 2009 Press Release teaches the secondary dose(s) "followed by one or more tertiary doses of the VEGF antagonist."¹³⁹ In the first study, the 2009 Press Release teaches 8–week tertiary doses of VEGF Trap-Eye following the week four

¹³⁸ *Id.* at ¶ 107.

¹³⁹ Id. at ¶ 108.

dose and, in the second, the press release teaches 8–week tertiary dosing teaches following the three monthly loading doses. Thus, the 2009 Press Release teaches "one or more tertiary doses."

The 2009 Press Release's 8–week tertiary doses do not meet "each tertiary dose is administered 12 weeks after the immediately preceding dose." Shams teaches an effective treatment of "intraocular neovascular disease" with tertiary doses every three months.¹⁴⁰ It would have been natural for one of skill in the art to look at Shams' teachings when considering the 2009 Press Release's 4 + 8 week dosing: (1) Shams was assigned to Genentech, who was a research leader in the early stages of VEGF antagonist treatment; (2) Regeneron's clinical trials use Genentech's ranibizumab as the control dose; and (3) Shams lists Regeneron's VEGF Trap as a suitable antagonist for its 12 week tertiary dosing program.¹⁴¹

One of skill in the art would have been motivated at least by market forces to extend the 2009 Press Release's 8–week tertiary dosing. As recognized in the 2009 Press Release, "monthly office visits and examinations . . . are inconvenient for these often elderly patients."¹⁴² The inconvenience included the physical

¹⁴⁰ E.g., Ex. 1004, Page 23, Lines 9-11.

¹⁴¹ Ex. 1003 at ¶ 112.

¹⁴² Ex. 1005 at 1, Third Paragraph.

discomfort of an intraocular injection and the mobility limitations of some elderly patients.¹⁴³ The monthly injections also inconvenienced retinal specialists because their practices could quickly fill with monthly maintenance injections.¹⁴⁴ Also, the high price (\$2,000 per injection) of Lucentis was a significant market force that drove longer tertiary dosing.¹⁴⁵

Thus, claim 1 is nothing more that the simple substitution of Shams' 12– week tertiary dose for the 2009 Press Release's 8–week tertiary dose.¹⁴⁶ Similarly, the known work by Genentech (12–week tertiary dosing) would prompt variations in the 2009 Press Release for at least the reason that market forces provided an incentive to extend the 8–week tertiary dosing.¹⁴⁷ Further, the modification merely combines prior art elements (Shams' 12–week tertiary dosing) to a known method (the 2009 Press Release's 4–week secondary dosing plus 12–week tertiary dosing) to arrive at a predicate result (a successful treatment of angiogenic eye disorders). The success was predictable because Shams teaches a successful 4 +

 144 Id.

¹⁴⁵ Id.

¹⁴⁶ Id. at 115.

¹⁴⁷ *Id.* at 111-12.

¹⁴³ Ex. 1003 at ¶ 110.

12 week dosing program, and also because Regeneron publically announced that

VEGF Trap was successful in quarterly doses.¹⁴⁸

Thus, the 2009 Press Release in view of Shams renders obvious "each tertiary dose is administered 12 weeks after the immediately preceding dose."

3. Dependent claims

a) Claim 2: The 2009 Press Release and Shams teach the claimed drug

The 2009 Press Release in view of Shams teaches "wherein the VEGF antagonist is aflibercept." The 2009 Press Release describes studies related to VEGF Trap-Eye, which is also called "aflibercept."¹⁴⁹

b) Claims 3 and 4: The 2009 Press Release and Shams teach the claimed modes of administration

The 2009 Press Release in view of Shams teaches "wherein all doses of the VEGF antagonist are administered to the patient by intraocular administration" (claim 3) and "the intraocular administration is intravitreal administration" (claim 4).¹⁵⁰ For example, the 2009 Press Release teaches: "In each study of the VIEW (VEGF Trap-Eye: Investigation of Efficacy and Safety in Wet AMD) program,

¹⁵⁰ *Id.* at 117.

¹⁴⁸ Ex. 1003 at ¶ 114 (citing Regeneron SEC Form 10-Q (May 4, 2007) at 17-18).

¹⁴⁹ Ex. 1003 at ¶ 116.

VEGF Trap-Eye is being evaluated for its effect on maintaining and improving vision when dosed as an intravitreal injection."¹⁵¹ Shams teaches "[t]he therapeutic compound for treatment of an intraocular neovascular disease is typically administered by ocular, intraocular, and/or intravitreal injection."¹⁵²

c) Claims 5-7: The 2009 Press Release and Shams teach the claimed dose amounts

The 2009 Press Release in view of Shams teaches "wherein all doses of the VEGF antagonist comprise from about 0.5 mg to about 2 mg of the VEGF antagonist" (claim 5), "wherein all doses of the VEGF antagonist comprise 0.5 mg of the VEGF antagonist" (claim 6), and "wherein all doses of the VEGF antagonist comprise 2 mg of the VEGF antagonist" (claim 7).¹⁵³ For example, the 2009 Press Release teaches: "In the first year of the studies, the safety and efficacy of VEGF Trap-Eye at doses of 0.5 mg and 2.0 mg administered at four week intervals and 2.0 mg at an eight-week dosing interval following one additional 2.0 mg dose at week four are being evaluated" and "VEGF Trap-Eye is also in Phase 2 development for the treatment of Diabetic Macular Edema (DME). VEGF Trap-Eye dosed at 0.5 mg or 2 mg monthly, 2 mg every eight weeks after three monthly

¹⁵¹ Ex. 1005 at 1, First Paragraph.

¹⁵² Ex. 1004, Page 25, Lines 15-16.

¹⁵³ Ex. 1003 at ¶ 118.

loading doses."¹⁵⁴ Shams also teaches administering the claimed doses (*see* Ground 1 above). Thus, the 2009 Press Release in view of Shams teaches the specific dosing regimens of claims 5-7.

d) Claims 8-11: The 2009 Press Release and Shams teach the disorders treated

The 2009 Press Release in view of Shams teaches "wherein the angiogenic eye disorder is selected from the group consisting of: age related macular degeneration, diabetic retinopathy, diabetic macular edema, central retinal vein occlusion, branch retinal vein occlusion, and corneal neovascularization" (claim 8), "wherein the angiogenic eye disorder is age related macular degeneration" (claim 9), "wherein the angiogenic eye disorder is diabetic retinopathy" (claim 10), and "wherein the angiogenic eye disorder is diabetic macular edema" (claim 11).¹⁵⁵ For example, the 2009 Press Release teaches "Phase 3 clinical trials evaluating VEGF Trap-Eye in the treatment of the neovascular form of age-related macular degeneration (wet AMD),"¹⁵⁶ and "VEGF Trap-Eye is also in Phase 2 development for the treatment of Diabetic Macular Edema (DME)."¹⁵⁷ Shams provides

¹⁵⁴ Ex. 1015 at 1, Fourth Paragraph and 2, Second Paragraph.

¹⁵⁵ Ex. 1003 at ¶ 119.

¹⁵⁶ Ex. 1015 at 1, First Paragraph.

¹⁵⁷ *Id.* at 2, Second Paragraph.

examples of "intraocular neovascular disease[s]" treatable by the disclosed VEGF antagonists, including "age-related macular degeneration (AMD), diabetic and other ischemia-related retinopathies, diabetic macular edema." ¹⁵⁸

VIII. GROUND 3: CLAIMS 1-11 FAIL TO SATISFY THE WRITTEN DESCRIPTION REQUIREMENT

Petitioner respectfully requests the Board cancel all claims of the '345 patent on the following Ground 3: Claims 1-11 fail the written description under pre-AIA 35 U.S.C. § 112, first paragraph and post-AIA 35 U.S.C. § 112(a).

The '345 patent fails to show Patent Owner's possession for all claims because the dosing regimen required by claim 1 is not supported. "The purpose of the written description requirement is to prevent an applicant from later asserting that he invented that which he did not; the applicant for a patent is therefore required to recount his invention in such detail that his future claims can be determined to be encompassed within his original creation." *Agilent Techs., Inc. v. Affymetrix, Inc.*, 567 F.3d 1366, 1383 (Fed. Cir. 2009) (quoting *Amgen Inc. v. Hoechst Marion Roussel Inc.,* 314 F.3d 1313, 1330 (Fed. Cir. 2003)). "In order to satisfy the written description requirement, the disclosure as originally filed does not have to provide *in haec verba* support for the claimed subject matter at issue." *Purdue Pharma L.P. v. Faulding Inc.,* 230 F.3d 1320, 1323 (Fed. Cir. 2000) (citing

¹⁵⁸ Ex. 1004, Page 21, Lines 1-6.

Fujikawa v. Wattanasin, 93 F.3d 1559, 1570 (Fed. Cir. 1996)). "Nonetheless, the disclosure 'must . . . convey with reasonable clarity to those skilled in the art that . . . [the inventor] was in possession of the invention." *Purdue*, 230 F.3d at 1323 (quoting *Vas-Cath Inc. v. Mahurkar*, 935 F.2d 1555, 1563-64, 19 U.S.P.Q.2D (BNA) 1111, 1117 (Fed. Cir. 1991) (alteration in original)).

The '345 patent fails to show possession of a dosing regimen with initial, secondary, and tertiary dosing where "each secondary dose is administered 4 weeks after the immediately preceding dose," and "each tertiary dose is administered 12 weeks after the immediately preceding dose" as required by all claims. "12 week" tertiary dosing is mentioned in two places in the '345 patent, but neither provides support for the claimed 4 week secondary doses and 12 week tertiary doses. The '345 patent's "Dosing Regimens" section lists myriad combinations of secondary and tertiary dosing and does not differentiate 4 week secondary dosing followed by 12–week tertiary dosing. Example 4 describes PRN tertiary dosing with an upper limit of 12 weeks, which Regeneron labelled insufficient during prosecution. Neither provides support for the claimed 4–week plus 12–week dosing regimen.

A. The '345 Patent's Disclosure of 12–Week Dosing

The '345 patent has little discussion of 12-week tertiary dosing. The 'Brief Summary' discusses tertiary doses, but it characterizes the inventor's discovery

broadly as tertiary dosing "once every 8 or more weeks."¹⁵⁹ The sole figure in the '345 patent describes tertiary dosing, but it is a fixed 8–week dosing regimen: "In this regimen, a single 'initial dose' of [VEGFT] is administered at the beginning of the treatment regimen (i.e. at 'week 0'), two 'secondary doses' are administered at weeks 4 and 8, respectively, and at least six 'tertiary doses' are administered once every 8 weeks thereafter."¹⁶⁰

When a 12–week tertiary dose is mentioned, the '345 patent includes it as one of 14 "or more" possible tertiary dosing frequencies: ¹⁶¹

In one exemplary embodiment of the present invention, each secondary dose is administered 2 to 4 (e.g., 2, 2¹/₂, 3, 3¹/₂, or 4) weeks after the immediately preceding dose, and each tertiary dose is administered at least 8 (e.g., 8, 8¹/₂, 9, 9¹/₂, 10, 10¹/₂, 11, 11¹/₂, **12**, 12¹/₂, 13, 13¹/₂, 14, 14¹/₂, or more) weeks after the immediately preceding dose.

This range has only one limit; the tertiary doses must be "at least 8 weeks." There is no upper bound on the range. Further, this disclosure is not limited to "each secondary dose is administered 4 weeks after the immediately preceding dose,"

¹⁵⁹ Ex. 1001, Col. 2:16.

¹⁶⁰ *Id.* at Col. 2:64-3:2.

¹⁶¹ Id. at Col. 3:57-62.

and "each tertiary dose is administered 12 weeks after the immediately preceding dose," as required by claim 1. Combinations of different tertiary doses, for example, are included in the scope of this description—the '345 patent explains that "each tertiary dose may be administered at the same frequency as the other tertiary doses" or, alternatively, the frequency at which "tertiary disease are administered to a patient can vary over the course of the treatment regimen."¹⁶²

When the '345 patent next mentions a 12–week dose frequency, the tertiary 12–week dose(s) is preceded by four 8–week "tertiary doses" and 12 weeks is again only one choice, among many, for the tertiary dose: ¹⁶³

[E]ach secondary dose may be administered to the patient 4 weeks after the immediately preceding dose. ... [F]ollowed by at least 5 tertiary doses of the VEGF antagonist, wherein the first four tertiary doses are administered 8 weeks after the immediately preceding dose, and wherein each subsequent tertiary dose is administered from 8 to 12 (e.g., 8, $8\frac{1}{2}$, 9, $9\frac{1}{2}$, 10, $10\frac{1}{2}$, 11, $11\frac{1}{2}$, 12) weeks after the immediately preceding dose. The frequency of administration may also be adjusted during the course of treatment by a physician depending on the needs of the individual patient following clinical examination.

¹⁶² *Id.* at Col. 4:23-34.

¹⁶³ Id. at Col. 23-43 (emphasis added).

Because the tertiary dosing in this description includes 8–week dosing followed by 12–week dosing, it does not describe "each tertiary dose is administered 12 weeks after the immediately preceding dose."¹⁶⁴

The remainder of the '345 patent includes no discussion of a dosing regimen where "each tertiary dose is administered 12 weeks after the immediately preceding dose." After the "Dosing Regimens" section, the patent continues with listing VEGF antagonists,¹⁶⁵ angiogenic eye disorders,¹⁶⁶ pharmaceutical formulations,¹⁶⁷ modes of administration, ¹⁶⁸ VEGF dosing amounts, ¹⁶⁹ and

¹⁶⁵ Ex. 1001, Col. 4:47-5:20.

- ¹⁶⁷ *Id.* at Col. 5:40-6:7.
- ¹⁶⁸ *Id.* at Col. 6:8-24.

¹⁶⁴ This discussion in the '345 patent corresponds to Claim 5 of the parent PCT Application. That claim does not teach "sequentially administering" secondary doses and tertiary doses of the same frequency for the same reasons as the corresponding paragraph in the '345 patent.

¹⁶⁶ *Id.* at Col. 5:21-39.

¹⁶⁹ *Id.* at Col. 6:25-7:25.

treatment population and efficacy.¹⁷⁰ These sections do not mention 12–week dosing frequencies.

The patent then describes seven examples. Examples 1 and 2 correspond to Regeneron's Phase 1 and 2 Aflibercept trials (described in Regeneron's 2006 and 2007 press releases¹⁷¹) and do not include any tertiary dosing.¹⁷² Example 3 describes a Phase 1 trial of a single dose, and similarly does not include any tertiary dosing.¹⁷³ Example 4 corresponds to the Phase 3 clinical trial (described in Regeneron's 2009 Press Release¹⁷⁴) and describes tertiary dosing, including an 8–week fixed tertiary dose and PRN tertiary dosing with a maximum of 12–weeks; this cannot provide support for the claimed dosing regimen because Regeneron argued during prosecution that the Press Release's identical disclosure does not teach "each tertiary dose is administered 12 weeks after the immediately preceding dose."¹⁷⁵ Example 5 corresponds to a Phase 2 clinical trial in diabetic macular

- ¹⁷⁰ Id. at Col. 7:26-44.
- ¹⁷¹ Ex. 1003 at ¶¶ 74-75.
- ¹⁷² Id. at ¶ 76.

¹⁷³ Id..

¹⁷⁴ Id. at ¶ 77.

¹⁷⁵ Ex. 1002, Response, 03/16/2020, Pages 4-5.

edema, respectively, and, like Example 4, describes tertiary dosing but is limited to administering each tertiary dose 8 weeks after the immediately preceding dose. Example 6 describes a Phase 3 dosing study in central retinal vein occlusion and does not include any tertiary dosing. Example 7 lists 20 "examples of dosing regimens within the scope of the present invention."¹⁷⁶ Although Example 7 discloses tertiary dosing, the dosing frequency is described as either "once every 8 weeks," "less frequent" than the secondary dosing, or PRN.¹⁷⁷ None of the 20 exemplary dosing regimens provided in Example 7 include a 12–week tertiary dose.¹⁷⁸

B. Regeneron's Discussion of 12-week Dosing During Prosecution

Regeneron overcame a double patenting rejection by arguing, in part, that the '345 patent was non-obvious over Patent Owner's earlier patents.¹⁷⁹ Specifically, Regeneron argued that the "standard of care for the treatment of [AMD] was to administer an antibody formulation (ranibizumab) by injection to

¹⁷⁷ Id.

¹⁷⁸ Id.

¹⁷⁹ Ex. 1002, Response, 06/28/2019.

¹⁷⁶ Ex. 1003 at ¶ 80.

the eye once per month^{"180} and characterized a paper by Heier¹⁸¹ as "showing improved unexpected results" that supports nonobviousness of the claimed 12– week tertiary dosing.¹⁸² According to Patent Owner, "the PRN treatment protocol [disclosed in Heier] as encompassed by . . . the 12–week dosing of claim [1] achieves results which would be surprisingly as good or better than the results obtained with monthly treatment."¹⁸³ Equating Heier with claim 1, Regeneron stated that "the Heier *et al.* results suggest that by administering the VEGF antagonist in accordance with a dosage regimen as claimed in independent [claim 1], it is possible to treat angiogenic eye disorder…on a less frequent basis than previously thought possible."¹⁸⁴

Heier published in December 2012 and describes a study to determine effect on "neovascular age-related macular degeneration (AMD) compared monthly and every-2-month dosing of intravitreal aflibercept injection . . . with monthly

¹⁸¹ Id.

¹⁸² Ex. 1002, Response, 06/28/2019 (citing 1012.)

¹⁸³ Id. at 9.

¹⁸⁴ Id.

 $^{^{180}}$ Id.

ranibizumab.^{*185} Heier states that all patients "received a minimum of dosing every 12 weeks with interim as-needed monthly intravitreal injections.^{** 186} As Regeneron indicated, Heier teaches the same regimen as Example 4 of the '345 patent.¹⁸⁷

In the next action, the PTO withdrew the double patenting rejection and added a rejection of (now) claims 1-11 as anticipated by the 2009 Press Release.¹⁸⁸ The PTO correctly described the Press Release as "teach[ing] that patients received/intravitreal doses of 0.5 mg or 2g VEGF Tap-Eye [sic] at 4–week intervals in the first year, followed by continual treatment for another year on a flexible, PRN regiment, with a dose administered at least every 12 weeks."¹⁸⁹ The 2009 Press Release has essentially the same description as Heier, the journal article that Regeneron relied on to overcome the double patenting rejection, and Example 4 of the '345 patent.

¹⁸⁷ Ex. 1002, Response, 06/28/2019, Page 8.

¹⁸⁸ Ex. 1002, Rejection, 10/01/2019, Pages 4-5.

 189 Id.

¹⁸⁵ Ex. 1012 at 2537.

¹⁸⁶ *Id.* at 246.

In response to the 2009 Press Release. Regeneron changed its characterization of 12-week dosing regimens, now arguing that as-needed tertiary dosing between 4 and 12 weeks does not explicitly disclose a tertiary 12-week dosing frequency component because "[a] practitioner of ordinary skill in the art would not have understood a PRN dosing regimen with 4 and 12-week limits as encompassing a [12-week dosing] regimen.¹⁹⁰ Regeneron also argued that the claims were not inherently disclosed in the Press Release because "[t]hough the Press Release discussed a PRN dosing regimen wherein a dose interval may extend out as far as 12 weeks, the dosages administered to patients were not necessarily this infrequent. For this reason, the Press Release was insufficient as an inherently anticipating reference."¹⁹¹ That exact dosing regimen is taught by Heier, the journal article that Regeneron cited to overcome the double patenting rejection by proving "unexpected results." Regeneron did not attempt to reconcile the inconsistency between its reliance on Heier (to overcome the first rejection) as sufficiently disclosing the claimed regimen to support unexpected results with its later critique of the 2009 Press Release as insufficient to disclose that same regimen.

¹⁹⁰ Ex. 1002, Response, 01/23/2020, Pages 6-8.

¹⁹¹ *Id.* (emphasis in original).

An examiner interview followed on March 6, 2020.¹⁹² Later that month, Regeneron filed a supplemental response presenting more arguments to distinguish claims 32-42 (issued claims 1-11) from the 2009 Press Release:¹⁹³

> Claims 32-42 relate to a method requiring tertiary dosing administered 12 weeks after the immediately preceding dose. There is a single appearance of the words "12 weeks" within the fourth paragraph of the Press Release. However, this paragraph is referring to a "flexible, criteria-based extended PRN regimen with a dose administered at least every 12 weeks, but not more often than every four weeks". As explained in our January 23, 2020 Response, this is not a disclosure of a regimen having 12– week tertiary dosing as specified in the claims. Mere mention of a prospective possibility of dosing at 12 weeks does not specifically indicate or teach towards a method where 12–week dosing would be undertaken, let alone successful.

The Examiner explicitly relied on this argument to allow claims 32-42,¹⁹⁴ which issued on November 10, 2020, as claims 1-11 of the '345 patent.¹⁹⁵

¹⁹³ Id.

¹⁹⁴ Ex. 1002, Notice of Allowance, 04/01/20, Pages 2-3.

¹⁹⁵ Ex. 1001, Cover Page.

¹⁹² Ex. 1002, Response, 03/16/2020, Page 4.

C. The '345 Patent Lacks Written Description Support for the Same Reasons Regeneron Articulated in its Critique of the 2009 Press Release

Before the PTO, Regeneron argued that the 2009 Press Release's teaching (which corresponds to Example 4)¹⁹⁶ of 12–week tertiary dosing was insufficient to teach a 12–week dosing regimen: ¹⁹⁷ "Mere mention of a prospective possibility of dosing at 12 weeks . . . does not specifically indicate or teach toward a method where 12–week dosing would be undertaken, let alone successful." ¹⁹⁸ This reasoning also applies to claim 1 and demonstrates that the claim lacks written description support.

The 2009 Press Release states that "[a]fter the first year of treatment [of 4– week secondary doses], patients will continue to be followed and treated for another year on a flexible, criteria-based extended PRN regimen with a dose administered at least every 12 weeks, but not more often than every four weeks." Example 4 discloses the same regimen: "During the second year of the study, . . . [tertiary doses] may be given as frequently as every 4 weeks, but no less frequently

¹⁹⁸ Id.

¹⁹⁶ Ex. 1003 at ¶¶ 77-78.

¹⁹⁷ Ex. 1002, Response to Office Action 03/16/2020, Page 4.

than every 12 weeks."¹⁹⁹ Given that the '345 patent and the 2009 Press Release disclose the same tertiary dosing possibilities, and that Regeneron critiqued the 2009 Press Release (to thereby obtain allowance of the '345 patent) as insufficient to teach a 12–week dosing regimen, the '345 patent's disclosure is also insufficient to teach "each tertiary dose is administered 12 weeks after the immediately preceding dose."

D. The '345 Patent's Undifferentiated Disclosure of Various Dosing Regimens Is Insufficient to Support a Claim to a Specific 4-Week Secondary and 12-Week Tertiary Dosing Regimen

In *Novozymes*, the Federal Circuit held that an "application's undifferentiated description" of a specific invention is insufficient unless the disclosure "provide[s] sufficient 'blaze marks' to guide a reader through the forest of disclosed possibilities towards the claimed" elements. 723 F.3d at 1346 (quoting *In re Ruschig*, 379 F.2d 990, 994-95 (C.C.P.A. 1967)). Thus, "one cannot disclose a forest in the original application, and then later pick a tree out of the forest and say here is my invention." *Purdue Pharma L.P. v. Faulding Inc.*, 230 F.3d 1320, 1326 (Fed. Cir. 2000); *see also Boston Sci. Corp. v. Johnson & Johnson*, 647 F.3d 1353, 1367 (Fed. Cir. 2011); *Fujikawa*, 93 F.3d at 1570-71.

¹⁹⁹Ex. 1001, Col. 10:12-14.

In *Purdue Pharma*, the claims required a method of administering an opioid so that the patient had specific plasma concentrations of the opioid at specific times. 230 F.3d at 1324. The original application described seven examples of administering an opioid, two of which included the required concentrations at the required times. *Id.* at 1326. But all seven examples also described numerous other parameters related to administering the opioid. *Id.* And nothing in the application's description suggested to one skilled in the art that the later-claimed concentrations (added during prosecution) were "an important defining quality" for the invention. *Id.* at 1327. Rather, the patentees appeared to have simply "pick[ed] a characteristic possessed by" some examples in the original application—a "characteristic that is not discussed even in passing in the disclosure." *Id.* The court explained that this was "exactly the type of overreaching the written description requirement was designed to guard against." *Id.*

Like *Purdue Pharma*, claim 1 of the '345 patent is specific—a 4-week secondary dosing frequency followed by a 12-week tertiary dosing frequency. And like *Purdue Pharma*, the '345 patent "discloses a forest in the original application, and [Patent Owner] then later pick[ed] a tree out of the forest and [said] here is my invention." *Purdue Pharma*, 230 F.3d at 1326. The '345 patent mentions 12-week tertiary dosing, but does so in an unbounded range of dosing

frequencies, with the only limit being that each tertiary dose must be "at least 8 weeks" frequency: "each tertiary dose is administered at least 8 (e.g., 8, 8¹/₂, 9, 9¹/₂, 10, 10¹/₂, 11, 11¹/₂, 12, 12¹/₂, 13, 13¹/₂, 14, 14¹/₂, or more)."²⁰⁰ This disclosure is much broader than claim 1's required "each secondary dose is administered 4 weeks after the immediately preceding dose," and "each tertiary dose is administered 12 weeks after the immediately preceding dose." The '345 patent explicitly teaches that the different secondary and tertiary doses can be administered at the same or different frequencies.²⁰¹ The '345 patent's combinations of secondary $(2, 2\frac{1}{2}, 3, 3\frac{1}{2}, or 4$ weeks) and tertiary doses (e.g., 8, $8\frac{1}{2}$, 9, $9\frac{1}{2}$, 10, $10\frac{1}{2}$, 11, $11\frac{1}{2}$, 12, $12\frac{1}{2}$, 13, $13\frac{1}{2}$, 14, $14\frac{1}{2}$, or more) therefore allows for secondary dosing at varying frequencies and tertiary dosing at varying frequencies, yielding unlimited combinations of secondary and tertiary dosing frequencies. For example, assume that a secondary/tertiary dosing regimen consisted of two secondary doses followed by three tertiary doses and that the '345 patent described just five possible secondary dose frequencies and 14 possible

²⁰⁰ Ex. 1001, Col. 3:60-62.

²⁰¹ Id. at Col. 4:23-34.

tertiary dose frequencies (i.e., ignore the "or more" tertiary doses), the possible dosing regimens gives 68,000²⁰² possible combinations.

Prosecution of the '345 patent reinforces that that the number of treatment options is vast. In overcoming the double patenting rejection, Regeneron described the treatment options as "virtually infinite:"²⁰³

There are virtually an infinite number of different treatment protocols that could be tested. A drug could be administered more frequently, or less frequently, relative to the accepted standard of care. Further, different variations in timing between dosing events are possible. Due to the virtually infinite number of combinations, applicants do not believe that the claimed treatment protocol is *prima facie* obvious in view of the prior art standard of care which is administration of the drug once per month.

Because the '345 patent also discloses a "virtually infinite" number of undifferentiated dosing regimens, the specific regimen of claim 1 is not reasonably supported by the disclosure.

three doses. 5x5x14x14x14=68,600.

²⁰³ Ex. 1002, Response, 06/28/2019.

²⁰² Five choices for each of the first two doses and 14 choices for each of the last

Even if the '345 patent's vast disclosure had been limited to secondary doses of the same frequency and tertiary doses of the same frequency (which it is not). the number of combinations are still too many for one of skill in the art to recognize possession of the specific invention. In the '345 patent, the 4-week secondary dosing is one of five explicit options and the 12-week dosing is one of 14 "or more" options, yielding 70 or more combinations for the claimed initial, secondary, and tertiary dosing frequencies. In *Ruschig*, the claim at issue was directed to a single compound. 379 F.2d at 994-95. The examiner there found the specification yielded over 1,000 combinations encompassing the specific claim, but the Patent Owner argued that the total number was 46. The Federal Circuit did not find this persuasive, holding: "Specific claims to single compounds require reasonably specific supporting disclosure and while . . . naming is not essential, something more than the disclosure of a class of 1000, or 100, or even 48, compounds is required." Id. at 994. In the same way here, even if the '345 patent disclosed just 70 combinations, something more is required to reasonably support the specifics of claim 1. Such a "laundry list" disclosure "would not 'reasonably lead' those skilled in the art to any particular" dosing frequency. Fujikawa, 93 F.3d at 1571; see also FWP IP ApS v. Biogen MA, Inc., 749 Fed. Appx. 969, 973 (Fed. Cir. 2018) (unpublished) (finding a specific dose of 480 mg unsupported

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where the specification only mentioned 480 mg three times, twice in a paragraph listing possible doses).

The '345 patent is different than those cases where the Federal Circuit found support for a claimed invention where the specification described the invention among various combinations. In those cases, the court found that one of skill in the art would recognize possession of the invention based on background knowledge in the art. Those cases cannot save the '345 patent and be consistent with Regeneron's prosecution arguments. For example, Regeneron argued during prosecution that one of skill in the art would understand a non-monthly tertiary dosing schedule to be the recognized option in the art: "At the time of the invention the standard of care for the treatment of the neovascular (or wet) form of age-related macular degeneration (AMD) was to administer an antibody formulation (ranibizumab) by injection to the eye once per month."²⁰⁴ As evidenced by Regeneron's statement in prosecution, one of skill in the art would not look at the undifferentiated lists of dosing frequencies and recognize possession of the "non-standard" specific regimen disclosed in claim 1.

For the above reasons, the '345 patent is invalid for failing the written description requirement of 35 U.S.C. § 112(a).

²⁰⁴ Ex. 1002, Response, 06/28/2019, at 7.

IX. THE BOARD SHOULD NOT EXERCISE DISCRETIONARY DENIAL UNDER 35 U.S.C. § 325(D)

Section 325(d) is inapplicable to this proceeding because the Petition does not raise substantially the same art or arguments in the same way as the examination of the '345 patent and, to the extent the Petition does, the Office erred in a material manner. *Advanced Bionics, LLC v. MED-EL Elektromedizinische Geräte GmbH*, IPR2019-01469, Paper 6, at 7-11 (P.T.A.B. Feb. 13, 2020) (precedential).

A. Shams Was Not Considered on the Record, Is Not Cumulative of any Reference Considered on the Record, and, Even if It Was Considered, the Office Materially Erred by Allowing the '345 Patent over Shams

"The Board has consistently declined exercising its discretion under Section

325(d) when the only fact a Patent Owner can point to is that a reference was

disclosed to the Examiner during the prosecution." Amgen Inc. v. Alexion

Pharma., Inc., IPR2019-00740, Paper 15 at 65-66 (P.T.A.B. Aug. 30, 2019).

Although Regeneron identified Shams in an Information Disclosure Statement

("IDS")²⁰⁵, the Examiner did not apply Shams in an anticipation or obviousness

rejection. Like the patent owner in *Amgen*, Regeneron here can only point to an IDS.

²⁰⁵ Ex. 1002, IDS filed 2019-06-09.

Shams is not cumulative of any art considered during prosecution. The Office issued two principal rejections of claim 1 during prosecution—obviousness-type double patenting and anticipation by the 2009 Press Release—and Regeneron overcame the rejections by arguing that the prior art was limited to monthly dosing or the prior art did not specifically teach 12 week tertiary dosing. Crediting Regeneron's arguments, the Office withdrew both rejections because no reference taught 4 week dosing of VEGF antagonist followed by 12–week dosing; Shams teaches this purportedly novel dosing regimen. Thus, Shams 12–week dosing is not cumulative to any reference discussed on the record.

Even if the Office had considered Shams, "the Office erred in a manner material to the patentability of challenged claims." *Advanced Bionics, LLC v. MED-EL Elektromedizinische Geräte GmbH*, IPR2019-01469, Paper 6, at 8 (P.T.A.B. Feb. 13, 2020) (precedential). While the Examiner may have considered Shams in an IDS, the touchstone is "the extent of such consideration" and whether there is "evidence of record indicating why the Examiner" did not reject the claims. *William Hill US Holdco, Inc. v. CG Tech. Dev.*, LLC, IPR2019-00317, Paper 14 at 35 (P.T.A.B. May 30, 2019). There is no evidence on the record that the Examiner here considered Shams substantively. "[I]f the record of the Office's previous consideration of the art is not well developed or silent, then a petitioner may show the Office erred by overlooking something persuasive." *Advanced*

Bionics at 10. Assuming the Examiner did consider Shams substantively but did not include the analysis on the record, the materiality of Shams—and the error in Office's '345 prosecution—is confirmed by prosecution in a third-party application. In U.S. Application No. 14/934,731, (the "'731 application"), Novartis filed claim 1 to cover "three individual doses of a VEGF antagonist at 4– week intervals" followed by "an additional dose of the VEGF antagonist once . . . every 12 weeks (q12 regimen) if [certain] criteria are not met."²⁰⁶ The Examiner in the '731 application rejected the claimed dosing regimens as anticipated by the European national stage application (EP 2311433) of Shams' PCT, comparing the '731 application's dosing regimen to Shams' disclosure:²⁰⁷

> [Shams] teaches methods for administering a mammal suffering from, or at risk for, an intraocular neovascular disorder with regular dosing of a therapeutically effective amount of VEGF antagonist, followed by less frequent dosing of a therapeutically effective amount of VEGF antagonist (abstract). The methods include administering to a mammal a number of first individual doses of a VEGF antagonist (ranibizumab), followed by administering to the mammal a number of second individual doses of the antibody, while the second individual doses are

²⁰⁶ Ex. 1014, claims filed 11-06-2015.

²⁰⁷ Ex. 1014, Non-Final Rejection mailed 11-14-2016 (citing EP 2311433).

administered less frequently. The mammal in need of may be a human. The administration of the VEGF antagonist is intravitreal. The first individual doses are administered at one month intervals (e.g., about 3 individual doses). In another embodiment the second individual doses are administered at three month intervals (e.g., about 6 individual doses).

To the extent the Office considered Shams in the '345 patent's prosecution, the Office materially erred for not rejecting the '345 patent claims under Shams as shown by the Office's analysis of the same dosing regimen in the '731 application.

For all the foregoing reasons, the Board should decline to exercise discretion under § 325(d) for Ground 1, the '345 Patent is anticipated by Shams.

B. The 2009 Press Release in View of Shams Was Not Considered on the Record nor Were any Similar Arguments Considered

"The Board frequently holds that a reference that was neither applied against the claims nor discussed by the Examiner does not weigh in favor of exercising the Board's discretion under § 325(d) to deny a petition." *Amazon Inc. v. M2M Sols. LLC, IPR2019-01204*, Paper 14 at 17 (P.T.A.B. Jan. 23, 2020) (internal quotations omitted). As discussed above, Shams was not considered substantively on the record nor cumulative of any reference discussed on the record. Thus, the 2009 Press Release in view of Shams was not considered, and the Board should not exercise its discretion under § 325(d) to deny the present petition.

Further, the Office only considered the 2009 Press Release's 4 week plus 12 week PRN dosing regimen on the record; the press release's 4 week plus 8 week dosing regimen—relied upon in Ground 2—was never discussed. In the '345 prosecution, the third party submission only raised the 2009 Press Release's 4 week and 12 week PRN dosing²⁰⁸ and only as a 102 argument; the Examiner did likewise. There is no evidence to suggest that the Office considered the 2009 Press Release's 4 week plus 8 week dosing regimen, much less consider modifying that dosing regimen to a 4 week plus 12 week dosing regimen. Thus, not only is the 2009 Press Release in view of Shams newly presented because Shams was not previously considered, but also because the 2009 Press Release was not previously considered on the record as a 103 reference nor was the 2009 Press Release's 4 week plus 8 week dosing regimen considered in a rejection.

For all the foregoing reasons, the Board should decline to exercise discretion under § 325(d) for Ground 2, the '345 Patent is obviousnes over the 2009 Press Release in view of Shams.

²⁰⁸ The third party submission introduced evidence from a second reference, Dixon, as evidence that the Press Release's "VEGF-Trap" was the claimed VEGF antagonist and, more particularly, Regeneron's aflibercept.

C. No Written Description Arguments Were Considered on the Record

The Office issued no rejections under 35 U.S.C. § 112 during prosecution of the '345 patent. Further, the Office never raised the 12 week tertiary dosing plan as lacking support. Because no 12–week dosing rejections or similar arguments were raised during prosecution, the Board should decline to deny Ground 3 under § 325(d).

X. MANDATORY NOTICES

A. Real Party-in-Interest

Pursuant to 37 C.F.R. § 42.8(b)(1), the real parties-in-interest in this proceeding are Chengdu Kanghong Biotechnology Co., Ltd. (Petitioner), Chengdu Kanghong Pharmaceutical Group Co., Ltd. (the parent company of Petitioner), and Beijing Kanghong Biomedical Co., Ltd. (a wholly-owned subsidiary of Petitioner's parent company). No other party has funded or exercises control over this Petition.

B. Related Matters Under 37 C.F.R. § 42.8(b)(2)

Petitioner is unaware of any related federal court or PTAB proceedings.

C. Lead and Back-Up Counsel and Service Information

Pursuant to 37 C.F.R. § 42.8(b)(3), Petitioner identifies the following counsel (and a power of attorney accompanies this Petition).

Lead Counsel for Petitioner	Backup Counsel for Petitioner
Matthew I. Kreeger	Jian Xiao
mkreeger@mofo.com	JXiao@mofo.com
Registration No.: 56,398	Registration No.: 55,748
MORRISON & FOERSTER LLP	MORRISON & FOERSTER LLP
425 Market Street	755 Page Mill Road
San Francisco, CA 94105	Palo Alto, CA 94304-1018
Tel: (415) 268-6467	Tel: (650) 813-5736
Fax: (415) 268-7522	Fax: (650) 494-0792
	Desmond O'Sullivan
	dosullivan@mofo.com
	Registration No.: 67,576
	MORRISON & FOERSTER LLP
	12531 High Bluff Drive, Suite 100
	San Diego, CA 92130
	Tel: (858) 314-7794
	Fax: (858) 720-5125

Pursuant to 37 C.F.R. § 42.8(b)(4), service information for lead and back-up counsel is provided above. Petitioner consents to electronic service by email to

CHENGDU-PGR@mofo.com.

XI. CONCLUSION

Because there is a reasonable likelihood that Kanghong will prevail on at least one of its asserted grounds with respect to at least one claim, Kanghong respectfully requests that the Board institute post-grant review of claims 1–11 of the '345 patent. Upon review, claims 1–11 should be held unpatentable.

The PTO is authorized to charge any required fees, including the fee as set

forth in 37 C.F.R. § 42.15(a) and any excess claim fees, to Deposit Account

No. 03-1952 referencing Docket No. 77688-00000.15.

Respectfully submitted,

Dated: January 7, 2021

/Matthew I. Kreeger/

By: Matthew I. Kreeger mkreeger@mofo.com Registration No.: 56,398 MORRISON & FOERSTER LLP 425 Market Street San Francisco, CA 94105 Tel: (415) 268-6467 Fax: (415) 268-7522

Counsel for Petitioner

CERTIFICATION OF WORD COUNT UNDER 37 C.F.R. § 42.24(d)

The undersigned hereby certifies that the foregoing petition contains 14,087 words according to the word count of the word-processing software used to prepare the petition, excluding the table of contents, listing of exhibits, mandatory notices under 37 C.F.R. § 42.8, certificate of service, and certificate of word count.

Dated: January 7, 2021

/Matthew I. Kreeger/

Matthew I. Kreeger Registration No.: 56,398

CERTIFICATE OF SERVICE (37 C.F.R. § 42.24)

I hereby certify that the attached Petition for Post Grant Review and

supporting materials were served as of the below date by UPS, which is a means at

least as fast and reliable as U.S. Express Mail, on the Patent Owner at the

correspondence address indicated for U.S. Patent No. 10,828,345:

Regeneron – Bozicevic, Field & Francis 201 Redwood Shores Parkway Suite 200 Redwood City, CA 94065

Dated: January 7, 2021

/Matthew I. Kreeger/

By: Matthew I. Kreeger mkreeger@mofo.com Registration No.: 56,398 MORRISON & FOERSTER LLP 425 Market Street San Francisco, CA 94105 Tel: (415) 268-6467 Fax: (415) 268-7522

UNITED STATES PATENT AND TRADEMARK OFFICE

BEFORE THE PATENT TRIAL AND APPEAL BOARD

CHENGDU KANGHONG BIOTECHNOLOGY CO., LTD. Petitioner

V.

REGENERON PHARMACEUTICALS, INC. Patent Owner

Case PGR2021-00035 Patent 10,828,345

PRELIMINARY RESPONSE OF PATENT OWNER REGENERON PHARMACEUTICALS, INC.

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I. INTRODUCTION

Patent Owner Regeneron Pharmaceuticals, Inc. ("Regeneron") is an innovative U.S. biotechnology company that invents life-changing medicines for people with serious diseases. Regeneron was founded and has been led for more than 30 years by physician-scientists and has developed nine FDA-approved medicines, including EYLEA®. The active agent in EYLEA®, aflibercept, is a novel fusion protein developed by Regeneron that binds to and neutralizes Vascular Endothelial Growth Factor (VEGF), a key contributor to angiogenesis. By binding and neutralizing VEGF, aflibercept is able to prevent blood vessel leakage and block the growth of abnormal blood vessels in the back of the eye and effectively treat angiogenic eye disorders. Since its approval by FDA in 2011, EYLEA® has revolutionized the treatment of angiogenic eye disorders including age-related macular degeneration (AMD), macular edema, and diabetic retinopathy.

Before the development of EYLEA®, the standard of care for treatment of angiogenic eye disorders was monthly intravitreal injections of ranibizumab (an anti-VEGF antibody fragment) or off-label use of bevacizumab (an anti-VEGF antibody). The great treatment burden of monthly eye injections led to extensive efforts in the art to decrease injection frequency and physician monitoring. Ex. 1012 at 1, 9. However, before EYLEA, fixed quarterly or "as needed" (*pro re nata*) dosing regimens with existing VEGF inhibitors, without monthly monitoring visits, were not effective at maintaining vision. Ex. 1012 at 1.

Regeneron's Phase III clinical trial results surprisingly demonstrated "remarkably similar improvement in vision and anatomic measures can be achieved" with less frequent dosing of aflibercept as compared to monthly injections of ranibizumab. Ex. 1012 at 10-11. Indeed, the Examiner relied on this evidence of unexpected results during prosecution of U.S. Patent No. 10,828,345 (the "345 Patent"). Not surprisingly, given the long-felt need and repeated failures in the art to reduce treatment burden and injection frequency, EYLEA has enjoyed great commercial success.

Petitioner Chengdu Kanghong Biotechnology Co., Ltd.'s ("Kanghong" or "Petitioner") seeks to capitalize on Regeneron's hard-earned success by commercializing conbercept, a "me too" fusion protein, in the United States.¹ Petitioner seeks to invalidate Regeneron's '345 Patent claims to extended (12week) dosing regimens for treating angiogenic eye disorders using the claimed VEGF antagonist fusion proteins, by arguing that Regeneron's claims are

¹ Petitioner seeks to invalidate the '345 Patent even before the safety and efficacy of its infringing conbercept product has been demonstrated. Indeed, Petitioner recently suspended one of its two Phase III pivotal clinical trials for conbercept in the United States based on a mid-term review of data generated in the study. In addition, the National Agency for the Safety of Medicines and Health Products of France recently halted a Phase III trial of conbercept in Europe. *See* Ex. 2032; Ex. 2033; Ex. 2034.

anticipated or obvious based on a prior art dosing regimen, and are not adequately described. However, the Shams prior art dosing regimen on which Petitioner relies was an acknowledged failure and Petitioner assiduously avoids any discussion of Regeneron's demonstration of unexpected results in prosecution, on which the Examiner relied in allowing the '345 Patent. Moreover, Petitioner overlooks the fact that the dosing regimen claimed is described as a specific example in the specification.

Patent Owner submits this preliminary response pursuant to 35 U.S.C. § 323 and 37 C.F.R. § 42.207 to Petitioner's request for post-grant review ("PGR") of Claims 1-11 of the '345 Patent, Ex. 1001. This preliminary response is timely filed within three months of the Patent Trial & Appeal Board's ("Board") notice (Paper No. 3), mailed January 15, 2021, indicating that the Petition was accorded a filing date. As set forth herein and in the accompanying exhibits, the Petition should be denied for at least the following reasons:

First, the '345 Patent is not eligible for PGR because its effective filing date is before March 16, 2013.

Second, the prior art asserted by Petitioner in Grounds 1 and 2, and the §112 disclosure challenged in Ground 3, were previously before the Examiner, and Petitioner has not sufficiently alleged that the Examiner erred in a manner material to the patentability of the challenged claims in considering the art and arguments,

warranting discretionary denial under 35 U.S.C. § 325(d). See Pharmacosmos A/S v. Am. Regent, Inc., PGR2020-00009, Paper 17 at 27-28 (Aug. 14, 2020).

Third, Petitioner has failed to meet its threshold burden under 35 U.S.C. §§ 324(a) and 322(a)(3) to show that it is more likely than not that at least one of the claims of the '345 Patent is unpatentable because (1) Shams does not anticipate the '345 Patent claims, (2) Shams and the 2009 Press Release do not render the '345 Patent claims obvious, and (3) the '345 Patent claims are adequately supported by the pre-March 16, 2013 priority applications.

II. BACKGROUND

A. The '345 Patent Claims

The '345 Patent's single independent claim, Claim 1, recites a method for treating an angiogenic eye disorder in a patient by administering a single dose of a VEGF antagonist followed by one or more secondary doses that are administered four weeks after the preceding dose, followed by tertiary or maintenance doses that are administered twelve weeks apart. Ex. 1001 at 21:55-22:56. Claim 1 also recites that the claimed VEGF antagonist is a receptor-based chimeric molecule comprising an immunoglobulin-like (Ig) domain 2 of VEGF receptor Flt1, Ig domain 3 of the VEGF receptor Flk1, and a multimerizing component. *Id.* In other words, Claim 1 recites a method of treating an angiogenic eye disorder by administering a recited

VEGF antagonist fusion protein at a twelve-week dosing regimen following an initial set of doses administered four weeks apart. *Id.*

The '345 Patent has ten dependent claims, Claims 2-11. Claim 2 recites that the particular VEGF antagonist fusion protein is affibercept, the fusion protein in Regeneron's Eylea[®] product. *Id.* at 22:56-57. Claims 3 and 4 recite that the VEGF antagonist fusion protein is administered intraocularly and intravitreally, respectively. *Id.* at 22:58-62. Claim 5 recites administering 0.5 to 2 mg, Claim 6 recites administering 0.5 mg, and Claim 7 recites administering 2 mg of VEGF antagonist. *Id.* at 22:63-23:2. Claim 8 recites that the angiogenic eye disorder is one of: age related macular degeneration (also known as "wet AMD"), diabetic retinopathy, diabetic macular edema, central retinal vein occlusion, branch retinal vein occlusion, and corneal neovascularization; Claim 9 recites that the angiogenic eye disorder is diabetic retinopathy and Claim 11 recites that the angiogenic eye is diabetic macular edema. *Id.*

III. THE LEVEL OF ORDINARY SKILL IN THE ART

For purposes of this Preliminary Response, Patent Owner has used Petitioner's definition of the person of ordinary skill in the art ("POSA"). Paper 2 at 16. Patent Owner reserves the right to propose another definition if this post-grant review is instituted.

IV. CLAIM CONSTRUCTION

The claims in a post-grant review are to be construed using the same standard that applies in district court proceedings, which is set forth in *Phillips v. AWH Corp.*, 415 F.3d 1303 (Fed. Circ. 2005) (*en banc*); see Changes to the Claim Construction Standard for Interpreting Claims in Trial Proceedings Before the Patent Trial and Appeal Board, 83 Fed. Reg. 51,340 (Oct. 11, 2018) (applicable to post-grant reviews filed on or after November 13, 2018).

Petitioner's challenge should be disposed of under 35 U.S.C. § 325. However, should the Board consider it necessary to decide whether Petitioner satisfied its threshold burden under 35 U.S.C. § 324, the proposed claim constructions are relevant to Petitioner's Ground 1 and 2 challenges.² As explained below, Patent Owner respectfully submits that "A method for treating an angiogenic eye disorder in a patient" is a positive limitation of Claim 1 that requires a therapeutically effective method for treating an angiogenic eye disorder and that the term "tertiary dose" means "dose(s) that maintain(s) a therapeutic

² Petitioner did not propose any constructions and, indeed, argued that no construction is required for any of the terms used in the '345 Patent. Paper 2 at 16-17.

effect throughout the course of treatment."3

A. Claim 1's Preamble Is a Positive Limitation That Requires A Therapeutically Effective Method of Treatment

The preamble of Claim 1 — "A method of treating an angiogenic eye disorder in a patient" — is limiting because it breathes life and meaning into the claim. Further, it provides an antecedent basis for terms in the body of the claim and dependent claims.

The preamble of Claim 1 gives the claim life and meaning. *See, e.g., Griffin v. Bertina*, 285 F.3d 1029, 1033 (Fed. Cir. 2002); *see also Jansen v. Rexall Sundown, Inc.*, 342 F.3d 1329, 1333 (Fed. Cir. 2003); *Novartis Pharms. Corp. v. Accord Healthcare Inc.*, 387 F. Supp. 3d 429, 436 (D. Del. 2019). It sets forth the essence of the claimed invention — "treat[ment] of an angiogenic eye disorder in a patient." Ex. 1001 at Claim 1; *see also* Ex. 1001 at Abstract ("The present invention provides methods for treating angiogenic eye disorders …."); *id.* at 2:3-18 (same); *Griffin*, 285 F.3d at 1033 (construing preamble that recites a "method for diagnosing" as limiting because "[d]iagnosis is … the essence of th[e] invention; its appearance in the count gives 'life and meaning' to the manipulative steps").

³ Patent Owner reserves the right to propose additional or different constructions for claim terms in this proceeding in response to a Decision on Institution or any arguments raised by Petitioner in any future submission.

Moreover, enforcing the preamble limitation grounds the claim in its obvious utility—treating subjects suffering from angiogenic eye disorders. *See*, *e.g., Boehringer Ingelheim Vetmedica, Inc. v. Schering-Plough Corp.*, 320 F.3d 1339, 1345 (Fed. Cir. 2003) (construing the preamble as limiting because without the preamble, "the claimed method reduces to nothing more than a process … whose absence of fathomable utility" is "nothing but an academic exercise."); E.I. *Du Pont de Nemours & Co. v. Monsanto Tech. LLC*, IPR2014-00333, 2014 WL 3507803, at *4-5 (July 11, 2014) (construing the preamble as limiting because a POSA "would not understand the utility of the process" "without construing the preamble language of the claim as limiting"). Thus, the preamble is a limitation of the claim requiring that the recited dosing regimen must *treat* a patient with an angiogenic eye disorder.

The Claim 1 preamble (which recites "a patient" and "an angiogenic eye disorder") provides an antecedent basis for "the patient" who is treated and for the "angiogenic eye disorders" that are specified in dependent Claims 8-11. The method comprises "sequentially administering *to the patient*" doses of VEGF antagonist. Ex. 1001 at Claim 1 (emphasis added). This "sequentially administering" step depends upon the preamble. Without the preamble, it would be unclear *who* is receiving sequentially administered doses. Likewise, dependent Claims 8-11 rely on the preamble for their antecedent basis because they recite the particular "angiogenic eye disorder[s]" to be treated. *See id.* at Claims 8-11.

Because the preamble of Claim 1 provides an antecedent basis on which other claim limitations rely, it is a positive limitation of the claim. *See, e.g., Sanofi Mature IP v. Mylan Labs. Ltd.*, 757 F. App'x 988, 993 (Fed. Cir. 2019) (finding the preamble — "a method of increasing survival" — to be limiting because it provides an antecedent basis for which a later limitation — "a patient in need thereof" — relied); *Rapoport v. Dement*, 254 F.3d 1053, 1059 (Fed Cir. 2001); *GlaxoSmithKline LLC v. Glenmark Pharms. Inc. USA*, No. 14-877-LPS-CJB, Dkt. 133 at 14-15 (D. Del. June 3, 2016) *adopted*, 2017 WL 658468 (construing preamble — "decreasing mortality caused by congestive heart failure in a patient in need thereof" — to be limiting because term in claim body — "said patient" — "relies on and derives antecedent basis" therefrom).

Thus, the preamble of Claim 1, "A method for treating an angiogenic eye disorder in a patient" is limiting because it breathes life and meaning into the claim and provides an antecedent basis for other limitations in the body of Claim 1 and the dependent claims, thus requiring a therapeutically effective method for treating an angiogenic eye disorder.

B. The "Tertiary Dose" Must Maintain the Therapeutic Effect During Treatment

Patent Owner respectfully submits that the claim term "tertiary dose" means "dose(s) that maintain(s) a therapeutic effect throughout the course of treatment."

Under the *Phillips* standard, claim terms are afforded "their ordinary and customary meaning," which is "the meaning that the term would have to a person

of ordinary skill in the art in question at the time of the invention." *Phillips*, 415 F.3d at 1312-13. But where a term has "no previous meaning to those of ordinary skill in the prior art," one looks "[elsewhere] in the patent." *Irdeto Access, Inc. v. Echostar Satellite Corp.*, 383 F.3d 1295, 1300 (Fed. Cir. 2004); *see also MyMail, Ltd. v. Am. Online, Inc.*, 476 F.3d 1372, 1376 (Fed. Cir. 2007) (construing "coined term" that was "without a meaning apart from the patent" in view of the specification). The specification is highly relevant and often dispositive to the claim construction analysis; it is "the single best guide to the meaning of a disputed term." *Phillips*, 415 F.3d at 1315; *Vitronics Corp. v. Conceptronic, Inc.*, 90 F.3d 1576, 1582 (Fed. Cir. 1996).

The plain language of Claim 1 conveys that one or more "tertiary doses" are to be administered 12 weeks after the preceding dose. However, the term "tertiary dose" does not have a "previous meaning to those of ordinary skill in the art," (Ex. 2001 ¶ 21), "apart from the patent." *Irdeto Access, Inc.*, 383 F.3d at 1300; *MyMail, Ltd.*, 476 F.3d at 1376. Accordingly, the Board must look to the '345 Patent specification to construe the term. *Irdeto Access, Inc.*, 383 F.3d at 1300; *MyMail, Ltd.*, 476 F.3d at 1376.

The '345 Patent specification explains that, at the time of patent filing in January 2011, therapies for the treatment of angiogenic eye disorders using VEGF antagonists existed in the art. Ex. 1001 at 1:57-63. Nonetheless, the '345 Patent recognized that there remained a need for less frequent dosing regimens that could

maintain a high degree of efficacy. *Id.* at 1: 64-67. The '345 Patent successfully addressed this long-felt need:

The present inventors have *surprisingly* discovered that *beneficial therapeutic effects* can be achieved in patients suffering from angiogenic eye disorders by administering a VEGF antagonist to a patient at a frequency of once every 8 or more weeks, especially when such doses are preceded by about three doses administered to the patient at a frequency of about 2 to 4 weeks.

Id. at 2:3-18 (emphases added). Indeed, the '345 Patent discloses that a key benefit of the claimed dosing regimens is that for "most of the course of treatment (*i.e.* the *tertiary doses*)," *id.* at 2:24-31 (emphasis added), patients may be treated less frequently as compared to therapies that existed in the art. Simply put, the disclosed dosing regimens were a significant advance over existing therapies because they enabled less frequent dosing while maintaining a high degree of therapeutic efficacy. Accordingly, read in view of the specification, the term "tertiary dose" means "dose(s) that maintain(s) a therapeutic effect throughout the course of treatment."

V. THE '345 PATENT IS NOT ELIGIBLE FOR POST-GRANT REVIEW

The '345 Patent claims priority to three provisional applications filed in January of 2011 through a series of continuation applications and one continuation-

in-part application.⁴ Petitioner offers two arguments in support of its assertion that the '345 Patent is eligible for PGR. Paper 2 at 5, 23. First, Petitioner argues that dependent Claim 8, which is directed to treatment of BRVO, is supported only by a continuation-in-part ("CIP") application filed July 12, 2013. Second, Petitioner argues that the quarterly (12-week) dosing regimen recited in each of the challenged claims lacks adequate written description support. However, for the reasons discussed below, all of challenged claims are supported by the pre-AIA 2011 Provisional Applications and, therefore, the '345 Patent is not eligible for PGR.

A. The Examiner Already Determined That the '345 Patent Should Be Reviewed Under Pre-AIA Standards

During prosecution, the Examiner explicitly examined the '282 application, which issued as the '345 Patent, under pre-AIA patentability standards. On each Office Action Summary sheet, the examiner noted "No" in the AIA (FITF) Status box, and began the remarks by stating that the "present application is being examined under the pre-AIA first to invent provisions." *See* Ex. 1002 at 113-14, 143-44, 224-

⁴ Provisional Application 61/432,245 was filed on January 13, 2011 (Ex. 1045) ("the '245 Application"), provisional application 61/434,836 was filed on January 21, 2011 (Ex. 1046) ("the '836 Application"), and provisional application 61/561,957 was filed on November 21, 2011 (Ex. 1047) ("the '957 Application") (collectively, the provisional applications are the "2011 Provisional Applications").

25, 322-23. The examiner issued and analyzed rejections under "pre-AIA 35 U.S.C. 102(b)." *Id.* at 144-45, 227. The "effective filing date" for the purposes of patent prosecution and PTAB proceedings rely on the same statutory definition, thus the priority analysis is identical. *See* AIA § 3(n)(1).

The Board has previously declined to find a patent PGR-eligible where an Examiner explicitly examined the challenged patents under the pre-AIA first-toinvent provisions. *See Mylan Pharms. Inc. v. Yeda Research & Dev. Co. Ltd.*, PGR2016-00010, Paper 9 at 6-10 (Aug. 15, 2016) (finding patent was not PGR eligible based in part on "the Examiner's findings as to the effective filing date of the [challenged patent] during prosecution."); *see also Merck Sharp & Dohme Corp. v. Wyeth LLC*, PGR2017-00016 & PGR2017-00017, Paper 9 (Oct. 20, 2017) (relying on the Examiner's marking of "No" in the AIA status box in making the determination that the patent was not eligible for post-grant review). Although the Board has said that it does not treat these designations made during prosecution as entirely dispositive of the issue of PGR eligibility, it has reaffirmed the relevance of these designations in deciding eligibility. *See e.g., Commonwealth Sci. & Indus. Research v. BASF Plant Sci. GMBH*, PGR2020-0003, Paper 11 (Sept. 10, 2020).

During prosecution, Regeneron relied upon the Examiner's pre-AIA finding in formulating its arguments for patentability, and should be entitled to rely on that determination now. For this reason alone, the Board should decline to find that the '345 is eligible for post-grant review.

B. The 2011 Provisional Applications Describe Treatment of Branch Retinal Vein Occlusion (BRVO)

1. Because the 2011 Provisional Applications Support the Challenged Claims, the '345 Patent Is Not PGR Eligible

Petitioner ignores the support provided in the 2011 Provisional Applications for treatment of all angiogenic eye disorders, which a skilled artisan would have known included BRVO. It also ignores that BRVO was known to be treatable with certain VEGF antagonists and that, based on the 2011 Provisional Applications' disclosure of demonstrated efficacy in wAMD, DME, and CRVO, a skilled artisan would understand the '345 Patent inventors to be in possession of a method of treating BRVO with a VEGF antagonist fusion protein as claimed.

Instead of contending with this support and the knowledge of the skilled artisan at the time, Petitioner simply asserts that because Claim 8 specifically recites treating "BRVO," the claim was not supported until the term "BRVO" was added to the specification by continuation-in-part application No. 13/940,370 (the '370 Application), filed on July 12, 2013. Paper 2 at 18. Claim 8, however, need not rely upon the post-AIA '370 Application for support because the priority applications'

disclosure of treating angiogenic eye disorders in general and central retinal vein occlusion ("CRVO") in particular adequately supports Claim 8.⁵

Under the AIA, if an application is filed after March 16, 2013 and claims the benefit of an application filed before March 16, 2013, the application is considered to be a "transition application" for purposes of the AIA. *See* MPEP § 210; AIA § 3(n)(1). Whether the AIA applies to a transition application is determined by the claims' effective filing date, as defined by 35 U.S.C. § 100(i)(1); *Merck Sharp & Dohme*, PGR2017-00016 & PGR2017-00017, Paper 9 at 6-7. The "effective filing date" in this context is the filing date of the earliest application for which the patent or application is entitled, as to such invention, to a right of priority under section 119, 365(a), 365(b), 386(a), or 386(b) or to the benefit of an earlier filing date under section 120, 121, 365(c), or 386(c)." 35 U.S.C. § 100(i)(1)(B). The fact that descriptive matter is added in a CIP application does not presumptively determine priority for the contents of the application; rather, priority is determined on a claim-by-claim basis and depends on compliance with 35 U.S.C. § 120 and

⁵ See No. 61/432,245 (Ex. 1045) filed on January 13, 2011 and No. 61/561,957 (Ex. 1047) filed on November 21, 2011.

112(a).⁶ Because the CIP subject matter is not relied upon by any of the challenged claims for § 112 support, *see infra* pp. 19-24, all claims of the '345 Patent have an effective filing date that pre-dates the AIA. Congress did not intend such patents to be subject to PGR, and instead explicitly provided for alternative mechanisms, such as *inter partes* review, for invalidity challenges to these pre-AIA patents.

2. Treatment of Branch Retinal Vein Occlusion (BRVO) Was Described by the 2011 Provisional Applications

The stated purpose of the first provisional application, filed January 13, 2011, is to "treat any angiogenic eye disorder," explaining that an angiogenic eye disorder means "*any* disease of the eye which is caused by or associated with the growth or

a. <u>By January 2011, A POSA Would Have Understood</u> <u>BRVO to Be Within the Patent's Disclosure of</u> "Angiogenic Eye Disorders"

⁶ See MPEP § 211.05 I.B. ("claims of the continuation-in-part application that are disclosed in the manner provided by 35 U.S.C. 112(a) in the prior-filed application are entitled to the benefit of the filing date of the prior filed application"); *see also Therma-Tru Corp. v. Peachtree Doors Inc.*, 44 F.3d 988, 992 (Fed. Cir. 1995) ("A claim in a CIP application is entitled to the filing date of the parent application when the claimed invention is described in the parent specification in a manner that satisfies, inter alia, the description requirement of 35 U.S.C. § 112.") (citing *Kennecott Corp. v. Kyocera Int'l Inc.*, 835 F.2d 1419, 1421 (Fed. Cir. 1987)).

proliferation of blood vessels or by blood vessel leakage." Ex. 1045 at [0024] (emphasis added); *see also* [0031] ("The methods of the present invention are useful for treating angiogenic eye disorders in patients that have been diagnosed with or are at risk of being afflicted with an angiogenic eye disorder.").

Petitioner ignores the fact that a POSA in 2011 would have known that "angiogenic eye disorders" was a well-defined class of diseases that included branch retinal vein occlusion ("BRVO"). Ex. 2001 ¶¶ 26-28; *see also* Ex. 2003 at 2 (a 2009 listing of "angiogenic eye disorders," including "branch RVO"). Likewise, by 2011, a POSA would have known that angiogenic eye disorders may be treated by VEGF antagonists. Ex. 2001 ¶ 29; Ex. 1001 at 1:54-56 ("[I]nhibiting the angiogenic-promoting properties of VEGF appears to be an effective strategy for treating angiogenic eye disorders"). The etiology of angiogenic eye disorders and the rationale for VEGF therapy had been widely-recognized before 2011. Ex. 2001 ¶ 26-27, 35, 38; Ex. 2004 at 23.

Additionally, by the earliest 2011 priority filing date, a POSA would have known that certain VEGF antagonists had proven effective, and were even FDA-approved, for the treatment of BRVO. Ex. 2001 ¶¶ 38-43. Accordingly, when the priority applications taught that the claimed VEGF antagonists could be used to treat angiogenic eye disorders, a POSA would have understood and immediately recognized that "angiogenic eye disorders" specifically included BRVO as a known

angiogenic eye disorder that was treatable using a VEGF antagonist. Ex. 2001 ¶¶ 44-46.

Petitioner contends that the skilled artisan "would not understand that successful treatment of one vascular disease ... means another (*e.g.*, BRVO) is *necessarily* treated." Paper 2 at 21 (emphasis added). This argument ignores the priority applications' teaching that the claimed method could be used for *any* angiogenic eye disorder. Ex. 1001 at 1:54-56. It also ignores the fact that BRVO had already been shown to be successfully treated by anti-VEGF agents before 2011. Ex. 2001 ¶¶ 39-43.

Based on the known etiology of BRVO by 2011 and the specific demonstration in the art that anti-VEGF agents had successfully treated BRVO, a POSA would have understood that BRVO was "an angiogenic eye disorder" "which is caused by or associated with the growth or proliferation of blood vessels or by blood vessel leakage," and understood the 2011 Provisional Applications to disclose methods of treating BRVO.

b. <u>The 2011 Provisional Applications' Disclosure of CRVO</u> <u>Treatment Would Have Provided a POSA With</u> <u>Confirmation That BRVO Was Among the Angiogenic</u> <u>Eye Disorders That Could Be Treated By the Claimed</u> <u>Dosing Regimens</u>

Not only would a POSA have understood from the general disclosure that BRVO is an "angiogenic eye disorder" that could be treated with the claimed antiVEGF fusion proteins, but the '245 Application specifically describes treatment of CRVO as one type of angiogenic eye disorder and thus confirms that the disclosed treatment methods include treatment of BRVO. Ex. 1045 at [0024] ("Non-limiting examples of angiogenic eye disorders that are treatable using the method of the present invention include ... central retinal vein occlusion"). CRVO, like BRVO, is a type of retinal vein occlusion. Retinal vein occlusions result from the development of thrombus in the retinal vein resulting in reduced blood flow and exhibit other similar pathologies, including VEGF up-regulation. Ex. 2001 ¶ 37. The principle distinction between the two diseases is the locus of the occlusion in the retinal veins, but in both cases the occlusion occurs from VEGF up-regulation and is treatable with an anti-VEGF agent. Ex. 2001 ¶ 37. By 2011, a POSA would have recognized a disclosure of treatment of CRVO with an anti-VEGF agent as an indicator for successful treatment of BRVO with the same type of therapy.

Petitioner argues that a POSA would not have recognized that a CRVO treatment would be used for BRVO because the diseases have anatomic differences, affect different patient populations and, historically, had different standards of care. Paper 2 at 22; Ex. 1003 ¶¶ 126-127. None of those differences were relevant by 2011 because, by that time, the skilled artisan understood that the two diseases shared a common etiology rooted in VEGF upregulation and, further, that both could be treated successfully with anti-VEGF therapies. Indeed, by 2011, clinical trials

showed successful treatment of CRVO and BRVO by anti-VEGF antagonists ranibizumab (Lucentis) and bevacizumab (Avastin). Ex. 2001 ¶ ¶ 39-42.⁷ In fact, clinicians had successfully treated patients with both types of retinal vein occlusions with off-label Avastin since approximately 2006, and by June 2010, Lucentis had received FDA approval for treatment of both indications. Ex. 2001 ¶¶ 43, 48; Ex. 2005. Further, the retina community frequently described these developments in the treatment of BRVO and CRVO in tandem. Ex. 2001 ¶ 51; Ex. 2006 at 2 (discussing Lucentis phase III BRVO and CRVO trials together).

⁷ As with all other evidentiary questions at the institution stage, the burden is on Petitioner to show that it is more likely than not that at least one challenged claim is unpatentable. *See* 35 U.S.C. § 324(a); *see also Hulu, LLC v. Sound View Innovations*, *LLC*, IPR2018-01039, Paper 29, at 16-20 (Dec. 20, 2019). Consistent with this statutory framework and Petitioner's ultimate burden of proof, the Board should consider all evidence and apply no evidentiary presumption for testimonial evidence favoring Petitioner. *See also* 85 Fed. Reg. 79120 (Dec. 9, 2020) (consistent with this statutory framework, USPTO has revised its rules to ensure any testimonial evidence submitted with a POPR will be taken into account in the totality of the evidence).

In addition to the success of other anti-VEGF agents, Regeneron's own anti-VEGF therapy had demonstrated efficacy in treating CRVO and was disclosed in the '957 Application, filed on November 21, 2011. The '957 Application disclosed the 24 and 52 week results of Regeneron's Phase III trial in CRVO, which demonstrated statistically significant improvements in visual acuity as compared to sham control. Ex. 2001 ¶ 55; Ex. 1047 at [0064]-[0066].

Accordingly, the knowledge of the skilled artisan combined with the disclosures in the '345 Patent's earliest provisional applications adequately support Claim 8.

C. The Claimed Twelve-Week Dosing Regimen Is Fully Supported by the 2011 Provisional Applications

For the reasons discussed below, *infra* Section VII.C, the 12-week dosing regimen recited in the '345 Patent claims is supported by the 2011 Provisional Applications. Because both written description challenges fail to establish PGR eligibility, the Petition should be denied.

VI. THE BOARD SHOULD DENY INSTITUTION UNDER 35 U.S.C. § 325(d)

The Board should exercise its discretion and deny institution under 35 U.S.C. § 325(d) because Petitioner relies on the same art and arguments that were considered by the Examiner during prosecution of the '345 Patent and fails to show that, in considering that art, the Examiner made any error material to the patentability of the challenged claims.

The Board applies a two-part framework to analyze discretionary denial under § 325(d): "(1) whether the same or substantially the same art previously was presented to the Office or whether the same or substantially the same arguments previously were presented to the Office; and (2) if either condition of [the] first part of the framework is satisfied, whether the petitioner has demonstrated that the Office erred in a manner material to the patentability of challenged claims." *Advanced Bionics, LLC v. MED-EL Elektromedizinische Geräte GmbH*, IPR2019-01469, 2020 WL 740292, at *3-4 (Feb. 13, 2020) (precedential), citing *Becton, Dickinson & Co. v. B. Braun Melsungen AG*, IPR2017-01586, Paper 8 (Dec. 15, 2017) (precedential as to § III.C.5, first paragraph).

Petitioner misstates this standard and incorrectly suggests that the art and arguments must have been raised *in the same way* as it was during prosecution for \$325(d) to apply. Paper 2 at 71. That is not correct. Rather, the Board has held that \$325(d) discretionary denial is appropriate where the same or substantially the same art or argument was previously presented to the Office, regardless of how it was applied or considered by the Examiner. *Advanced Bionics*, 2020 WL 740292, at *3. If the art or argument were previously presented, the Board moves on to determine if Petitioner has shown that the Office materially erred. *Id*.

A. The Examiner Considered the Same or Substantially the Same Art and Arguments During Prosecution (*Becton, Dickinson* factors (a), (b), and (d)⁸)

Petitioner's Grounds rely upon the same art and arguments that were presented to, and considered by, the Examiner during prosecution of the '345 Patent, thus satisfying step one of the *Advanced Bionics* framework. In Ground 1, Petitioner argues that the '345 Patent claims are anticipated by Shams (Ex. 1004); in Ground 2, Petitioner argues that the '345 Patent claims are rendered obvious by the 2009 Press Release (Ex. 1005) in view of Shams; and in Ground 3, Petitioner argues that the '345 Patent claims lack adequate written description.

1. Shams (Ex. 1004)

Shams WO 2006/047325 (Ex. 1004) is an abandoned Genentech, Inc. ("Genentech") patent application, which published on May 4, 2006 from PCT/US2005/038006. Regeneron presented Shams to the Office in an Information

⁸ Becton, Dickinson factors: (a) similarities and material differences between the asserted art and prior art involved during examination; (b) cumulative nature of the asserted art and prior art evaluated during examination; and (d) extent of overlap between arguments made during examination and the manner in which petitioner relies on the prior art. *See Becton, Dickinson*, IPR2017-01586, Paper 8 at 17–18 (precedential as to § III.C.5, first paragraph).

Disclosure Statement ("IDS") that was considered by the Examiner during prosecution of the '345 Patent. Ex. 1002 at 225, 239. Shams is cited on the face of the '345 Patent. Ex. 1001 at 1. Petitioner admits that Shams was submitted in an IDS and marked "considered" by the Examiner during prosecution.⁹ Paper 2 at 71.

Citing a single pre-*Advanced Bionics* decision, Petitioner argues that "[t]he Board has consistently declined exercising its discretion under Section 325(d) when the only fact a Patent Owner can point to is that a reference was disclosed to the Examiner during the prosecution." Paper 2 at 71. However, the Board has expressly rejected this argument, stating that *Advanced Bionics* provides that "previously presented art includes art made of record ... such as on an [IDS]." *See, e.g., Biocon Pharma Ltd. v. Novartis Pharms. Corp.*, IPR2020-01263, 2021 WL 608300, at *4 (Feb. 16, 2021) (citing *Advanced Bionics*, 2020 WL 740292, at *3); *see also Philip Morris Prods., S.A. v. Rai Strategic Holdings, Inc.*, IPR2020-00921, 2020 WL 6750120, at *5 (Nov. 16, 2020) ("the art presented in the Petition is the same as the art previously presented to the Office during examination because all of Petitioner's

⁹ Ex. 1002 at 225 ("The information disclosure statement (IDS) submitted on 19 June 2019 ... [has] been considered by the Examiner."); *id.* at 239 ("ALL REFERENCES CONSIDERED EXCEPT WHERE LINED THROUGH") (Shams not lined through).

references were cited in an IDS and are listed as cited art on the front face of the '268 Patent."); *Mylan Pharms. Inc. v. Merck Sharp & Dohme Corp.*, IPR2020-00040, 2020 WL 2478503, at *6 (May 12, 2020) ("Petitioner's emphasis on the absence of any prior art rejection as if dispositive on the 325(d) inquiry is, thus, misplaced; the first part of the § 325(d) framework may be met when relied-upon art is presented in an IDS but never discussed or cited in a rejection by the Examiner..."). Thus, Shams was previously presented to and considered by the Office.

2. 2009 Press Release (Ex. 1005)

The 2009 Press Release is entitled, "Enrollment Completed in Regeneron and Bayer HealthCare Phase 3 Studies of VEGF Trap-Eye in Neovascular Age-Related Macular Degeneration (Wet AMD)," and labeled with a date of September 14, 2009. The 2009 Press Release appears on the face of the '345 Patent. Ex. 1001 at 8. Petitioner admits that the 2009 Press Release was the basis for a rejection under 35 U.S.C. § 102 during the prosecution of the '345 Patent, which was overcome by Regeneron. Paper 2 at 2, 75.

Since *Advanced Bionics*, the Board has consistently found that a reference was previously provided to the Office when it is part of the basis of a rejection during prosecution. *E.g., Balt USA, LLC v. Microvention, Inc.*, IPR2020-01259, 2021 WL 219251, at *8 (Jan. 21, 2021); *Gofire, Inc. v. Canopy Growth Corp.*, IPR2020-00044, 2020 WL 5991725, at *4 (Oct. 9, 2020); *Flex Logix Techs., Inc. v. Konda*, IPR202000262, 2020 WL 4462127, at *4 (Aug. 3, 2020); *GlaxoSmithKline Consumer Healthcare Holdings (US) LLC v. Cipla Ltd.*, IPR2020-00371, 2020 WL 4390665, at *6 (July 31, 2020); *Samsung Elec. Co., Ltd. v. Neodron Ltd.*, IPR2020-00334, 2020 WL 3892132, at *5 (Jul. 10, 2020); *Boragen, Inc. v. Syngenta Participations AG*, IPR2020-00124, 2020 WL 2206972, at *6 (May 5, 2020). Thus, the 2009 Press Release was previously presented to and considered by the Office.

3. Written Description

Petitioner asserts that the Board should decline to exercise its discretion under § 325(d) with respect to Ground 3 (written description) because "no 12-week dosing regimen rejections or similar rejections were raised during prosecution." Paper 2 at 76. Following *Advanced Bionics* and its progeny, the Board is entitled to find that the same or substantially the same arguments or issues were presented to the Examiner during prosecution even in the absence of an express rejection. *See Universal Imaging Indus., LLC v. Lexmark Int'l Inc.*, IPR2019-01387, 2020 WL 2201770, at *3-4 (May 4, 2020).¹⁰ Moreover, the prosecution history reveals that the Examiner reviewed the specification of the '345 Patent and considered written description support in the instant application and related applications in the priority

¹⁰ The Board has not addressed whether, post-Advanced Bionics, an Examiner is presumed to have considered the adequacy of written description support for claims during prosecution. However, such a presumption is consistent with the burdenshifting framework of Advanced Bionics. As Advanced Bionics explains, "[a]t bottom, this framework reflects a commitment to defer to previous Office evaluations of the evidence of record unless material error is shown." 2020 WL 740292, at *3. While the Board in Hybridgenics v. Forma Therapeutics found that the absence of a written description rejection did not establish that an Examiner had considered written description arguments, that decision is inapplicable here. See PGR2018-00098, Paper 10 at 20-21 (Mar. 20, 2019). Not only is Hybrigenics a nonprecedential pre-Advanced Bionics decision, but the concern that animated the Board's decision in *Hybrigenics* — that "[t]o find otherwise would potentially suggest that we should apply our discretion under 325(d) to deny review in every post-grant review where written description is challenged..." (Id. at 20) — ignores the fact that Advanced Bionics creates a rebuttable presumption that a Petitioner can overcome by showing material error.

chain. In light of Patent Owner's evidence that the Examiner considered the specification, Petitioner should be required to show error material to patentability. Petitioner fails to even so allege.

Petitioner relies only on the absence of a written description rejection as evidence that the Examiner did not consider §112. However, consistent with Advanced Bionics, the Examiner should be presumed to have reviewed and understood the '345 Patent's disclosures relating to the claimed 12-week dosing regimen. The Board has noted "filt is reasonable to assume that the Examiner read the Specification and understood these statements as part of examining" the '345 Patent. Universal Imaging Indus., LLC v. Lexmark Int'l, Inc., IPR2019-01387, 2020 WL 959375, at *6 (Feb. 27, 2020). Likewise, there is a presumption that an adequate written description of the claimed invention is present in the specification as filed. MPEP § 2163. By analogy, in the context of *inter partes* review, the Board has held that an Examiner is presumed to be aware of the substantive disclosure of the material incorporated by reference into and which effectively becomes part of the specification. Free Stream Media Corp. v. Alphonso Inc., IPR2019-00762, 2019 WL 4200650, at *5 (Sept. 4, 2019) (citing Telemac Cellular Corp. v. Topp Telecom, Inc., 247 F.3d 1316, 1329 (Fed. Cir. 2001); see Monolithic Power Sys., Inc. v. Volterra Semiconductor LLC, IPR2020-01348, 2021 WL 838428, at *5 (Mar. 4, 2021).

Furthermore, the prosecution history of the '345 Patent shows that the Examiner did consider the adequacy of the specification's disclosure. In particular, Patent Owner submitted the now-issued '345 Patent claims by way of a preliminary amendment, adding new claims that included the 12-week dosing regimen. Ex. 1002 at 386. In the arguments/remarks presented to the Examiner, Patent Owner asserted that support for the new 12-week dosing claims could be found in the originallypending claims and in the specification. Ex. 1002 at 389. Pursuant to \$1.121(f), no amendment may introduce new matter into the disclosure of an application. 37 CFR §1.121(f); see Kolmes v. World Fibers Corp., 107 F.3d 1534, 1538-39 (Fed. Cir. 1997). Additionally, an examiner is obligated to review the specification and determine whether the invention as claimed complies with all statutory requirements, including §112. MPEP § 2103 ("Examiners will review the complete specification."). In a Non-Final Rejection dated April 3, 2019, the Examiner confirmed that the preliminary amendment had been entered in full, *i.e.*, the Examiner found adequate support in the specification for the newly-filed claims that included the 12-week dosing regimen. Ex. 1002 at 323. Thus, the issue of the §112 sufficiency of the 12-week dosing claims was previously considered by the Examiner, as reflected by the Examiner's entry of the preliminary amendment.

In addition, the prosecution history of related family members of the '345 Patent confirms that the Examiner reviewed and considered the disclosure of the ³⁴⁵ specification. In several applications leading to the ³⁴⁵ Patent, including the '282 Application, after entering the preliminary amendment, the Examiner objected to the specification because it did not include "[a]n updated status of the parent nonprovisional application" as "the first sentence." Ex. 1002 at 323. Regeneron amended the specification to address the Examiner's objection. Ex. 1002 at 278. In light of the amendment, the Examiner withdrew his objection. Ex. 1002 at 225. In several other applications in the priority chain, the Examiner made informality-based objections to the specification's disclosure that Regeneron similarly overcame. E.g., Ex. 2007 at 151 (objecting to specification because "[a]n updated status of the parent nonprovisional application should be included in the first sentence"); Ex. 2008at 104 (same). And, in S.N. 13/940,370, another application in the priority chain that issued as U.S. Patent No. 9,254,338, the Examiner rejected pending claims for lack of written description. Ex. 2009 at 262-264. Accordingly, the prosecution history reflects that the Examiner considered the specifications as well as potential § 112 issues in reviewing Regeneron's dosing regimen applications.

B. Petitioner Fails to Show That the Examiner Erred in a Manner Material to Patentability (*Becton, Dickinson* factors (c), (e), and (f)¹¹)

As step one has been satisfied, Petitioner must show that the Office erred in a manner material to the patentability of the challenged claims. "An example of a material error may include misapprehending or overlooking specific teachings of the relevant prior art where those teachings impact patentability of the challenged claims." *Advanced Bionics*, 2020 WL 740292, at *4 n.9. "If reasonable minds can disagree regarding the purported treatment of the art or arguments, it cannot be said that the Office erred in a manner material to patentability." *Id.* at *4.

1. Petitioner Fails to Show That the Examiner Erred in a Manner Material to Patentability in His Analysis of Shams (Ground 1)

Petitioner does not identify any "material error" that the Examiner committed *in this case*. Petitioner cannot demonstrate material error simply because Shams

¹¹ Becton, Dickinson factors: (c) the extent to which the asserted art was evaluated during examination, including whether the prior art was the basis for rejection; (e) whether Petitioner has pointed out sufficiently how the Examiner erred in its evaluation of the asserted prior art; and (f) the extent to which additional evidence and facts presented in the Petition warrant reconsideration of the prior art or arguments. See Becton, Dickinson, IPR2017-01586, Paper 8 at 17-18.

was not substantively discussed in the prosecution record. *Universal Imaging*, 2020 WL 959375, at *5 (finding that the absence of a rejection based upon the petitioned grounds "is not the end of [the Board's] analysis" on material error). Instead, the Board considers the Examiner's familiarity with the substance of the petitioned reference. *Id.* As shown above, the prosecution record indicates that the Examiner was familiar with Shams. *Supra* Section VI.A.1; *see Husky Injection Molding Sys., Ltd. v. Plastipak Packaging, Inc.,* IPR2020-00438, 2020 WL 4353621, at *7 (July 29, 2020) (in determining petitioner did not meet its burden under step two, finding statement that "all references considered except where lined through" in prosecution record indicated that the Examiner substantively considered asserted reference).

Petitioner points to the prosecution of an entirely unrelated patent application, involving a different therapeutic agent, specification and claims, where a different examiner applied Shams' alleged disclosure of a 12-week dosing regimen to reject the claims as-filed.¹² Paper 2 at 73-74. But, the Petition is silent as to what error the Examiner purportedly committed during the prosecution of the '345 Patent. At best, this extrinsic evidence indicates that "reasonable minds disagree[d] regarding the

¹² Notably, Novartis overcame the rejection based on the European equivalent of Shams, and a patent issued with claims reciting the 12-week dosing element. *E.g.*, U.S. Patent No. 10,035,850 at 17:43-18:43.

purported treatment of the art or arguments," which is insufficient to show "that the Office erred in a manner material to patentability." *Advanced Bionics*, 2020 WL 740292, at *3;¹³ see also Regeneron Pharms., Inc. v. Kymab Ltd., IPR2020-00389, 2020 WL 2738613, at *7 (May 26, 2020) (petitioner only offered "a different interpretation" of prior art, which is not material error). The mere fact that one examiner applied Shams in a rejection, while another marked it as cited and considered, is not enough for Petitioner to meet its burden to show that the Examiner committed material error. *Sony Interactive Entm't LLC v. Terminal Realty, Inc.*, IPR2020-00711, 2020 WL 6065188, at *5 (Oct. 13, 2020) (finding that Petitioner's argument that the asserted references were not evaluated by the examiner failed to

¹³ Petitioner's reliance on *Advanced Bionics* is inapposite. Paper 2 at 72-73. There, the claims at issue were rejected in view of the petitioned reference during prosecution. 2020 WL 740292 at *8. In dicta, the Board postulated that a petitioner may be able to show error if the record is silent or not well developed with respect to a reference. *Id.* at *4. But a silent record is not dispositive; Petitioner is still required to identify Examiner error. *Id.* at *3 ("At bottom, this framework reflects a commitment to defer to previous Office evaluations of the evidence of record *unless material error is shown*.") (emphasis added). Likewise, a disagreement with the Examiner is not material error. *Id.*

sufficiently identify Examiner error). In *Sony Interactive*, the Board noted that "Sony [Petitioner] was provided the opportunity to provide explanation [of material error], but Sony was silent in this regard.... Accordingly, *Becton, Dickinson* Factor (e) favors exercising our discretion to deny institution." *Id.* Similarly, here Petitioner fails to identify any material error that the Examiner purportedly committed in his consideration of Shams.

2. Petitioner Does Not Argue That the Examiner Erred in a Manner Material to Patentability as to the 2009 Press Release or Written Description (Grounds 2 and 3)

Petitioner does not allege that the Examiner committed any error during prosecution with respect to its analysis of the 2009 Press Release or the written description requirement. In fact, Petitioner is entirely silent in that regard.

The Board has found a petitioner's failure to allege material error to be a sufficient basis to determine that petitioner did not carry its burden to meet step two. *E.g., Balt,* 2021 WL 219251, at *10 (Petitioner's own independent analysis of prior art without reference to or discussion of Examiner's analysis is insufficient to show material error); *NXP USA, Inc. v. Impinj, Inc.*, IPR2020-00519, 2020 WL 4805424, at *5 (Aug. 17, 2020) (no discussion of material error); *GlaxoSmithKline Consumer Healthcare Holdings (US) LLC v. Cipla Ltd.*, IPR2020-00369, 2020 WL 4390663, at *5 (July 31, 2020) (step two not met when, in part, petitioner did not "explicitly allege error in the Examiner's previous consideration of the prior art or arguments").

Petitioner asserts only that the combination of the 2009 Press Release in view of Shams was not considered during prosecution, and that the 2009 Press Release was not considered for a portion of its disclosure (8-week dosing) or as a §103 reference. Paper 2 at 75. Petitioner is splitting hairs. Petitioner does not dispute that the substance of the 2009 Press Release was expressly considered by the Examiner as the basis for a §102(b) rejection. Ex. 1002 at 227-228. Moreover, the 2009 Press Release is two pages long. Ex. 1005 at 1-2. The disclosure of the 8week dosing regimen appears in the same paragraph as the PRN regimen capped at 12 weeks. Ex. 1002 at 1. In light of this, it is not credible for Petitioner to allege that the Examiner was not aware of or did not consider the 2009 Press Release, including its disclosure of 8-week dosing. Paper 2 at 75.

Additionally, Petitioner does not substantively address the evidence of unexpected results presented during the prosecution of the '345 Patent. In response to a double patenting rejection, Regeneron argued that even if the claimed invention were *prima facie* obvious, that finding would be overcome by the fact that the

claimed invention exhibits unexpected results.¹⁴ Ex. 1002 at 284-286. The Examiner withdrew his rejection, in pertinent part because of Regeneron's "persuasive arguments as they pertain to the rejection...." Ex. 1002 at 225-226. Noticeably absent from the Petition is any argument that the Examiner committed material error when he found this evidence persuasive. The Board has found that petitioner did not meet its burden under step two when it failed to show that the examiner's reliance on the unexpected results was material error. *Biocon Pharma*, 2021 WL 608300, at *6-7; *see also Apotex Inc. v. Celgene Corp.*, IPR2018-00685, 2020 WL 2095846, at *4-5 (Apr. 30. 2020) (step two not met when petitioner did not advance "any argument or evidence that the Examiner erred in evaluating or

¹⁴ During prosecution, Regeneron argued that monthly administrations of ranibizumab were "(1) expensive; (2) painful to the patient; (3) inconvenient for the patient as well as the patient's family; (4) psychologically and physically traumatic to the patient; and (5) subjects the patient to potential adverse effects such as infection with each treatment visit," and thus there was a need in the art for alternative treatment protocols. Ex. 1002 at 283-284. Regeneron explained, however, that the claimed regimen was not *prima facie* obvious, and in fact, a 2012 paper by Heier et al demonstrated that dosing less frequently than every month was surprisingly noninferior to monthly dosing. *See infra* Sections VII.B.1, VII.C.3.

balancing the evidence of unexpected results."). Here, because Petitioner utterly fails to address the Examiner's consideration of the unexpected results that eventually led to the issuance of the '345 Patent, Petitioner has failed to meet its burden under *Advanced Bionics* step two.

Because the same or substantially the same art and arguments were previously presented to the Office and were considered by the Examiner, and Petitioner has failed to show that the Office materially erred in its consideration of that art or argument, the Board should exercise its discretion and deny institution under §325(d).

VII. THE BOARD SHOULD DENY INSTITUTION BECAUSE PETITIONER FAILS TO MAKE ITS THRESHOLD SHOWING THAT AT LEAST ONE CHALLENGED CLAIM IS UNPATENTABLE

In a post-grant review proceeding, the Petitioner must "demonstrate that it is more likely than not that at least 1 of the claims challenged in the petition is unpatentable." 35 U.S.C. § 324(a). And the Petition must "identif[y], in writing and with particularity, each claim challenged, the grounds on which the challenge to each claim is based, and the evidence that supports the grounds for the challenge to each claim." *Id.*, § 322(a)(3). Where Petitioner fails to meet its threshold burden, the Board "may not authorize a post-grant review to be instituted." 35 U.S.C. § 324(a). Indeed, the Board denies institution where a petitioner has failed to demonstrate that at least one challenged claim is unpatentable. *See, e.g., One World Techs., Inc. v.*

Chervon (HK) Ltd., PGR2020-00059, 2020 WL 7222691 (Dec. 7, 2020); *Eton Pharms., Inc. v. Exela Pharma Scis., LLC*, PGR2020-00064, Paper 12 (Nov. 18, 2020); *Align Tech., Inc. v. 3Shape A/S*, PGR2018-00103, 2019 WL 2112182 (May 13, 2019). For the reasons discussed below, Petitioner has failed to "demonstrate that it is more likely than not that at least 1 of the" '345 Patent claims is unpatentable for Grounds 1, 2, and 3, and thus, denial of the petition is warranted. 35 U.S.C. § 324(a).

A. Ground 1: Petitioner Fails to Demonstrate That It Is More Likely Than Not That At Least One of the Challenged Claims Is Anticipated

Petitioner fails to show that any challenged claim is more likely than not unpatentable for anticipation based on Shams (Ground 1).

Shams (Ex. 1004) is a Genentech patent application that published on May 4, 2006 and was abandoned. The Shams specification discloses a single prophetic example, Example 1, that corresponds to a study that Genentech conducted with its VEGF antibody fragment, Lucentis[®] (ranibizumab), called PIER. Ex. 2002 ¶ 29-30. The PIER trial began in August 2004 and was completed in March 2007, nearly a year after Shams was published. Ex. 2010. The purpose of the study was to "evaluate the efficacy and safety of ranibizumab administered monthly for three months and then quarterly" in subjects with AMD. Ex. 1026 at 1. As shown in Figure 2 of Shams, subjects would receive three monthly doses of ranibizumab

followed by doses every 3 months for a period of 24 months. Ex. 1004 at 32:8-13. Shams does not include any data, nor does it report any results.

1. Petitioner Fails to Demonstrate That Shams' Treatment Schema Discloses the Recited Fusion Protein

Claim 1 requires using a VEGF antagonist that is a "receptor-based chimeric molecule comprising an immunoglobin-like (Ig) domain 2 of a first VEGF receptor which is Flt1 and Ig domain 3 of a second VEGF receptor which is Flk1, and a multimerizing component." Ex. 1001 at 21:65-22:54. The Petition asserts that Shams discloses this limitation, but provides no support whatsoever for this proposition. Paper 2 at 31. It relies on the "treatment schema" shown in Figure 2, which Petitioner repeatedly depicts in the Petition. Paper 2 at 26-28, 31, 33, 38. The treatment schema, however, only shows a prospective method of treatment using ranibizumab, an antibody fragment, not a VEGF antagonist fusion protein as required by the claims of the '345 Patent.¹⁵ That method of treatment does not anticipate the '345 Patent claims.

The Petition argues that Shams teaches that the treatment schema of Figure 2 could be used with any VEGF antagonist "includ[ing] Regeneron's fusion protein."

¹⁵ Even for ranibizumab, Shams merely discloses its plan to assess the efficacy and safety of administering ranibizumab according to the Figure 2 treatment schema, but says nothing about the treatment schema actually being effective.

Paper 2 at 31-32. But nothing in the "treatment schema" of Figure 2, on which Petitioner relies, states that one should use that schema with "VEFG-Trap (Regeneron)." The only compound Genentech identifies for evaluation in Figure 2 is its own ranibizumab. Ex. 1004 at 31-34, Fig. 2. And even for ranibizumab, Shams merely discloses a prophetic plan to assess ranibizumab using the Figure 2 treatment schema.¹⁶ Figure 2 says nothing about treating angiogenic eye disorders with a fusion protein, let alone the specific type of "receptor-based chimeric molecule" required by the '345 Patent, using the Figure 2 treatment schema. To the contrary, as discussed *infra*, Shams expressly discloses that anti-VEGF antibodies are preferred, and that ranibizumab in particular, an antibody fragment, is the most preferred VEGF antagonist disclosed by Shams. Ex. 1004 at 7:6, 13-14. The VEGF antagonist fusion protein molecules required by the claims of the '345 Patent are not antibody fragments. *See infra*. Thus Shams not only fails to disclose the claimed dosing method using a VEGF antagonist fusion protein as required by

¹⁶ Petitioner disingenuously suggests that Groups 1, 2, and 3 depicted in Shams' Figure 2 treatment schema "received" a 0.3 mg dose 0.5 mg, or sham injection. Paper 2 at 26. However, Shams does not report any actual administration and its sole example, Example 1, is a prophetic example drafted in present tense. *See* Ex. 1004 at 31:1-19.

the claims, but it teaches that such molecules are the least preferred to use in any treatment method.

For this reason alone, Petitioner fails to carry its burden to show that Shams anticipates.

2. Petitioner Fails to Demonstrate That Shams' Reference to "VEGF-Trap (Regeneron)" Discloses the Recited Fusion Protein

Petitioner also fails to show that Shams expressly or inherently discloses the VEGF antagonist fusion protein recited in Claims 1-11 of the '345 Patent. Petitioner relies on Shams' reference to a "VEGF-Trap (Regeneron)" as allegedly disclosing use of a "receptor-based chimeric molecule" meeting the specific molecular requirements of the '345 Patent claims. The Petition fails to satisfy its burden of proving any aspect of this empty assertion. There is no evidence that "VEGF-Trap (Regeneron)" refers to any particular molecule, nor is there evidence that the term refers to a category of molecules that necessarily satisfies the requirements of the '345 Patent.

Petitioner does not even attempt to meet its burden of explaining what the term "VEGF-Trap (Regeneron)" denoted to a skilled artisan at the time of filing. Nothing in Shams discloses the amino acid sequence or component parts of "VEGF-Trap (Regeneron)," nor does it identify any references that provide this information.

Instead, the Petition asserts that the term "VEGF-Trap (Regeneron)" discloses a genus of compounds that "includes" compounds that satisfy the specific VEGF antagonist fusion protein limitations of the '345 Patent claims. Paper 2 at 31 ("One of skill in the art in 2006 would understand that 'VEGF Trap (Regeneron)' includes Regeneron's fusion protein, which, in 2006, included 'a receptor-based chimeric molecule comprising an immunoglobin-like (Ig) domain 2 of a first VEGF receptor which is Flt1 and Ig domain 3 of a second VEGF receptor which is Flk1, and a multimerizing component."") (citing Ex. 1003 ¶89). Notably, paragraph 89 of Dr. Wu's declaration (Petitioner's cited support) relies on the '345 patent itself (Ex. 1001), and not on Shams to purportedly evidence this limitation. Thus, Petitioner fails to show that a POSA would have understood Shams' reference to "VEGF Trap (Regeneron)" to be the recited fusion proteins of Claim 1. Shams therefore fails to identify by name or otherwise a single molecule satisfying the requirement of the '345 Patent claims and, at best, discloses a vast sea of molecules that *could* include chimeric molecules comprising the specific domains of a VEGF receptor specified by the '345 Patent claims, but could also include molecules that do not.

Indeed, as of January 2011, a POSA would have known that there were numerous Regeneron VEGF-Trap molecules, including many that do not satisfy the requirements of the '345 Patent claims. By the early 2000s, Regeneron had developed, tested and published on a variety of engineered VEGF fusion proteins that it called "VEGF Trap" molecules. For instance, a 2002 PNAS article published by Holash *et al.*¹⁷ describes a number of different Regeneron's VEGF-Traps, many of which fall outside of the scope of the '345 Patent claims. Ex. 2011 at 1.¹⁸ Likewise, by 2006, a Regeneron published patent application to Daly *et al.*, PCT/US2004/021059, titled "VEGF Traps and Therapeutic Uses Thereof," discloses multimeric VEGF-binding proteins comprising two or more fusion polypeptides (also called VEGF 'trap' molecules), which include molecules that fail

¹⁸ Holash discloses that VEGF Trap_{parental} was created by fusing the first three Ig domains of Flt1 (VEGFR1) to the Fc region; VEGF-Trap $_{\Delta B1}$ was created by removing a highly basic 10-aa stretch from the third Ig domain of the parental VEGF-Trap; and VEGF-Trap $_{\Delta B2}$ was created by removing the entire first Ig domain from VEGF-Trap $_{\Delta B1}$. *Id.* None of these disclosed VEGF-Traps, which contain only domains of VEGF receptor 1 and no domains from VEGF receptor 2, satisfies the fusion protein limitation of the '345 Patent claims.

¹⁷ Petitioner acknowledges that Holash 2002 was in the prior art since it relies on this publication as a ground for challenge in its concurrently filed IPR challenging Regeneron Patent No. 10,464,992 in IPR2021-00402, Petition at 4.

to satisfy the compound requirements of the '345 Patent claims.¹⁹ Thus, a POSA would not have understood Shams' disclosure of "VEGF-Trap (Regeneron)" — a genus of fusion proteins — to necessarily satisfy the recited fusion protein limitation of '345 Patent, Claim 1.

As the Federal Circuit has explained, a prior art reference may anticipate without disclosing a feature of the claimed invention only "if that missing characteristic is necessarily present, or inherent, in the single anticipating reference." *Schering Corp. v. Geneva Pharms.*, 339 F.3d 1373, 1377 (Fed. Cir. 2003). "Inherency, however, may not be established by probabilities or possibilities. The mere fact that a certain thing may result from a given set of circumstances is not sufficient." *In re Robertson*, 169 F.3d 743, 745 (Fed. Cir. 1999).

The mere possibility that "VEGF-Trap (Regeneron)" could comprise a chimeric fusion protein molecule meeting the limitation of Claim 1 is insufficient to demonstrate inherency for anticipation. *See Amgen, Inc. v. Alexion Pharms., Inc.,* IPR2019-00739, Paper 15, at 24-25 (Aug. 30, 2019) (rejecting inherent anticipation where "eculizumab" referred to at least two different proteins in the prior art,

¹⁹ Daly discloses that VEGF-traps can include receptor components from VEGFR3 (Flt-4), which fall outside the requirements of the '345 Patent claims. Ex. 2012 at [002].

including the unclaimed "Thomas IgG4 isotype eculizumab"); *see also Endo Pharms. Sols., Inc. v. Custopharm Inc.*, 894 F.3d 1374, 1383 (Fed. Cir. 2018) (finding incomplete prior art disclosure of a composition insufficient to inherently disclose the claimed composition).

Shams' recitation of a generic "VEGF-Trap (Regeneron)" does not expressly or inherently disclose a method of treating an angiogenic eye disorder using the recited VEGF antagonist fusion protein of the challenged claims. Disclosure of a method of using a genus of compounds does not anticipate a method of using one compound from that genus. *See Impax Lab'ys, Inc. v. Aventis Pharms., Inc.,* 468 F.3d 1366, 1383 (Fed. Cir. 2006). Thus, Petitioner has failed to show that Shams anticipates the challenged claims.

3. Shams Does Not Disclose the '345 Patented Invention As Arranged in the Claims

To anticipate, a reference "must not only disclose all elements of the claims within the four corners of the document, but must also disclose those elements arranged as in the claim." *Net MoneyIN, Inc. v. VeriSign, Inc.*, 545 F.3d 1359, 1369 (Fed. Cir. 2008) (internal quotations omitted). Petitioner fails to show that the disclosures in Shams upon which it relies are arranged as in the challenged claims of the '345 Patent.

Petitioner picks and chooses from different portions of the Shams' specification and different embodiments without identifying any language tying them together into a single, coherent disclosure of an anticipating method. For the claimed dosing regimen — "a single initial dose" followed by "secondary dose[s] … administered 4 weeks after the immediately preceding dose" followed by "tertiary dose[s] … administered 12 weeks after the immediately preceding dose," (Ex. 1001 at Claim 1) — Petitioner relies on Figure 2 of Shams, which illustrates the prophetic dosing regimen of Example 1. *See* Paper 2 at 30-38.

But, as explained above, because Example 1 and Figure 2 specifically disclose administering *ranibizumab*, which does not satisfy the '345 claims, the Petitioner is forced to cherry-pick from Shams' laundry list of "VEGF antagonists" to find one that is purportedly recited by the claims. *See* Paper 2 at 31.

"VEGF antagonists," according to Shams, refers to a wide range of molecules — any "molecule capable of neutralizing, blocking, inhibiting, abrogating, reducing or interfering with VEGF activities." Ex. 1004 at 6:27-29. These "include anti-VEGF antibodies" as well as "antigen-binding fragments" of such antibodies, "receptor molecules" and their "derivatives," "anti-VEGF receptor antibodies" and various "VEGF receptor antagonists" including "VEGFR tyrosine kinase inhibitors." Ex. 1004 at 6:29-33. The term also includes "antagonist variants of VEGF, antisense molecules directed to VEGF, RNA aptamers specific to VEGF, and ribozymes against VEGF or VEGF receptors." *Id.* at 6:33-7:1. None of these constitute the specific type of "receptor-based chimeric molecule" required by the '345 Patent claims.

Shams lists numerous examples of its preferred antibodies, none of which can be used to practice the '345 Patent claims, including A4.6.1, bevacizumab, ranibizumab, G6, B20, 2C3 "and others as described in" ten references Shams identifies. *Id.* at 7:8-13. And "[m]ore preferably," of all the disclosed anti-VEGF antagonists, is ranibizumab — the undisputed focus of Shams. *Id* at 7:13-14.

Shams' disclosure of "VEGF-Trap (Regeneron)" appears but once in Shams and only among an extensive list of VEGF antagonists. "VEGF-Trap (Regeneron)" is not among the most preferred or even the preferred VEGF antagonists on this list. Shams makes clear that "anti-VEGF antagonistic antibodies" are to be preferred over the other categories of disclosed VEGF antagonists. Ex. 1004 at 7:6. The '345 Patent, however, does not claim the use of any antibodies. Petitioner does not attempt to explain why a POSA would select the disfavored "VEGF-Trap (Regeneron)" molecules from among all the possible VEGF antagonists disclosed in Shams and then, from among those "VEGF-Trap (Regeneron)" molecules known in the prior art, select a particular type, as required by the '345 Patent claims, that was never even disclosed by Shams. Thus, Shams discloses "VEGF-Trap (Regeneron)" molecules only as a small, disfavored portion of a much larger genus of VEGF antagonists. *See Impax Lab'ys*, 468 F.3d at 1383 (finding no anticipation of method of using riluzole because "riluzole is just one of hundreds of compounds included in formula I" of the prior art patent).

Even if Shams had specifically called out "VEGF-Trap (Regeneron)" as the molecule to use in Figure 2 — instead of relegating it to the non-preferred backwater of a vast genus of VEGF antagonists — Petitioner points to nothing in Shams to suggest that a "VEGF-Trap (Regeneron)" molecule (let alone the precise type of fusion protein required by the '345 Patent claims) could simply be substituted into the dosing protocol set forth in Shams Figure 2. Petitioner points to no disclosure in Shams to suggest a corresponding dose or dosing regimen for a "VEGF-Trap (Regeneron)" molecule. Petitioner's attempt to rewrite Shams' Example 1 and Figure 2 to replace its single-minded focus on "ranibizumab" with "VEGF-Trap (Regeneron)" (*Id.*; Ex. 1004 at 6:33) is not supported by the disclosure of Shams or the law of anticipation.

The Federal Circuit and its predecessor court have explained that anticipation requires more than merely picking and choosing from a single prior art reference to arrive at a claimed invention. *See Net MoneyIN*, 545 F.3d at 1371 (finding that district court erred in "combin[ing] parts of the separate [examples] shown in the ... reference" to conclude that a challenged claim was anticipated because "it is not enough that the prior art reference ... includes multiple, distinct teachings that the artisan might somehow combine to achieve the claimed invention."); *In re Arkley*, 455 F.2d 586, 587-88 (CCPA 1972) (A "reference must clearly and unequivocally disclose the claimed [invention] or direct those skilled in the art to the [claimed invention] without *any* need for picking, choosing, and combining various disclosures not directly related to each other by the teachings of the cited reference. Such picking and choosing ... has no place in the making of a 102, anticipation rejection.") (emphasis in original).

The Board has also repeatedly denied institution where an allegedly anticipatory reference does not disclose the claim elements as they are arranged in the challenged claim. For example, in *Coherus Biosciences, Inc. v. AbbVie Biotechnology Ltd.*, the Board denied institution because the allegedly anticipatory reference required "picking and choosing with no guidance in the prior art" to arrive at the claimed invention. IPR2017-00822, 2017 WL 3974063, at *6 (Sept. 7, 2017) (rejecting the petitioner's argument that a POSA "reading the [PCT] would 'at once envisage' the claimed arrangement or combination" because the PCT did not "teach each of the limitations of the challenged claims arranged as in the claims"); *see also Endo Pharms. Inc. v. Depomed, Inc.*, IPR2014-00655, 2014 WL 4925714, at *7 (Sept. 29, 2014) (denying institution because "Petitioner cannot combine parts of separate embodiments disclosed in the [reference] to piece together the claimed invention."); *see also Reckitt Benckiser LLC v. GEMAK Tr.*, IPR2020-00184, 2020 WL 2511249 (May 15, 2020).

As in *Coherus Biosciences*, Shams discloses a broad genus of proteins and there is nothing in Shams directing a POSA to use "VEGF-Trap (Regeneron)" over any other disclosed VEGF antagonist. *See, e.g., Coherus Biosciences Inc.*, 2017 WL 3974063, at *7 (denying institution where the allegedly anticipatory reference "provides innumerable possibilities for proteins that may provide sufficient buffering capacity" and the recited protein is not identified "in any example or as a preferred antibody."). As the Board explained in *Coherus BioSciences*, "picking and choosing with no guidance in the prior art as to which choices to make is not anticipation." 2017 WL 3974063, at *6.

Because Petitioner has done no more than piece together different disclosures from separate embodiments in Shams to argue anticipation, the Board should decline to institute Petitioner's Ground 1 anticipation challenge.

4. Petitioner Fails to Show That Shams Discloses or Enables A Therapeutically Effective Method of Treating an Angiogenic Eye Disorder

Claim 1 of the '345 Patent requires a therapeutically effective method for treating an angiogenic eye disorder. *See supra* Section IV.A. Petitioner relies on Shams' disclosure of a prophetic dosing regimen for ranibizumab in Example 1 and Figure 2 ("Treatment Schema") for its anticipation challenge. However, Shams' disclosed 12-week dosing regimen was a failure. A prior art reference cannot anticipate a claimed invention "if the allegedly anticipatory disclosures cited as prior art are not enabled." *Amgen Inc. v. Hoechst Marion Roussel, Inc.*, 314 F.3d 1313, 1354 (Fed. Cir. 2003); *see also United States v. Adams*, 383 U.S. 39, 50 (1966) (finding that an invention that is inoperable or fails to achieve its intended result does not anticipate). In addition, a POSA would not have viewed Shams as disclosing an effective method for treating an angiogenic eye disorder in a patient. Because Shams is not enabled and does not disclose a recited claim limitation — *treatment* of an angiogenic eye disorder — Shams does not anticipate the '345 Patent claims.

a. <u>Petitioner is Not Entitled to a Presumption that Shams'</u> Disclosure of a 12-Week Dosing Regimen Is Enabled

The Petition presents no evidence that Shams discloses an *effective* quarterly dosing regimen. Shams itself does not supply this evidence as it discloses only a prophetic quarterly dosing regimen. Nor can Petitioner rely on a presumption of enablement to cure this deficiency because Shams is not an issued patent.

While disclosures of prior art patents enjoy a presumption of enablement in adversarial proceedings, the Federal Circuit has not held that this same presumption applies to non-patent printed publications. *Cubist Pharms., Inc. v. Hospira, Inc.*, 75 F. Supp. 3d 641, 661 n.10 (D. Del. 2014). Indeed, this presumption should *not* to apply to non-patent printed publications. Issued patents are presumed valid because of "the basic proposition that a government agency such as the [PTO] was presumed to do its job." *Am. Hoist & Derrick Co. v. Sowa*

& Sons, Inc., 725 F.2d 1350, 1359 (Fed. Cir. 1984). Patent prosecution is an "inquisitorial process between patent owner and examiner," *see SAS Inst., Inc. v. Iancu*, 138 S. Ct. 1348, 1353 (2018), which provides a level of confidence that the patent examiner has performed his or her job in issuing a patent. There is no such "inquisitorial process" for non-patent prior art publications, especially as to whether a disclosure was enabled.

In a PGR proceeding, Petitioner bears the burden to demonstrate that it is "more likely than not" that at least one of the challenged claims is unpatentable. 35 U.S.C. § 324(a). Such an adversarial proceeding presents different prudential considerations from patent prosecution²⁰ and thus the burden to make a *prima facie*

²⁰ The burden of proving non-enablement of an allegedly anticipating reference in prosecution is on the patent applicant, who "is in a better position to show, by experiment or argument, why the disclosure in question is not enabling or operative" because "an examiner, who has no access to experts or laboratories, is not in a position to test each piece of prior art for enablement in citing it, and requiring him to do so would be onerous, if not impossible." *In re Antor Media Corp.*, 689 F.3d 1282, 1288-89 (Fed. Cir. 2012). But this rationale does not extend to adversarial PGR proceedings where a Petitioner has the resources and

case of anticipation — including the enablement of an allegedly anticipatory nonpatent reference — should be placed firmly on the Petitioner seeking to invalidate an already issued patent. *See e.g., Takeda Pharm. Co., Ltd v. Handa Pharms., LLC*, No. C-11-00840 JCS, 2013 WL 9853725, at *64-65 (N.D. Cal. Oct. 17, 2013) (concluding that the ultimate burden of proving enablement of allegedly anticipatory non-patent prior art is on the patent challenger).

Just as Petitioner bears the initial burden for establishing the "printed publication" status of a prior art reference at the institution stage by identifying "evidence sufficient to establish a reasonable likelihood that the reference was publicly accessible before the critical date of the challenged patent," *see Hulu*, IPR2018-01039, Paper 29 at 16, so too here, to satisfy its threshold burden under 35 U.S.C. § 324(a), Petitioner should bear the burden to come forward with evidence that the allegedly anticipatory disclosures of Shams are enabled. Petitioner has made no such showing.

Moreover, the factual record, including the prosecution history of Shams, evidences that no such presumption is appropriate here. Ex. 2013; Ex. 2014. Shams' disclosure and claims were repeatedly rejected during prosecution for lack

wherewithal to develop evidence, using experts and laboratories, to show enablement of non-patent prior art.

of written description and enablement; these objections were never overcome and Shams was ultimately abandoned.²¹ Ex. 2013; Ex. 2014. Thus, the basic rationale for a presumption of enablement of an issued patent should not apply to non-patent prior art publications and, indeed, is counterfactual in the case of Shams.

> b. <u>Overwhelming Evidence Demonstrates That Shams Was</u> <u>Not a Therapeutically Effective Method for Treating an</u> <u>Angiogenic Eye Disorder</u>

Shams' disclosure of a prophetic quarterly (q4/q12) dosing regimen for ranibizumab corresponds to and was tested in Genentech's Phase IIIb clinical trial called the "PIER Study." Ex. 2002 ¶¶ 27, 29-30; *see also* Ex. 2001 ¶¶ 63-64. Genentech's PIER clinical trial results, and publications reporting on those results, show that Shams' disclosed 12-week dosing regimen for ranibizumab was a failure and not an effective method for treating an angiogenic eye disorder in a patient.

The PIER study was designed to test whether ranibizumab (Lucentis) could be dosed quarterly rather than monthly and still maintain its efficacy. Ex. 1026 at 1; Ex. 2015 at 1; Ex. 2002 ¶ 27. Patients in PIER were randomized to receive 0.3 mg ranibizumab, 0.5 mg ranibizumab, or sham control by intravitreal injections

²¹ Shams (Ex. 1004) was the priority utility filing leading to U.S. continuation applications S.N. 11/738, 284, S.N. 13/780,239, and 14, 307,233 (which never published), all of which were abandoned.

administered monthly for the first three months followed by quarterly injections (every 12 weeks) through month 24 of the study. Ex. 1026 at 2.

In the first year of PIER, patients in the treatment arm gained visual acuity during the first three monthly injections of ranibizumab, but then lost all visual acuity gains after moving to fixed quarterly dosing. Ex 1026 at 7; Ex. 2002 ¶ 31; Ex. 2001 ¶ 65. This vision loss was accompanied by anatomic changes, such as increase in vascular leakage and mean retinal thickness, that are hallmark characteristics of recurrence of wet AMD. Ex. 2002 ¶ 32.

While patients in PIER were losing vision, Genentech announced results from its pivotal Phase III ANCHOR and MARINA trials that showed that fixed monthly doses of ranibizumab could improve visual acuity, and maintain those vision improvements over the course of treatment. Ex. 2002 ¶¶ 17-24. After MARINA and ANCHOR results were made public, it was no longer considered appropriate or ethical treatment to simply permit the progression of a patient's wAMD (as was done with sham control). Ex. 2002 ¶¶ 20, 23, 33; Ex. 2001 ¶ 69-70. As a consequence, the PIER protocol was amended to allow sham control subjects to cross over to the treatment arm of monthly dosing of 0.5 mg ranibizumab for the remaining year of the study. Ex. 1026 at 2; Ex. 2002 ¶ 33; Ex. 2001 ¶ 66, 69.

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It is not merely the case that the PIER regimen was less effective at improving visual acuity as compared to a monthly regimen; following the monthly loading doses, quarterly injections in PIER led to vision loss, with ultimately *zero* gains in visual acuity as compared to baseline. Ex. 1026 at 5 (Figure 1). The oneyear PIER results were so disappointing that Genentech amended the PIER study protocol yet again —this time to allow *all patients* remaining in the study the opportunity to roll over from the *12-week (quarterly) dosing* to receive 0.5 mg ranibizumab *monthly dosing* for the remainder of the two-year study. Ex. 2016 at 2; Ex. 2002 ¶ 34; Ex. 2001 ¶ 66. The fact that Genentech amended its PIER Study protocol to allow the quarterly treatment arm to roll-over to monthly dosing was an acknowledgement that PIER quarterly dosing regimen did not work. Ex. 2002, ¶ 34; Ex. 2001 ¶ 66, 69.

The PIER study was recognized as a failure in the art. Ex. 2002 ¶¶ 35-38, 43; Ex. 2001 ¶¶ 68, 70. Genentech presented PIER's One Year results in late May/early June 2006 at the Retinal Physician Symposium. Ex. 2002 ¶ 31; Ex. 2015 at 1. Dr. David Brown, the PIER investigator who first presented the data, said it was "a shock to a lot of people" that patients in the PIER study did not maintain the improvements that were seen in the MARINA and ANCHOR trials. Ex. 2017 at 2. Dr. Brown noted PIER's key take-away: "This shows that we cannot just mandatorily treat on a quarterly basis and maintain the visual gains seen with the first three monthly injections." Ex. 2017 at 1; Ex. 2002 ¶ 37. In fact, industry publications reported: "The PIER data have led Genentech to recommend that patients receive either monthly injections of ranibizumab, or have their retreatment schedules determined through individualized testing." Ex. 2015 at 1. Indeed, as discussed in Section VII.B. below, PIER was not only regarded as a failure in the art, but also as a cautionary tale against fixed quarterly dosing.

Because Petitioner has failed to show that Shams enabled or discloses a therapeutically effective method of treating an angiogenic eye disorder, Shams cannot anticipate.

5. Shams Fails to Disclose A Tertiary Dose That Maintains the Therapeutic Effect Throughout the Course of Treatment When Administered 12 Weeks After the Immediately Preceding Dose

Claim 1 requires a tertiary dose that maintains therapeutic effect when administered 12 weeks after the immediately preceding dose. *Supra*, pp. 10-12. But Shams' dosing regimen failed to maintain a therapeutic benefit during the quarterly dosing phase of the regimen. *Supra*, pp. 62-64.

As noted in Section VI.4.3. above, when the Shams treatment schema of Figure 2 was tested in the PIER Study, all visual acuity gains from the first three monthly doses of ranibizumab were lost once injections were reduced to quarterly (12-week) administration. Ex. 1026 at 5 (Figure 1). As discussed above, the PIER

Study was a failure and was consistently characterized as an ineffective treatment regimen in the art.

Because Petitioner has failed to show that Shams discloses "a tertiary dose that maintains a therapeutic effect throughout the course of treatment when administered 12 weeks after the immediately preceding dose," Petitioner has failed to make its threshold showing for Ground 1.

B. Ground 2: Petitioner Fails to Demonstrate That It Is More Likely Than Not That At Least One of the Challenged Claims Is Obvious

According to the Petition, the 2009 Press Release in view of Shams renders the '345 Patent claims obvious. Not so.

As a threshold matter, Petitioner fails to address, let alone overcome, Regeneron's showing during prosecution that the claimed dosing regimen exhibits unexpected results. This omission is fatal to Petitioner's obviousness argument. To make matters worse, neither the 2009 Press Release nor Shams teaches an *effective* method of treating an angiogenic eye disorder with a quarterly dosing regimen of the recited VEGF antagonist. Nor does either provide a reasonable expectation of success for achieving an effective method of treatment. To the contrary, Shams' quarterly dosing regimen with Lucentis was an acknowledged failure. Accordingly, Petitioner has failed to "demonstrate that it is more likely than not that at least 1 of the claims challenged in the petition is unpatentable," and thus, denial of the petition is warranted. 35 U.S.C. §§ 322(a)(3), 324(a).

1. The Board Should Deny Institution Under 35 U.S.C. § 324(a) Because Petitioner Has Failed to Address, Let Alone Overcome, the Examiner's Finding of Unexpected Results

As discussed in the Petition, during prosecution, Regeneron overcame an obviousness-type double patenting rejection based on unexpected results demonstrating that extended dosing of aflibercept was noninferior to the existing therapy, which as of the priority filing date of the 345 Patent, was fixed monthly doses of ranibizumab. Ex. 1002 at 255. To overcome an obviousness-type double patenting rejection during prosecution, Regeneron relied on a 2012 paper by Heier *et al.* (Ex. 1012) to demonstrate that with extended dosing, *i.e.*, dosing less frequent than monthly, "it is possible to treat angiogenic eye disorders such as AMD while administering doses on a less frequent basis than previously thought possible." *Id.* at 256. Based on Regeneron's response, the Examiner withdrew the obviousness-type double patenting objection (*id.* at 225-226) and later allowed the claims. *Id.* at 145-46.

Although Petitioner acknowledges that Regeneron presented evidence of unexpected results during prosecution (*see, e.g.*, Paper 2 at, 36, 60, 62), nowhere does the Petition overcome, or even substantively address, these arguments. This omission is fatal to Petitioner's Ground 2. The Board has consistently denied institution where a petitioner fails to address secondary indicia of non-obviousness, such as unexpected results, in the prosecution history. *See, e.g., Coalition for*

Affordable Drugs V LLC v. Hoffmann-La Roche Inc., IPR2015-01792, Paper 14 at 18 (Mar. 11, 2016) (denying institution because "the unrebutted objective indicia of nonobviousness presented in the prosecution history of the [challenged] patent ... supports the non-obviousness of the challenged claims" and "[t]he Petition ... should have addressed the evidence of unexpected results as part of Petitioner's showing of a reasonable likelihood of success on the merits."); see also Omron Oilfield & Marine, Inc. v. MD/Totco, a division of Varco, L.P., IPR2013-00265, Paper 11 at 16 (Oct. 31, 2013) (denying institution where Petitioner did not in its Petition challenge the merits of commercial success evidence Patent Owner developed in reexamination involving different prior art references than those asserted in IPR); see Gilead Scis., Inc. v. United States, IPR2019-01453, Paper 14 at 30 (Feb. 20, 2020) (finding that "Petitioner's failure to persuasively address [the showing of unexpected results during prosecution] in the Petition means that Petitioner falls short of its burden to establish a reasonable likelihood of success in prevailing on its challenge"); Stryker Corp. v. KFXMed., LLC, IPR2019-00817, Paper 10 at 29 (Sept. 16, 2019) (finding Petitioner's "failure to address the known evidence of secondary considerations" in a related proceeding "weighs in favor of denying institution"); Robert Bosch Tool Corp. v. SD3, LLC, IPR2016-01751, Paper 15 at 23-24 (Mar. 22, 2017) ("We have cautioned petitioners in prior proceedings that known evidence of secondary considerations should be addressed in the petition.").

Here, Petitioner fails to address Regeneron's showing of unexpected results during prosecution. The only reference to Regeneron's showing of unexpected results in the Petition is an *acknowledgment* by the Petitioner that Regeneron put forth evidence of unexpected results during prosecution. *See, e.g.* Paper 2 at 36 ("Heier (which Patent Owner cited as evidence of 'unexpected results' during prosecution) ..."); *see also id.* at 60, 62. At least in *Coalition for Affordable Drugs* and *Omron Oilfield & Marine*, the petitioners *attempted*, albeit insufficiently, to overcome the record evidence of objective indicia of non-obviousness. Here, where Petitioner has utterly failed to address Regeneron's showing of unexpected results, the Board should decline to institute for the same reasons it declined to institute in *Coalition for Affordable Drugs*, *Omron Oilfield & Marine*, *Gilead Scis.*, *Stryker Corp*, and *Robert Bosch Tool Corp*.

2. A POSA Would Not Reasonably Expect to Treat an Angiogenic Eye Disorder Using the Claimed Regimen by Combining the 2009 Press Release and Shams

Petitioner argues that the '345 Patent claims are rendered obvious by the 2009 Press Release in view of Shams. But a claim is not rendered obvious "merely by demonstrating that each of its elements was, independently, known in the prior art." *KSR Int'l Co. v. Teleflex Inc.*, 550 U.S. 398, 418 (2007). Rather, Petitioner must "demonstrate ... that the skilled artisan would have had a reasonable expectation of success in" "combin[ing] the teachings of the prior art references." *Intelligent Bio*- *Sys., Inc. v. Illumina Cambridge Ltd.*, 821 F.3d 1359, 1367-68 (Fed. Cir. 2016); *see also KVK-Tech, Inc. v. Shire PLC*, IPR2018-00290, 2019 WL 2884463, at *7 (July 3, 2019) (same).

Petitioner fails to show that a POSA would have had a reasonable expectation of success in treating an angiogenic eye disorder with the claimed dosing regimen by combining the 2009 Press Release and Shams. In Ground 2, Petitioner relies on Shams for the tertiary dosing limitation only because Shams discloses "tertiary doses every three months" with ranibizumab. Paper 2 at 48. In Petitioner's own words, it is "merely combin[ing] prior art elements (Shams' 12-week dosing) to a known method (the 2009 Press Release's 4-week secondary dosing plus 12-week tertiary dosing) to arrive at a predicate result." Paper 2 at 49.

But the Board has denied institution where a petitioner fails to make a threshold showing that a POSA would have had a reasonable expectation of success in combining asserted prior art references to achieve the claimed invention. *See, e.g., Thermo Fisher Sci. Inc. v. Regents of the Univ. of Calif.*, IPR2018-01156, 2018 WL 6318146, at *10 (Dec. 3, 2018) (declining to institute review "[b]ecause an obviousness argument cannot succeed without Petitioner establishing a reasonable likelihood that a person of ordinary skill in the art would have had a reason to combine the relevant references" and "demonstrating that each of [a claim's] elements was, independently, known in the prior art" is not sufficient) (quoting *KSR*

Int'l, 550 U.S. at 403 (internal quotation marks omitted); *see also Illumina, Inc. v. Complete Genomics, Inc.*, IPR2020-00079, Paper 7 (Apr. 22, 2020) (denying institution in part because petitioner failed to "sufficiently show[] that one of ordinary skill in the art would have had a reasonable expectation of success"); *NOF Corp. v. Nektar Therapeutics*, IPR2019-01392, Paper 23 (Feb. 4, 2020) (same). Without demonstrating that a POSA would have had a reasonable expectation of success in treating an angiogenic eye disorder using the claimed dosing regimen, Petitioner fails to demonstrate that the '345 Patent claims are obvious.

Not only does Petitioner fail to make the requisite showing for a reasonable expectation of success, but it is clear that a POSA would *not* have had a reasonable expectation of success in treating an angiogenic eye disorder with the claimed dosing regimen based on the 2009 Press Release and Shams.

First, the 2009 Press Release does not disclose an effective method for treating an angiogenic eye disorder using a quarterly tertiary dosing regimen. In the 2009 Press Release, Regeneron announced that it had completed enrollment in two randomized, double-masked Phase III clinical trials, called VIEW 1 and VIEW 2, evaluating "VEGF Trap-Eye" for the treatment of wet AMD. Ex. 1005 at 1. The Press Release reported that the first year of the VIEW studies would involve dosing patients with VEGF Trap-Eye either monthly or every two months, after three monthly doses, and in the second year "patients will continue to be followed and

treated for another year on a flexible, criteria-based extended PRN [*pro re nata* — taken as needed] regimen with a dose administered at least every 12 weeks, but not more often than every four weeks until the end of the study." *Id.*

Notably, the Press Release does not disclose that any patient was dosed, nor does it disclose that any patient had started the second year of PRN dosing. *Id.* Additionally, the Press Release does not report any results but does report that "[o]ne-year primary endpoint data from both studies are expected in the fourth quarter of 2010." *Id.* Thus, not only does the 2009 Press Release fail to disclose quarterly dosing — which Petitioner acknowledges, Paper 2 at 48; Ex. 1003 ¶ 108 — but it does not disclose the *treatment* of any angiogenic eye disorder with any dosing regimen. The 2009 Press Release discloses a *prospective* study, for which enrollment is complete.²² Accordingly, the 2009 Press Release does not disclose

²² The Petition incorrectly suggests that the 2009 Press Release reports on a study that has been completed: "The 2009 Press Release teaches, among other arms, 'that patients *received* intravitreal doses of 0.5 mg or 2g VEGF Tap-Eye [sic] at 4–week intervals in the first year, *followed* by continual treatment for another year on a flexible, PRN regiment [sic], with a dose administered at least every 12 weeks." Paper 2 at 42 (emphases added). Although the statement includes a citation to the Press Release, the quoted language is not found in the Press Release.

treatment of any disease. In fact, results from the first year of the study were not even expected until "the fourth quarter of 2010," over a year after the publication of the Press Release. Moreover, the 2009 Press Release does not make any statements about the efficacy of VEGF-Trap-Eye. Nor could it. Indeed, FDA prohibits preapproval promotion of investigational drugs. 21 C.F.R. § 312.7(a) ("A sponsor or investigator, or any person acting on behalf of a sponsor or investigator, shall not represent in a promotional context that an investigational new drug is safe or effective for the purposes for which it is under investigation or otherwise promote the drug."). Accordingly, a POSA would not have understood the disclosure or announcement of a prospective Phase III trial to provide a reasonable expectation of success.

Second, a POSA would not have had a reasonable expectation of success in combining the 2009 Press Release and Shams to achieve the claimed treatment regimen because a POSA would have recognized that the 12-week ranibizumab dosing regimen disclosed in Shams would not treat an angiogenic eye disorder.²³ Petitioner argues that "[i]t would have been natural for one of skill in the art to look at Shams' teachings when considering the 2009 Press Release's 4 + 8 week dosing" Paper 2 at 48. But Petitioner does not demonstrate that a POSA would

²³ Petitioner asserts that "success was predictable because Shams teaches a successful 4 + 12 week dosing regimen..." (Paper 2 at 49-50) but fails to cite evidence to support this assertion. Moreover, as detailed in Ex. 2002 (¶¶ 29-43) and Ex. 2001 (¶¶ 63-70), Shams' 4 +12 week dosing regimen was a failure and recognized as such in the art. Petitioner also asserts that "Regeneron publicly announced that VEGF Trap was successful in quarterly doses" and cites paragraph 114 of Dr. Wu's declaration in support. Paper 2 at 50 & n.148. But Dr. Wu's testimony does not support a finding of obviousness. *First*, Dr. Wu's declaration at paragraph 114 does not appear to be cite the correct document (Ex. 1030, not Ex. 1031). Regardless, neither Exhibit is properly cited in the Petition. *Second*, even if Petitioner had properly cited Exs. 1030 and 1031, these documents merely purport to provide results for patients who received *a single dose* of VEGF Trap Eye, not the claimed dosing regimen. Reported success after a single administration does not render obvious the claimed dosing regimen.

have had a reasonable expectation of success in combining Shams with the 2009 Press Release to achieve an effective treatment regimen, nor could it.

As discussed *supra*, Shams' prophetic disclosure of a quarterly dosing regimen for ranibizumab was tested in Genentech's PIER trial, which was widely-regarded as a failure. *See supra* Section VII.A.4. Moreover, following the disclosure of PIER clinical trial results, peer-reviewed publications reflect the perception that fixed quarterly dosing and even fixed extended dosing should be approached with caution in view of the clinical results in PIER:

- "A recent analysis of the ANCHOR, MARINA, and PIER data demonstrated that monthly intravitreal ranibizumab dosing significantly reduced the frequency of macular hemorrhages... The effect was lost when patients were switched from monthly to quarterly dosing in the PIER study. *Reducing the frequency of injections should, therefore, be done with caution.*" Ex. 2018 at 5, emphasis added.
- "In the PIER trial, when patients were switched from monthly to quarterly injections of ranibizumab, they subsequently lost the vision they had gained with monthly injections." Ex. 2019 at 1.
- "In PIER, no benefit of ranibizumab over sham was observed after the patients were switched to the quarterly protocol. ... As seen in PIER, switching from monthly to quarterly injection intervals may not have the same beneficial effect and could put the patient at an increased risk for vision threatening complications." Ex. 2020 at 5, emphasis added.
- "However, fixed quarterly [citing PIER] or 'as needed' (pro re nata [PRM]) dosing regimens, without requiring monthly monitoring visits were not effective at maintaining vision." Ex. 2021 at 1.

After PIER, no retina physician would treat his or her wAMD patient with fixed quarterly dosing of ranibizumab. Ex. 2001 ¶ 70; Ex. 2002 ¶ 43.

Until Regeneron conducted its Phase III pivotal trials on aflibercept in wAMD, there remained an unmet need for extended dosing regimens of VEGF antagonists for the treatment of angiogenic eye disorders, despite "extensive efforts to decrease injection and monitoring frequency." Ex. 2021 at 9. Shams not only fails to provide a reasonable expectation that a fixed quarterly dosing regimen would work, but rather, is evidence of long-felt need and failure of others to develop such an extended dosing regimen. Simply put, a POSA would not have had a reasonable expectation of success in using quarterly dosing to treat angiogenic eye disorders and would have been discouraged by the results of the PIER study. Thus, Petitioner is incorrect to suggest that "Shams teaches a *successful* 4 + 12 week dosing regimen." Paper 2 at 49-50 (emphasis added).

Accordingly, neither the 2009 Press Release nor Shams, alone or together, would have provided a POSA with a reasonable expectation of success that a 12-week dosing regimen would work for the VEGF antagonists recited by the '345 Patent claims. Without more, merely plucking the tertiary quarterly dosing regimen from Shams and combining it with a prospective trial in the 2009 Press Release, does not render the '345 Patent claims obvious.

* * *

For the reasons discussed above, Petitioner has not met its burden under 324(a) and 322(a) to demonstrate that it is more likely than not that at least one claim will be found unpatentable as obvious over the 2009 Press Release in view of Shams. Accordingly, the Board should deny institution.

C. Ground 3: Petitioner Fails to Demonstrate That It Is More Likely Than Not That At Least One of the Challenged Claims Lacks Written Description

Claim 1, the only independent claim of the '345 Patent, requires that "each tertiary dose is administered 12 weeks after the immediately preceding dose." Ex. 1001 at Claim 1. Petitioner concedes that the "345 patent mentions 12–week tertiary dosing." Paper 2 at 66. This "mention" also appears in the priority applications. Although Petitioner argues that the '345 Patent is invalid for lack of written description because it discloses too many different dosing regimens, Petitioner fails to meet its burden of demonstrating that it is more likely than not that at least one of the challenged claims lacks written description.

Petitioner does not cite any case invalidating claims merely because the specification discloses too many examples. Paper 2 at Sec. VIII. Instead, Petitioner relies on cases where the specification did not disclose the claimed embodiment and, at best, support could be found only by selectively piecing together portions of the specification with no blazemarks that pointed to the claimed embodiment. *Id.* at 65-71 (citing *Novozymes, Purdue Pharma L.P., Ruschig, Fujikawa, FWP IP ApS*,

Boston Sci. Corp.). Those cases have no applicability to the '345 Patent because the priority applications here disclose as one example the precise embodiment covered by the claims. If a patent specification discloses multiple examples, and one of those examples is the claimed embodiment, as it is here, the claim is supported. *Erfindergemeinschaft UroPep v. Eli Lilly & Co.*, 276 F. Supp. 3d 629, 656 (E.D. Tex. 2017), *aff'd*, 739 F. App'x 643 (hereinafter "*UroPep*").

Accordingly, because the challenged claims are adequately supported by the specification, Petitioner has not demonstrated at least one of the challenged claims is unpatentable for lack of written description.

1. Petitioner Fails to Demonstrate That the Claims More Likely Than Not Lack Written Description Because the '345 Patent Specification Discloses as a Specific Example the Exact Dosing Regimen Claimed

Section 112 does not prescribe the manner in which a specification must support a claim, so long as the specification "clearly allow[s] persons of ordinary skill in the art to recognize that [the inventor] invented what is claimed." *Ariad Pharms.*, *Inc. v. Eli Lilly & Co.*, 598 F.3d 1336, 1351 (Fed. Cir. 2010) (en banc) (alteration in original) (internal quotation marks omitted).

The Petitioner cannot carry its burden because the '345 Patent specification discloses that "[i]n one *exemplary* embodiment of the present invention, *each* secondary dose is administered 2 to 4 (e.g., 2, 2 $\frac{1}{2}$, 3, 3 $\frac{1}{2}$, or 4) weeks after the *immediately preceding dose, and each tertiary dose is administered* at least 8 (e.g.,

8, 8 ¹/₂, 9, 9 ¹/₂, 10, 10 ¹/₂, 11, 11 ¹/₂, *I2*, 12 ¹/₂, 13, 13 ¹/₂, 14, 14 ¹/₂, or more) *weeks after the immediately preceding dose*." Ex. 1001 at 3:57-62 (emphases added). Accordingly, the '345 Patent specification specifically discloses that a "secondary dose is administered ... 4 ... weeks after the immediately preceding dose, and each tertiary dose is administered ... 12 ... weeks after the immediately preceding dose." *Id.* This is also known as a q4/q12 dosing regimen. Thus, the specification discloses the q4/q12 dosing regimen of Claim 1 as an "exemplary embodiment." *Id.* at 3:57. This exact same disclosure is found in the 2011 Provisional Applications to which the '345 Patent claims priority; therefore, there can be no doubt that the claims are supported by the 2011 Provisional Applications. *See* Ex. 1045 at [0016]; Ex. 1046 at [0016]; Ex. 1047 at [0016].

Petitioner argues that this disclosure is insufficient because the patent discloses other dosing regimens as well. Paper 2 at 68. A court addressing a very similar set of circumstances explained that "[a] patentee is free to selectively claim one particular embodiment without running afoul of the written description requirement." *UroPep*, 276 F. Supp. 3d at 656, *aff'd* 739 F. App'x 643. Indeed, "[i]t is common for patentees to disclose a range of possible embodiments" and "a patentee need not indicate that one embodiment is 'of special interest' in order to claim it." *Id.*

In *UroPep*, the patent claimed a method for prophylaxis or treatment of benign prostatic hyperplasia (BPH) with a phosphodiesterase (PDE) 5 inhibitor. *Id.* The patent challenger argued that the written description was not sufficient because the specification described other PDE inhibitors, such as PDE1 and PDE4 inhibitors, as well as using all three PDE inhibitors (PDE1, PDE4, and PDE5) to treat BPH; and the specification taught the treatment of other prostatic diseases. *Id.* at 655. Thus, the challenger argued that the specification lacked written description because it did not provide blazemarks leading one specifically to the treatment of *BPH* with a *PDE s* inhibitor. *Id.* at 655. But, because "selectively claim[ing] one particular embodiment" does not "run[] afoul of the written description requirement," the court found the claims not invalid for lack of written description. *Id.* at 656.

Petitioner does not cite a single case that runs counter to *UroPep*, invalidating or rejecting a claim supported by a specific example in the specification. Here, there can be no dispute that disclosing the q4/q12 dosing regimen permits a skilled artisan to recognize that the inventors possessed the claimed q4/q12 dosing regimen. Accordingly, Petitioner has failed to meet its burden of demonstrating that at least one challenged claim is unpatentable for lack of written description.

2. None of Petitioner's Cases Involve a Specification That Actually Discloses the Claimed Species

Petitioner relies on cases where courts found the specification lacked sufficient "blazemarks" to direct the skilled artisan to the claimed invention because it disclosed a genus. See Paper 2 at 65-70. For example, in *Ruschig*, the challenged claim recited a chemical compound. The specification, however, merely disclosed a broad genus that included the claimed compound along with "half a million" others.²⁴ In re Ruschig, 379 F.2d 990, 993 (CCPA 1967). The '345 priority applications, unlike the specification in *Ruschig*, actually disclosed the claimed q4/q12 dosing regimen as an "exemplary" embodiment.

Petitioner argues that the '345 Patent discloses a genus of tens of thousands of dosing regimens, but that is not correct.²⁵ The specification does not merely say

-SO3-NH-CO-NH-Ra

²⁵ Petitioner misleadingly cites to a statement that Regeneron made during prosecution to argue that the '345 patent specification discloses a "virtually infinite" number of dosing regimens. Paper 2 at 68. During prosecution, to overcome a double patenting rejection, Regeneron explained that as of the filing date, the standard of care was monthly dosing but that there was a need in the art to extend that dosing and there were "virtually an infinite number of different treatment

²⁴ The specification in *Ruschig* disclosed a genus that includes multiple unspecified substituents (R, R₁, R₂), as shown below, as opposed to a single compound:

the secondary dose must be between 2 and 4 weeks and the tertiary dose must be between 8 or more weeks. It provides specific examples which include q4/q12, as explained above.²⁶ Because the '345 patent discloses q4/q12 as an example, there is no need for "blazemarks." *UroPep*, 276 F. Supp. 3d at 656. To use Petitioner's analogy, the '345 Patent specification does not "disclose[] a forest" (Paper 2 at 66), it discloses many "tree[s]."

All of Petitioner's other cases are likewise inapplicable. *See, e.g., Fujikawa v. Wattanasin*, 93 F.3d 1559, 1571 (Fed. Cir. 1996) (finding patent invalid for lack of written description because there was no disclosure of the recited sub-genus, a

²⁶ Moreover, as Petitioner acknowledges, in addition to teaching that the secondary and tertiary doses can be administered at "different frequencies," the patent teaches they can be administered at the "same" frequency. Paper 2 at 67; Ex. 1001 at 4:23-34. In the latter case, where the timing of secondary and tertiary doses are fixed, the patent discloses only 70 examples: 5 exemplary secondary doses and 14 exemplary tertiary doses (5 x 14 = 70).

protocols that could be tested." Ex. 1002 at 255. But, in making this statement, Regeneron was not talking about specific dosing regimens disclosed in the '345 specification, as Petitioner implies, but all dosing regimens that theoretically could be tested. *Infra* Section VII.C.3.

species within the sub-genus, or even a suggestion that the sub-genus was "of special interest"); *Boston Sci. Corp. v. Johnson & Johnson*, 647 F.3d 1353, 1367 (Fed. Cir. 2011) (same); *see also FWP IP ApS v. Biogen MA, Inc.*, 749 F. App'x 969, 975 (Fed. Cir. 2018) (finding claims invalid for lack of written description because specification did not teach that the recited dose could effectively treat multiple sclerosis, as recited by the limitation at issue); *Novozymes A/S v. DuPont Nutrition Bioscis. APS*, 723 F.3d 1336, 1348, 1341 (Fed. Cir. 2013) (finding claims directed to enzyme with particular properties invalid for lack of written description because there was no disclosure of a variant enzyme in the specification with all recited properties); *Purdue Pharma L.P. v. Faulding Inc.*, 230 F.3d 1320, 1327 (Fed. Cir. 2000) (finding written description not adequate where claims recited ratio of two pharmacokinetic parameters and nowhere was the recited ratio "discussed even in passing in the disclosure").²⁷

²⁷ The *Uropep* court similarly distinguished many of the cases on which Petitioner relies. *UroPep*, 276 F. Supp. 3d at 655-656 (distinguishing *Novozymes*, *Boston Scientific*, *Fujikawa*, and *Ruschig*).

3. Petitioner's Argument Based on the Prosecution History Is Legally Irrelevant to Written Description and Factually Unsupported

Petitioner argues that (1) Regeneron took inconsistent positions during prosecution and (2) the '345 Patent claims lack written description for the same reason that Regeneron argued the 2009 Press Release did not anticipate during prosecution. Paper 2 at Sec. VIII.C. Neither argument has merit.

Regeneron's statements made during prosecution to overcome prior art are not legally relevant to the issue of whether the claimed dosing regimen is adequately supported by the '345 Patent specification. *Ariad*, 598 F.3d at 1351 ("the test [for written description] requires an objective inquiry into the four corners of the specification from the perspective of a person of ordinary skill in the art."). Petitioner cites no cases to support that proposition, nor does it even attempt to explain how they could be relevant. Paper 2 at Sec. VIII.B. This alone disposes of the issue.

Even if Regeneron's prosecution history statements were legally relevant, they are not inconsistent. During prosecution, the Examiner rejected the claims challenged by the Petition based on non-statutory double patenting. Ex. 1002 at 324-329. Regeneron argued that the pending claims to particular dosing regimens were nonobvious over some of the patents. *Id.* at 282-287. It argued that there was a need in the art to improve on the standard of care for wAMD by providing a method requiring less frequent dosing than the prevailing fixed monthly dosing of ranibizumab. *Id.* at 283. Less frequent dosing would reduce the treatment burden of monthly intravitreal injections, including the expense, pain and inconvenience to the patient and physician. *Id.* Regeneron argued its less frequent dosing method was not *prima facie* obvious because there were "virtually an infinite number of different treatment protocols that could be tested." *Id.* Furthermore, even if the claimed invention were *prima facie* obvious, that finding would be overcome because the claimed invention exhibited unexpected results, as demonstrated by a 2012 paper by Heier *et al.* (Ex. 1012) ("Heier").²⁸ *Id.* at 284-285. Heier demonstrated that extended dosing of aflibercept (*i.e.*, less frequent than every month) "would be surprisingly as good or better than the results obtained with monthly treatment" [of prior art anti-VEGF therapies]. *Id.* at 285. The Examiner withdrew the obviousness-type double patenting rejection in view of Regeneron's unexpected results arguments. *Id.* at 225-226.

The Examiner later rejected the 12-week dosing claims as anticipated by the 2009 Press Release. According to the Examiner, the 2009 Press Release's disclosure of a "flexible PRN schedule" "would include at least one tertiary dose at 12 weeks from the immediately preceding dose." *Id.* Regeneron, however, explained that the

²⁸ Heier was published in 2012 and thus is not prior art to the '345 Patent claims.

2009 Press Release did not disclose the claim limitation "tertiary dosing administered 12 weeks after the immediately preceding dose." *Id.* at 210-211. The only reference to "12 weeks" in the Press Release was a reference "to a flexible, criteria-based extended PRN regimen with a dose administered at least every 12 weeks, but not more often than every four weeks"; the "[m]ere mention of a prospective possibility of dosing at 12 weeks does not explicitly indicate or teach towards a method where 12-week dosing would be undertaken, let alone successful." *Id.* at 176-177. The Examiner allowed the claims.

Petitioner argues that Regeneron took "inconsisten[t]" positions when it relied on "Heier (to overcome the first rejection) as sufficiently disclosing the claimed regimen to support unexpected results" and later critiqued "the 2009 Press Release as insufficient to disclose the same regimen" to overcome the second rejection. Paper 2 at 62. Petitioner's argument is wrong for several reasons.

Regeneron never argued that Heier "disclose[s] the claimed regimen." *Id.* Rather, Regeneron explained in prosecution that Heier supported the proposition that extended dosing regimens (such as those covered by the then-pending claims) were unexpectedly noninferior to the prevailing standard of care (*i.e.*, monthly injections of ranibizumab). *Supra*, p. 59. Moreover, even if Regeneron had argued that Heier disclosed the claimed regimen — contrary to the prosecution history — that would not have been inconsistent with Regeneron's statements regarding the 2009 Press Release. Petitioner incorrectly asserts that the "2009 Press Release has essentially the same description as Heier." Paper 2 at 61. Petitioner ignores that Heier 2012 reports on the results of a clinical trial that had not been completed and were not known in 2009. By Heier 2012, both the frequency of dosing in the PRN regimen and the clinical trial results demonstrating efficacy were known, whereas at the time of the 2009 Press Release, neither of those things were known. Accordingly, there was nothing inconsistent in Regeneron's explanation that the Press Release failed to disclose 12-week dosing and that Heier demonstrated the success of extended dosing.

* * *

Petitioner's argument that the '345 Patent claims are invalid for lack of written description are not persuasive and Petitioner has not established that it is more likely than not that any claim challenged in Ground 3 will be found unpatentable.

VIII. CONCLUSION

For the reasons discussed *infra*, because Petitioner has failed to show: (1) that the '345 Patent claims are PGR-eligible; (2) that the Examiner materially erred in considering the same art or argument in prosecution; and (3) that any challenged claim is more likely than not unpatentable based on Grounds 1, 2, or 3, the Board should deny institution of the Petition. Dated: April 15, 2021

Respectfully Submitted,

/s/ Deborah E. Fishman

Deborah E. Fishman (Reg. No. 48,621) Arnold & Porter Kaye Scholer LLP 5 Palo Alto Square, Suite 500 3000 El Camino Real Palo Alto, CA 94306

Counsel for Patent Owner, Regeneron Pharmaceuticals, Inc.

CERTIFICATE OF COMPLIANCE

The undersigned certifies that this preliminary response complies with the type-volume limitations of 37 C.F.R. 42.24(a)(1)(i). This preliminary response contains 18,047 words as calculated by the "Word Count" feature of Microsoft Word 2010, the word processing program used to create it.

The undersigned further certifies that this preliminary response complies with the typeface requirements of 37 C.F.R. § 42.6(a)(2)(ii) and typestyle requirements of 37 C.F.R. § 42.6(a)(2)(iii). This preliminary response has been prepared in a proportionally spaced typeface using Microsoft Word 2010 in Times New Roman 14 point font.

/s/ Deborah E. Fishman

Deborah E. Fishman (Reg. No. 48,621) Arnold & Porter Kaye Scholer LLP 5 Palo Alto Square, Suite 500 3000 El Camino Real Palo Alto, CA 94306

CERTIFICATE OF SERVICE

Pursuant to 37 CFR §§42.6(e)(4)(i) et seq. and 42.205(b), the undersigned

Certifies that on April 15 2021, a true and entire copy of this PRELIMINARY

RESPONSE OF PATENT OWNER REGENERON PHARMACEUTICALS,

INC., and all supporting exhibits, were served via e-mail to the Petitioner at the

following email addresses:

CHENGDU-IPR@mofo.com mkreeger@mofo.com jxiao@mofo.com dosullivan@mofo.com

/s/ Deborah E. Fishman

Deborah E. Fishman (Reg. No. 48,621) Arnold & Porter Kaye Scholer LLP 5 Palo Alto Square, Suite 500 3000 El Camino Real Palo Alto, CA 94306 Patent No. 10,828,345 Petition for Post Grant Review

UNITED STATES PATENT AND TRADEMARK OFFICE

BEFORE THE PATENT TRIAL AND APPEAL BOARD

CHENGDU KANGHONG BIOTECHNOLOGY CO., LTD., Petitioner,

v.

REGENERON PHARMACEUTICALS, INC., Patent Owner.

Patent No. 10,828,345

Post Grant Review No. PGR2021-00035

DECLARATION OF DAVID WU, M.D., PH.D

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EXHIBITS

Exhibit Description	<u>No.</u>
U.S. Patent No. 10,828,345	1001
File History of U.S. Application No. 16/159,282 (U.S. Patent No. 10,828,345)	
International Publication No. WO 2006/047325 (May 4, 2006) to Shams	
Regeneron "Press Release Dated September 14, 2009" (September 14, 2009)	1005
Genentech "News Release dated June 30, 2006" (June 30, 2006)	1006
Regeneron "Press Release dated November 18, 2011" (November 18, 2011)	1007
"Safety and Tolerability Study of Intravitreal VEGF-Trap Administration in Patients With Neovascular AMD" (NCT00320775)	1008
Nguyen et al., "Results of a Phase 1, Dose-Escalation, Safety, Tolerability, and Bioactivity Study of Intravitreous VEGF Trap in Patients with Neovascular Age-Related Macular Degeneration" ARVO Annual Meeting Abstract (May 1 2006)	1009
Benz et al., "CLEAR-IT-2: Interim Results Of The Phase II, Randomized, Controlled Dose-and Interval-ranging Study Of Repeated Intravitreal VEGF Trap Administration In Patients With Neovascular Age-related Macular Degeneration (AMO)" ARVO Annual Meeting Abstract (May 2007)	1010
U.S. Application No. 13/940,370	1011
<i>Heier et al.</i> , "Intravitreal Aflibercept (VEGF Trap-Eye) in Wet Age- related Macular Degeneration," Ophthalmology, Volume 119, Number 12, December 2012, Page 2538	1012

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Exhibit Description	<u>No.</u>
<i>Heier et al</i> , "The 1-year Results of CLEAR-IT 2, a Phase 2 Study of Vascular Endothelial Growth Factor Trap-Eye Dosed As-needed After 12–week Fixed Dosing," Ophthalmology, Volume 118, Number 6, June 2011, at 1110	1013
File History of U.S. Application No. 14/934,731	1014
Curriculum Vitae of Dr. David Wu, M.D., Ph.D	1015
PCT Application No. PCTUS1220855	1016
Macular Photocoagulation Study, G. (1991). "Laser photocoagulation of subfoveal neovascular lesions in age-related macular degeneration. Results of a randomized clinical trial. Macular Photocoagulation Study Group." <u>Arch Ophthalmol</u> 109 (9): 1220-1231	1017
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Gragoudas, E. S., A. P. Adamis, E. T. Cunningham, Jr., M. Feinsod, D. R. Guyer and V. I. S. i. O. N. C. T. Group (2004). "Pegaptanib for neovascular age-related macular degeneration." <u>N Engl J Med</u> 351 (27): 2805-2816.	1019
Michels, S., P. J. Rosenfeld, C. A. Puliafito, E. N. Marcus and A. S. Venkatraman (2005). "Systemic bevacizumab (Avastin) therapy for neovascular age-related macular degeneration twelve-week results of an uncontrolled open-label clinical study." <u>Ophthalmology</u> 112 (6): 1035-1047.	1020
Rosenfeld, P. J., A. A. Moshfeghi and C. A. Puliafito (2005). "Optical coherence tomography findings after an intravitreal injection of bevacizumab (avastin) for neovascular age-related macular	1021

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Exhibit Description	<u>No.</u>
degeneration." Ophthalmic Surg Lasers Imaging 36(4): 331-335	
Avery, R. L., D. J. Pieramici, M. D. Rabena, A. A. Castellarin, M. A. Nasir and M. J. Giust (2006). "Intravitreal bevacizumab (Avastin) for neovascular age-related macular degeneration." <u>Ophthalmology</u> 113 (3): 363-372 e365	1022
Wall Street Journal 2007 – "Genentech's Big Drug for Eyes Faces a Rival"	1023
Fung, A. E., G. A. Lalwani, P. J. Rosenfeld, S. R. Dubovy, S. Michels, W. J. Feuer, C. A. Puliafito, J. L. Davis, H. W. Flynn, Jr. and M. Esquiabro (2007). "An optical coherence tomography-guided, variable dosing regimen with intravitreal ranibizumab (Lucentis) for neovascular age-related macular degeneration." <u>Am J Ophthalmol</u> 143 (4): 566-583.	1024
Gupta, O. P., G. Shienbaum, A. H. Patel, C. Fecarotta, R. S. Kaiser and C. D. Regillo (2010). "A treat and extend regimen using ranibizumab for neovascular age-related macular degeneration clinical and economic impact." Ophthalmology 117(11): 2134-2140	1025
Regillo, C. D., D. M. Brown, P. Abraham, H. Yue, T. Ianchulev, S. Schneider and N. Shams (2008). "Randomized, double-masked, sham-controlled trial of ranibizumab for neovascular age-related macular degeneration: PIER Study year 1." <u>Am J Ophthalmol</u> 145 (2): 239-248.	1026
Schmidt-Erfurth, U., B. Eldem, R. Guymer, J. F. Korobelnik, R. O. Schlingemann, R. Axer-Siegel, P. Wiedemann, C. Simader, M. Gekkieva, A. Weichselberger and E. S. Group (2011). "Efficacy and safety of monthly versus quarterly ranibizumab treatment in neovascular age-related macular degeneration: the EXCITE study." <u>Ophthalmology</u> 118 (5): 831-839	1027
Regeneron SEC Form 8-K Exhibit: "Press Release of Regeneron Pharmaceuticals, Inc. dated May 1, 2006" (May 2, 2006)	1028

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Exhibit Description	<u>No.</u>
FDA, Lucentis, Initial US Approval: 2006.	1029
Regeneron SEC Form 10-Q (May 4, 2007)	1030
Regeneron SEC Form 8-K Exhibit: "Overheads for presentation at Regeneron's Annual Meeting of Shareholders to be held on June 8, 2007" (June 8, 2007)	1031
Regeneron SEC Form 8-K Exhibit: "Press Release dated October 1, 2007" (October 1, 2007).)	1032
Regeneron "Press Release dated April 28, 2008	1033
CMS, Local Coverage Determination (LCD) for Ranibizumab (Lucentis) (L29266, First Coast Service Options, Inc June 14, 2011	1034
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Retina Coding Q & A, Retinal Physician, 16: 18, 54. July/August 2019	1036
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Exhibit Description 102(10): 1425-1433	<u>No.</u>
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Brown, D. M., P. A. Campochiaro, R. B. Bhisitkul, A. C. Ho, S. Gray, N. Saroj, A. P. Adamis, R. G. Rubio and W. Y. Murahashi (2011). "Sustained benefits from ranibizumab for macular edema following branch retinal vein occlusion: 12-month outcomes of a phase III study." <u>Ophthalmology</u> 118 (8): 1594-1602.	1042
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Exhibit Description	<u>No.</u>
U.S. Provisional Application 61/432,245	1045
U.S. Provisional Application 61/434,836	1046
U.S. Provisional Application 61/591,657	1047
Dixon <i>et al.</i> , "VEGF Trap-Eye for the treatment of neovascular age- related macular degeneration," (2009) 18(10):1573-1580	1048

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I. INTRODUCTION

1. I have been retained by counsel for Chengdu Kanghong Biotechnology Co., Ltd. ("Kanghong") as a technical expert in connection with the proceeding identified above. I have been asked to provide my opinions and views on the materials I have reviewed in relation to U.S. Patent No. 10,828,345 (the "345 patent") (Ex. 1001) and the scientific and technical knowledge regarding the subject matter of the '345 patent before and at the earliest possible priority date. I submit this declaration in support of Kanghong's petition for post grant review of the '345 patent against Regeneron Pharmaceuticals, Inc. ("Patent Owner" or "Regeneron").

2. I am being paid at an hourly rate for my work on this matter. I have no personal or financial stake or interest in the outcome of the present proceeding.

II. PROFESSIONAL BACKGROUND

3. I am the Joan Whitten Miller Scholar in Retina and an Assistant Professor of Ophthalmology at Harvard Medical School. In 2005, I completed my MD/PhD in the selective Inteflex program at the University of Michigan Medical School, followed by a residency (2006-09) in ophthalmology and medical retina/research fellowship (2009-10) at the University of Michigan Kellogg Eye

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Center. I then pursued my vitreoretinal surgery fellowship at the Doheny Eye Institute / University of Southern California USC+LAC program. Afterward, I joined the faculty of the Massachusetts Eye and Ear Infirmary and Harvard Medical School (MEEI/HMS), serving until 2019 when I was recruited to the Wilmer Eye Institute at Johns Hopkins School of Medicine to serve in a similar capacity. In 2020, I was recruited back to MEEI/HMS as the Joan Whitten Miller Scholar in Retina.

4. I maintain an active medical and surgical retinal practice at the two major MEEI offices, in downtown Boston and in the Longwood Medical Area next to Harvard Medical School. I treat a variety of disorders including retinal detachments, macular holes, epiretinal membranes, age-related macular degeneration (AMD), retinal arterial and venous occlusions, and diabetic retinopathy. In my clinical practice, I frequently treat patients with VEGF antagonists, including Regeneron's aflibercept (Eylea), Genentech's ranibizumab (Lucentis), as well as off-label use of Genentech's bevacizumab (Avastin). Surgically, I have a particular interest in the repair of complex retinal detachments including those secondary to proliferative vitreoretinopathy and diabetic related traction. I have been named to Boston Magazine's honor roll of Top Doctors four times.

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5. In addition to my practice, I run a research lab. My lab uses advanced techniques such as RNA-seq and AAV gene therapy in order to study the molecular mechanisms of retinal disease and develop new therapies. I most recently showed that selective overexpression of Nrf2 in the RPE of a mouse model of photoreceptor degeneration protected the RPE and preserved visual function. I am a co-inventor of the AAV vector to overexpress Nrf2 in the RPE, which is under consideration for clinical trials. I recently received a Thome Foundation grant to study the role of retinal metabolism and how it may factor in the development of early age-related macular degeneration. My research has been funded by the NIH/NEI, and I was the inaugural recipient of the MEEI Iraty Award for retinal diseases in 2017. To date, I have received over \$1.25 million in funding for my research, with an additional \$0.5 million pending.

6. I also teach ophthalmology. I have won several teaching awards throughout my career, and was named the Division Educational Champion for Retina resident education while on the faculty at the Johns Hopkins Wilmer Eye Institute. I am a faculty member of the MEEI/HMS vitreoretinal surgery fellowship program, and teach clinical management of retina problems and advanced vitreoretinal surgical techniques. My teaching includes local instruction on the management of medical and surgical retina conditions, including the

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evaluation and management of medical retina disorders commonly treated by anti-VEGF agents such as wet AMD, diabetic retinopathy, BRVO, and CRVO. I have also lectured at regional and national continuing medical education events (CME) for Harvard Medical School and Johns Hopkins University. I have trained over 36 surgical and medical retina fellows who have gone on to careers in academic ophthalmology or private practice, as well as mentored several medical students and residents.

7. I am a member of the Association for Research in Vision and Ophthalmology, the American Academy of Ophthalmology, and the American Society of Retina Specialists; I am a diplomat of the American Board of Ophthalmology. I am a licensed physician in the states of Maryland, Massachusetts, and Michigan. I have served as an editor for the Journal of Visualized Experiments, JAMA Ophthalmology, American Journal of Ophthalmology, Ophthalmology, PLOS One, Investigative Ophthalmology & Visual Science, and Translational Vision Science and Technology. My research has been widely published; a full list of my publications can be found in my curriculum vitae (Ex. 1015).

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III. BASIS FOR OPINION

8. My opinions and views set forth in this report are based on my education, training, and experience in ophthalmology, the materials I reviewed in preparing this report, and the scientific knowledge regarding the same subject matter that existed prior to the earliest filing of a patent application in the '345 patent family.

9. I have considered information from various sources in forming my opinions. Besides drawing on my experience as a clinician and researcher, I have reviewed the following materials: (a) the '345 patent (Ex. 1001), (b) the prosecution history of the '345 patent (Ex. 1002), (c) all prior art references cited herein including all prior art relied upon in my analysis of each challenged claim set forth below (including Ex. 1004-05), (d) all other documents and references cited herein (including Ex. 1006-47), and (e) the petition for post grant review of the '345 patent to which my declaration relates.

IV. SUMMARY OF MY OPINIONS

The '345 patent covers a dosing regimen of a VEGF trap
 comprising an initial dose, secondary doses every 4 weeks, and tertiary doses every
 weeks. The '345 patent is anticipated by at least Shams, a Genentech patent
 application. Shams teaches the Regeneron's VEGF-Trap and the same dosing

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regimen in claim 1 of the '345 patent, 4 week secondary dosing followed by 12 week tertiary dosing. Further, Shams explicitly teaches the additional limitations of dependent claims 2-11.

11. Claim 1 of the '345 patent is also obvious over Regeneron's own press releases published more than one year before the filing of the earliest patent application in the '345 patent family. In a 2009 press release about its clinical trials, Regeneron disclosed a dosing program that included 4-week secondary doses followed by 8 week tertiary doses. One of skill in the art would have found claim 1's dosing regimen obvious over the press release's dosing regimen, especially in view of Shams' 12-week tertiary doses. One or both of the 2009 Press Release and Shams teaches each limitation of the dependent claims of the '345 patent.

12. The '345 patent's claim 8 is not supported by a pre-2013 application. Claim 8 lists a number of diseases treatable by the 4 week plus 12 week dosing regimen, including Branch Retinal Vein Occlusion ("BRVO"). BRVO was not included in a pre-2013 application in the '345 patent family; the disorder was added in a July 2013 patent filing.

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V. PATENT LAW STANDARDS

13. I am not an attorney. I have been informed about certain aspects of the law that are relevant to my opinions. My analysis and opinions are based on my expertise in this technical field, as well as the instructions for the legal standards relating to validity provided by counsel. My understanding of the law is as follows.

A. Claim Construction

14. I understand that before any invalidity analysis can be properly performed, the scope and meaning of the challenged claims must be determined by claim construction.

15. I understand that a patent may include two types of claims, independent claims and dependent claims. I understand that an independent claim stands alone and includes only the limitations it recites. I understand that a dependent claim depends from an independent claim or another dependent claim. I understand that a dependent claim includes all the limitations that it recites in addition to the limitations recited in the claim (or claims) from which it depends.

16. I understand that to determine how a person of ordinary skill would have understood a claim term, one should look to sources available at the time of the invention that show what a person of skill in the art would have

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understood disputed claim language to mean. It is my understanding that this may include what is called "intrinsic" evidence as well as "extrinsic" evidence.

17. I understand that, in construing a claim term, one should primarily rely on intrinsic patent evidence, which includes the words of the claims themselves, the remainder of the patent specification, and the prosecution history. I understand that extrinsic evidence, which is evidence external to the patent and the prosecution history, may also be useful in interpreting patent claims when the intrinsic evidence itself is insufficient. I understand that extrinsic evidence may include dictionaries and other resources available to those of skill in the art at the time of the invention.

18. I understand that words or terms should be given their ordinary and accepted meaning unless it appears that the inventors were using them to mean something else or something more specific. I understand that to determine whether a term has special meaning, the claims, the patent specification, and the prosecution history are particularly important, and may show that the inventor gave a term a particular definition or intentionally disclaimed, disavowed, or surrendered claim scope.

19. I understand that the claims of a patent define the scope of the rights conferred by the patent. I understand that because the claims point out and

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distinctly claim the subject matter, which the inventors regard as their invention, and claim construction analysis must begin with and is focused on the claim language itself. I understand that the context of the term within the claim as well as other claims of the patent can inform the meaning of a claim term. For example, because claim terms are normally used consistently throughout the patent, how a term is used in one claim can often inform the meaning of the same term in other claims. Differences among claims or claim terms can also be a useful guide in understanding the meaning of particular claim terms.

20. I understand that a claim term should be construed not only in the context of the particular claim in which the disputed term appears, but also in the context of the entire patent, including the entire specification. I understand that because the specification is a primary basis for construing the claims, a correct construction must align with the specification.

21. I understand that the prosecution history of the patent as well as art incorporated by reference or otherwise cited during the prosecution history are also highly relevant in construing claim terms. For instance, art cited by or incorporated by reference may indicate how the inventor and others of skill in the art at the time of the invention understood certain terms and concepts.

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Additionally, the prosecution history may show that the inventors disclaimed or disavowed claim scope, or further explained the meaning of a claim term.

22. With regard to extrinsic evidence, I understand that all evidence external to the patent and prosecution history, including expert and inventor testimony, dictionaries, and learned treatises, can also be considered. For example, technical dictionaries may indicate how one of skill in the art used or understood the claim terms. However, I understand that extrinsic evidence is considered less reliable than intrinsic evidence, and for that reason is generally given less weight than intrinsic evidence.

23. I understand that in general, a term or phrase found in the introductory words or preamble of the claim, should be construed as a limitation if it recites essential structure or steps, or is necessary to give meaning to the claim. For instance, I understand preamble language may limit claim scope: (i) if dependence on a preamble phrase for antecedent basis indicates a reliance on both the preamble and claim body to define the claimed invention; (ii) if reference to the preamble is necessary to understand limitations or terms in the claim body; or (iii) if the preamble recites additional structure or steps that the specification identifies as important.

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24. On the other hand, I understand that a preamble term or phrase is not limiting where a challenged claim defines a structurally complete invention in the claim body and uses the preamble only to state a purpose or intended use for the invention. I understand that to make this determination, one should review the entire patent to gain an understanding of what the inventors claim they invented and intended to encompass in the claims.

B. Anticipation

25. I understand that a challenged claim can be invalid for lacking novelty over the prior art, and that this concept is also known as "anticipation." I understand that a prior art reference anticipates a challenged claim, and thus renders it invalid by anticipation, if all elements of the challenged claim are disclosed in the prior art reference. I understand that the prior art reference does not have to use the same words as the challenged claim, but all of the requirements of the claim must be disclosed so that a person of ordinary skill in the art could make and use the claimed subject-matter.

26. I understand the disclosure in the prior art reference can be either explicit or inherent. I understand that a disclosure is inherent if it is necessarily present. I understand that inherency may not be established by possibilities or probabilities, and the mere fact that a certain thing may result is not sufficient. I

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also understand that an inherent disclosure need not be recognized by those skilled in the art at the time of invention.

27. I understand that when a challenged claim covers several structures, either generically or as alternatives, the claim is deemed anticipated if any of the structures within the scope of the claim is found in the prior art reference.

28. I understand that when a challenged claim requires selection of an element from a list of alternatives, the prior art teaches the element if one of the alternatives is taught by the prior art.

29. I understand that a claimed range is anticipated by a prior art reference if the reference discloses a point within the range.

C. Obviousness

30. I understand that a claim is invalid if the differences between the claimed subject matter and the prior art are such that the claimed subject matter would have been obvious to a person of ordinary skill in the pertinent art at the time of the alleged invention.

31. I understand that obviousness must be determined with respect to the challenged claim as a whole.

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32. I understand that one cannot rely on hindsight in deciding whether a claim is obvious.

33. I also understand that an obviousness analysis includes the consideration of factors such as (1) the scope and content of the prior art, (2) the differences between the prior art and the challenged claim, (3) the level of ordinary skill in the pertinent art, and (4) "secondary" or "objective" evidence of non-obviousness.

34. Secondary or objective evidence of non-obviousness includes evidence of: (1) a long felt but unmet need in the prior art that was satisfied by the claimed invention; (2) commercial success or the lack of commercial success of the claimed invention; (3) unexpected results achieved by the claimed invention; (4) praise of the claimed invention by others skilled in the art; (5) taking of licenses under the patent by others; (6) deliberate copying of the claimed invention; and (7) contemporaneous and independent invention by others. However, I understand that there must be a relationship between any secondary evidence of nonobviousness and the claimed invention.

35. I understand that a challenged claim can be invalid for obviousness over a combination of prior art references if a reason existed (at the time of the alleged invention) that would have prompted a person of ordinary skill

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in the art to combine elements of the prior art in the manner required by the challenged claim. I understand that this requirement is also referred to as a "motivation to combine," "suggestion to combine," or "reason to combine," and that there are several rationales that meet this requirement.

36. I understand that the prior art references themselves may provide a motivation to combine, but other times simple common sense can link two or more prior art references. I further understand that obviousness analysis recognizes that market demand, rather than scientific literature, often drives innovation, and that a motivation to combine references may come from market forces.

37. I understand obviousness to include, for instance, scenarios where known techniques are simply applied to other devices, systems, or processes to improve them in an expected or known way. I also understand that practical and common-sense considerations should be applied in a proper obviousness analysis. For instance, familiar items may have obvious uses beyond their primary purposes.

38. I understand that the combination of familiar elements according to known methods is obvious when it yields predictable results. For instance, obviousness bars patentability of a predictable variation of a technique even if the technique originated in another field of endeavor. This is because design

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incentives and other market forces can prompt variations of it, and predictable variations are not the product of innovation, but rather ordinary skill and common sense.

39. I understand that a particular combination may be obvious if it was obvious to try the combination. For example, when there is a design need or market pressure to solve a problem and there are a finite number of identified, predictable solutions, a person of ordinary skill has good reason to pursue the known options within his or her technical grasp. This would result in something obvious because the result is the product not of innovation but of ordinary skill and common sense. However, I understand that it may not be obvious to try a combination when it involves unpredictable technologies.

40. It is further my understanding that a proper obviousness analysis focuses on what was known or obvious to a person of ordinary skill in the art, not just the patentee. Accordingly, I understand that any need or problem known in the field of endeavor at the time of invention and addressed by the patent can provide a reason for combining the elements in the manner claimed.

41. Exemplary rationales that can support a conclusion of obviousness include:

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- Combining prior art elements according to known methods to yield predictable results;
- Simple substitution of one known element for another to obtain predictable results;
- Use of known technique to improve similar devices (methods, or products) in the same way;
- Applying a known technique to a known device (method, or product) ready for improvement to yield predictable results;
- Choosing from a finite number of identified, predictable solutions, with a reasonable expectation of success;
- Known work in one field of endeavor may prompt variations of it for use in either the same field or a different one based on design incentives or other market forces if the variations are predictable to a person of ordinary skill in the art; and
- Some teaching, suggestion, or motivation in the prior art that would have led one of ordinary skill to modify the prior art reference or to combine prior art reference teachings to arrive at the claimed invention.

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42. A person of ordinary skill in the art looking to overcome a problem will often use the teachings of multiple publications together like pieces of a puzzle, even though the prior art does not necessarily fit perfectly together. Therefore, I understand that references for obviousness need not fit perfectly together like puzzle pieces. Instead, I understand that obviousness analysis takes into account inferences, creative steps, common sense, and practical logic and applications that a person of ordinary skill in the art would employ under the circumstances.

43. I understand that a claim can be obvious in light of a single reference, if the elements of the challenged claim that are not explicitly or inherently disclosed in the reference can be supplied by the common sense of one of skill in the art.

44. I understand that when the general conditions of a claim are disclosed, it is presumptively obvious to discover the optimum or workable ranges by routine experimentation. I understand that if the prior art recognizes that a variable affects a relevant property or result, then the discovery of an optimum value of the variable is obvious.

45. I understand that obviousness also bars the patentability of applying known or obvious design choices to the prior art. One cannot patent

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merely substituting one prior art element for another if the substitution can be made with predictable results. Likewise, combining prior art techniques that are interoperable with respect to one another is generally obvious and not patentable.

46. In order for a claim to be found invalid based upon a modification or combination of the prior art, there must be reasonable expectation that a person of ordinary skill would have successfully modified or combined the prior art to arrive at the claimed arrangement. This does not mean that it must be certain that a person of ordinary skill would have been successful – the law only requires that the person of ordinary skill in the art would have perceived a reasonable expectation of success in modifying or combining the prior art to arrive at the claimed invention.

47. In sum, my understanding is that obviousness invalidates claims that merely recite combinations of, or obvious variations of, prior art teachings using understanding and knowledge of one of skill in the art at the time and motivated by the general problem facing the inventor at the time. Under this analysis, the prior art references themselves, or any need or problem known in the field of endeavor at the time of the invention, can provide a reason for combining the elements of or attempting obvious variations on prior art references in the claimed manner.

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D. Written Description

48. I am informed and understand that the written description must actually or inherently disclose each and every element of a claim in order to meet the written description requirement. I am also informed that, to meet this requirement, a patent application must reasonably convey to those skilled in the art that the inventor had possession of the full scope of the invention. I am further informed that the words of the claim need not appear *in haec verba* in the written description but it is insufficient that undisclosed subject matter would have been obvious to a POSITA at the time the patent application was filed.

VI. BACKGROUND TECHNOLOGY

49. Many sight-threatening disorders of the eye are due to the dysfunction of blood vessels. Blood vessels perform the critical role of supplying oxygen and nutrients to all of our tissues, and so when their function becomes disrupted by disease, the consequences can be severe. This is particularly true in the eye, where the delicate nature, transparency, and compactness demanded of the visual apparatus to perform its function renders it extremely vulnerable to vascular dysfunction. One simplistic, but broad perspective is to classify clinically significant vascular dysfunction into two categories. The first category is neovascularization, in which new vessels grow into a part of a tissue where they $\frac{26}{1000}$

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should not be, interfering with the function of and/or destroying the tissue into which it grows. The second category is loss of vascular integrity, in which the vessel walls become more leaky. As a result of vascular generation, there can be an escape of hematological components ranging from red blood cells to plasma components outside of the vascular lumens and into tissue that should remain "dry." This can be disruptive in the same way that it is destructive when water leaks into what should be a dry basement. Loss of integrity can also mean the insufficient transport of oxygen and nutrients, which in itself is detrimental to the health of the local tissue. In practice, these pathophysiological processes may overlap as vascular insufficiency can lead to neovascularization, and vessels borne of a neovascular process have less barrier integrity compared to healthy vessels.

50. Because pathophysiology of the vasculature is central to many eye conditions, it is an area that has been under intense study for many years. Through this work, it has become clear that the molecule, Vascular Endothelial Growth Factor (VEGF), plays a central role in this disease process as the key molecule mediating neovascularization and increase in vascular permeability. It is this recognition of the central role of VEGF as a promoter of angiogenesis (and cause of ocular disorders such as neovascular age-related macular degeneration) and the subsequent development of VEGF antagonists for clinical use that has

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revolutionized the treatment of many eye conditions, allowing the restoration of sight in many patients who two decades ago would have been resigned to blindness.

51. The story of how anti-VEGF agents came into clinical use for the retina is fascinating, not just from a medical and scientific point of view, but also from a socioeconomic perspective. I also believe it to be relevant in understanding the evolution of the dosing regimen of these agents that most retina specialists now utilize. It is instructive to examine these revolutionary events through the perspective of wet age-related macular degeneration, the first ocular disease against which anti-VEGF agents were employed.

52. If one thinks of the eye as a camera, the retina is the film of the camera; and just as the film in a traditional camera, it resides along the back wall and is where the lenses focus the light that enters the eye/camera. The retina is a transparent, multi-laminar structure of cells, including the photoreceptor cells that convert light to biochemical and electrical signals that we perceive as vision. The central part of the retina, called the macula, has a particularly high concentration of cone photoreceptors - although it is small in surface area relative to the rest of the retina, it is critical because this high concentration of cones gives us the high

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resolution vision that we take for granted as "sight" (such as the ability to read this document).

53. In many patients, the macula begins to degenerate with age (as one reaches the age of 50 and beyond), in a disease known as age-related macular degeneration (AMD). Approximately 11 million people in the US alone have some form of AMD, with the number projected to double by 2050 as modern medicine increases life expectancy. The degenerative changes typically begin with deposits of material forming under the retina in the absence of bleeding, which is a subtype of AMD known as "dry." Vision loss with dry macular degeneration can be slow, and in mild cases, of little clinical significance. However, in about 10% of these patients, the "dry" degenerative changes are supplanted by growth and dysfunction of blood vessels in the choroid. The choroid is a dense network of blood vessels residing just beneath the retina, whose function is to transport oxygen and nutrients to the photoreceptors. In some AMD patients, the choroidal vessels behind the macula undergo a neovascularization process, proliferating into a choroidal neovascular membrane (CNV) that subsequently leaks and bleeds. At this point, the patient is termed as converting to "wet" macular degeneration. This event can lead to abrupt vision loss, initially from the leakage of blood and fluid,

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but then ultimately death of the photoreceptors, eventual central scarring and central blindness, and loss of one's high resolution vision.

54. As recently as the early 2000's, wet macular degeneration meant the severe and irreversible loss of functional vision. The only treatment that had shown any benefit up to that point was thermal laser, in which an ophthalmologist would apply a laser directly into the macula to target the CNV. While this would destroy the CNV, it would also cause scarring and an immediate central blind spot (scotoma) that was very noticeable to the patient. However, the idea was that the patients would eventually be worse off if they did not undergo this sight-robbing treatment – i.e., the treatment would prevent the later development of an even larger scar and scotoma had the CNV been allowed to continue to flourish on its own. This is well-described in the reports of the Macular Photocoagulation Study (MPS) which demonstrated the efficacy of this now outdated technique. The authors advised that "Both the ophthalmologist and the patient selected for treatment of subfoveal CNV should be prepared for an immediate decrease in visual acuity, 3 lines on average, with relatively stable visual acuity thereafter. No substantial treatment benefit was observed until 18 months after treatment, on average." (Ex. 1017.)

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55. The next advance, photodynamic therapy (PDT), relied on the systemic infusion of a photosensitizing dye that preferentially accumulated in the CNV and whose toxic effects were then "activated" by application of a specific wavelength of light to the CNV itself in an attempt to limit the scope of the dye's toxicity. Like thermal laser, this treatment could still only slow the rate of vision loss in wet AMD patients. (Ex. 1018.) It is also worth mentioning that as limited as these treatments were, they worked best in only a subset of wet macular degeneration patients - those with a variant known as classic disease. Thus, for many years, the state of the art treatment for a wet AMD patient was to undergo examination and fluorescein angiography by a retina specialist. Those identified to have primarily classic disease were offered a treatment that might make things worse initially and would only slow their eventual vision loss; those without classic disease had even less hope. Needless to say, both retina specialists and patients were very eager to have a more effective treatment for this condition.

56. The anti-VEGF revolution for treatment of eye disease began in 2004. Thirty years after Dr. Judah Folkman at Harvard Medical School proposed the idea of specifically targeting an angiogenic factor to treat disease, Gragoudas, Adamis and colleagues published in the *New England Journal of Medicine* that intravitreal injections of Pegaptanib (Macugen), a ribonucleic acid aptamer that

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selectively blocked one specific isoform of VEGF-A (VEGF165), could prevent vision loss. (Ex. 1019.) It was necessary to repeat the injection about every 6 weeks, and like PDT it was only able to slow the loss of vision. However, unlike PDT, it appeared to be effective regardless of the angiographic subtype of CNV.

57. Pegaptanib represents an important medical milestone in the application of anti-VEGF agents, but was quickly relegated to a historical footnote when the results of the phase 3 trial of ranibizumab were reported at the 2005 American Society of Retina Specialists (ASRS) meeting in Montreal, Canada. The results showed that Genentech's anti-VEGF agent, ranibizumab (Lucentis), administered by monthly intravitreal injections, actually *improved* the vision of patients with wet macular degeneration. Like pegaptanib, the treatment appeared effective against all subtypes of wet AMD rather than just primarily "classic." This marked the first time in history that retina specialists could tell their wet AMD patients that a treatment was available to actually make their vision better. Unlike pegaptanib, which selectively blocked one isoform of VEGF that was thought to be most important in eye disease, ranibizumab blocked all isoforms of VEGF-A. Studies soon showed that pegaptanib's theoretical safety advantage of blocking a specific isoform never materialized, and pan-VEGF-A blockade proved more effective, opening the door wider for anti-VEGF agents that block multiple

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VEGF isoforms as well as other angiogenic molecules, including Regeneron's aflibercept (Eylea).

58. Not only was the 2005 ASRS meeting important for the presentation of the first clinical trial that improved vision in wet AMD patients, but it would also be remembered for the presentation of data regarding the off-label use of another anti-VEGF agent, bevacizumab (Avastin). Ironically also made by Genentech, bevacizumab had the same VEGF blocking domains as ranibizumab, but was a full length antibody. Bevacizumab had already been FDA-approved for intravenous administration to treat colorectal cancer as of 2004. Lucentis was still undergoing clinical trials at the time, but retina specialists were desperate for a more effective treatment. Phil Rosenfeld and colleagues at Bascom Palmer Eye Institute had recognized the similarities between bevacizumab and ranibizumab and had already begun to administer it to patients in an off-label manner. First, they systemically administered intravenous infusions of bevacizumab, showing wet AMD patients who received 2 to 3 intravenous infusions of bevacizumab over 6 months had equivalent results to those receiving monthly injections of ranibizumab. (Ex. 1020.) Secondly, they showed that an intravitreal injection of bevacizumab also resolved exudation from macular degeneration by OCT as seen

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with ranibizumab, proof of concept that bevacizumab could be delivered in the same manner and with the same result as Ranibizumab. (Ex. 1019.)

59. The simultaneous reports created a unique situation. As previously described, ranibizumab, an anti-VEGF agent specifically developed for the eye, had demonstrated a paradigm shifting result (for the first time a treatment that improved the vision of wet AMD patients) in a landmark phase III clinical trial. Typically, one would have expected ranibizumab and the clinical trial's monthly dosing regimen to set the standard for treatment once FDA-approval was obtained. Yet, the success reported with the off-label use of bevacizumab, albeit in studies of much more limited scope, fundamentally changed the situation. Bevacizumab closely resembled ranibizumab on a molecular basis, was already available off-label, was manufactured by the same company, and was now reported to be effective when compounded for intravitreal injections. For so long there was no treatment for wet AMD, and now the promise of anti-VEGF to improve wet AMD patients was obvious with the ranibizumab results and available "today" if one was willing to use off-label bevacizumab. Thus, even before ranibizumab was FDA-approved for wet AMD in June 2006, retina specialists were already using bevacizumab off-label and improvising their own regimens based on their best clinical judgment.

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60. One of the first reports on bevacizumab was presented at the AAO 2005 meeting in October 2005 and later published in the journal *Ophthalmology*. (Ex. 1022.) Although the anti-VEGF agent ranibizumab had been dosed at a 4 week interval for the clinical trials, the authors noted that for bevacizumab "The optimum dosing sequence for intravitreal bevacizumab is undetermined. We elected to defer reinjection into eyes when there was complete resolution of SRF, macular edema, and PEDs until there was a recurrence. Some patients have not recurred 15 weeks after a single injection." Not only did this report bolster the initial reports that bevacizumab was effective for wet AMD, but it also provided evidence that not all patients required monthly dosing and the rationale that dosing of anti-VEGF may be individualized.

61. This story illustrates that from the very beginning of the anti-VEGF revolution in the eye, retina specialists of ordinary skill and the art were already thinking of ways to test the durability of the treatment and extend the time between treatments. From the perspective of the retina specialist, as exciting as it was to suddenly have one, and maybe two treatments for a previously untreatable condition, the ground was shifting rapidly. Suddenly, patient outlook for wet AMD changed from having a blinding disease with poor treatment options to having a disease where vision could be restored with an injection. There was the

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logistical challenge of accommodating a sudden change in practice pattern, in which a patient that previously would only need to be seen every 6-12 months for something that had little chance of improvement was now returning regularly, and potentially every month. There was concern our clinics would be overwhelmed with "injection patients." In addition, many of these patients were elderly and had difficulty getting to our offices in the first place - to ask them to come in every month would be stressful for them. Furthermore, there was great concern that these new treatments, while groundbreaking in terms of saving vision, could bankrupt the healthcare system. The wholesale cost of ranibizumab was \$1950 per treatment, and one did not need to be an economist to understand the ramifications of potentially adding \$1950 to the monthly health care bill of a million patients. All of these factors, as well as the early hints that bevacizumab (which cost about \$100 per treatment) might have a similar effect over longer time periods, made it second nature to consider decreasing the frequency of anti-VEGF dosing in "real life." Consideration of these issues was not limited to retina specialists, as with the billions of dollars at stake, the esoterics of AMD treatment spilled over into the public eye (for example see Wall Street Journal "Genentech's Big Drug for Eyes Faces a Rival," (Ex. 1023)), and ultimately sparked the National Institutes of Health to embark on the Comparison of Age-related Macular Degeneration

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Treatment Trials (CATT), comparing ranibizumab vs bevacizumab and monthly vs as-needed dosing.

62. For these reasons, there were intensive efforts on the part of the retina community to discover the true durability of anti-VEGF agents. Amongst the key studies during this time was the PrONTO study, which described the results of an as-needed (as also known as "pro re nata," or "PRN") regimen and established the central role of the OCT in determining the dosing requirements for a patient on anti-VEGF treatment. (Ex. 1024.) Although PRN dosing reduced the number of injections administered, the frequency of visits remained the same. Many retina specialists began adopting an alternative regimen, deemed treat-andextend (TER). (Ex. 1025.) Essentially, this treatment strategy begins with regular fixed interval dosing, typically at the monthly interval, until the disease is under control. At that point, the intervals between exams and injections are extended, often by 1-2 week intervals. If the disease reactivates, the intervals between visits is then reduced to the previous successful interval with the goal of maintaining a fluid-free retina with the least number of office visits and injections as possible. Multiple surveys of the ASRS membership over the years suggest that TER remains by far the most popular regimen.

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63. Recognizing these factors, the use of ranibizumab at extended intervals was studied. The PIER study explored quarterly (dosing every 12 weeks) administration following a series of three monthly injections. (Ex. 1026.) The study demonstrated quarterly dosing was superior to sham. However, the study populations in PIER were not directly compared to a monthly dosing regimen within the same study. A prospective study (EXCITE) directly compared monthly to quarterly dosing. This study was consistent with the findings in PIER in that both monthly and quarterly dosing of ranibizumab was able to improve vision of wet AMD patients, but that the vision of those dosed monthly improved to a greater extent. (Ex. 1027.)

64. Regeneron entered the VEGF antagonist arena in 2005, beginning within clinical development of VEGF Trap for treatment of AMD with a Phase I, Dose-Escalation, Safety, Tolerability, and Bioactivity Study. The study included "a single dose of VEGF Trap-Eye at doses ranging of 0.05, 0.15, 0.5, 1, 2, and 4 milligrams (mg) intravitreally." (Regeneron SEC Form 8-K: "Press Release of Regeneron Pharmaceuticals, Inc. dated May 1, 2006" (May 2, 2006) at 4-5 (Ex. 1028).) Based on the results, Regeneron initiated a Phase 2 trial in AMD. (*Id.*) The Phase 2 AMD clinical trial of intravitreally administered VEGF Trap (called "CLEAR-IT 2") included two groups receiving monthly doses of 0.5 or 2.0 mg of

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VEGF Trap-Eye and three groups receiving quarterly doses of 0.5, 2.0, or 4.0 mg of VEGF Trap-Eye (at baseline and week 12). (Ex. 1013.) Thus, patients received an initial dose of VEGF Trap followed by either 3 secondary doses administered every 4 weeks until week 12, or one secondary dose administered at week 12. (Id.) In May 2007, Regeneron announced positive results of the trial, including for those patients dosed at 12 weeks only: "Moreover, patients in the dose groups that received only a single dose, on average, compared to baseline, demonstrated a decrease in excess retinal thickness ($p \le 0.0001$) and an increase in visual acuity (p= 0.012) at 12 weeks. There were no drug-related serious adverse events, and treatment with the VEGF Trap-Eye was generally well-tolerated." (Regeneron SEC Form 10-Q (May 4, 2007) (Ex. 1030) at 17-18; see also Regeneron SEC Form 8-K (Ex. 1031): "Overheads for presentation at Regeneron's Annual Meeting of Shareholders to be held on June 8, 2007" (June 8, 2007) at 24-25 (noting that monthly and quarterly dosing did not result in substantially different results at 8 weeks and that the results suggested the opportunity for a longer treatment interval using VEGF Trap-Eye).) An October 2007 Regeneron press release quoted Jeffrey Heier, M.D., a clinical ophthalmologist at Ophthalmic Consultants of Boston, a primary investigator in the Phase 2 study, and chair of the steering committee for the Phase 3 trial: "[the] results reaffirm the decision to study both the 0.5 mg and 2

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mg monthly doses in the Phase 3 program . . . The quarterly dosing arms seemed to sustain their effect on visual acuity out to eight weeks, providing the rationale for exploring an eight-week dosing schedule in the Phase 3 program." (Regeneron SEC Form 8-K (Ex. 1032): "Press Release dated October 1, 2007" (October 1, 2007).) In an April 2008 press release announcing "Encouraging 32-Week Follow-Up Results from a Phase 2 Study of VEGF Trap-Eye in Age-Related Macular Degeneration," Regeneron quoted Quan Dong Nguyen, M.D., M.Sc., Assistant Professor of Ophthalmology, Wilmer Ophthalmological Institute, the Johns Hopkins University School of Medicine, Baltimore, MD and a primary investigator in the Phase 2 study, as stating "it is anticipated that VEGF Trap-Eye may be able to be dosed at a frequency less than once monthly, especially on a chronic basis, without compromising visual acuity." (Regeneron "Press Release dated April 28, 2008" (Ex. 1033).)

65. On September 14, 2009 (the "2009 Press Release," Ex. 1005), Regeneron announced completion of patient enrollment in two randomized, double-masked, Phase 3 clinical trials evaluating VEGF Trap-Eye in the treatment of the neovascular form of age-related macular degeneration. In the trials, VEGF Trap-Eye was evaluated for its effect on maintaining and improving vision when dosed as an intravitreal injection on a schedule of 0.5 milligram (mg) every four

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weeks, 2.0 mg every four weeks, or 2.0 mg every eight weeks (following three monthly doses), as compared with intravitreal ranibizumab administered 0.5 mg every four weeks during the first year of the studies. The 2009 Press Release also described two other trials, one a Phase 3 trial for treatment of central retinal vein occlusion ("CRVO") with six monthly doses of 2 mg VEGF Trap-Eye followed by PRN for another six months and the other a Phase 2 trial for treatment of diabetic macular edema ("DME") which included VEGF Trap-Eye dosed at 0.5 mg or 2.0 mg monthly, 2 mg on an as-needed basis after three monthly loading doses, or 2 mg every eight weeks after three monthly loading doses.

VII. THE '345 PATENT

66. The '345 patent has one independent claim, repeated below:

A method for treating an angiogenic eye disorder in a patient, said method comprising

sequentially administering to the patient

a single initial dose of a VEGF antagonist,

followed by one or more secondary doses of the VEGF antagonist,

followed by one or more tertiary doses of the VEGF antagonist;

wherein each secondary dose is administered 4 weeks after the immediately preceding dose; and

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wherein each tertiary dose is administered 12 weeks after the immediately preceding dose;

wherein the VEGF antagonist is a receptor-based chimeric molecule comprising an immunoglobin-like (Ig) domain 2 of a first VEGF receptor which is Fltl and Ig domain 3 of a second VEGF receptor which is Flk1, and a multimerizing component.

67. Claim 1 has three sequential steps: (1) administer a single dose of a VEGF antagonist, (2) administer "secondary doses" of the VEGF antagonist every four weeks, and (3) administer "tertiary doses" of the VEGF antagonist every 12 weeks. The VEGF antagonist in the claimed dosing regimen is "a receptor-based chimeric molecule comprising an immunoglobin-like (Ig) domain 2 of a first VEGF receptor which is Fltl and Ig domain 3 of a second VEGF receptor which is Flk1, and a multimerizing component."

68. The dependent claims narrow independent claim 1 by specifying the drug administered (claim 2), modes of administration (claims 3 and 4), dose amount (claims 5-7), and the disorder(s) treated (claims 8-11). (Ex. 1001, Col. 22:56-23:13.)

69. Claim 2 requires the VEGF antagonist to be aflibercept.

70. Claim 3 requires all doses are administered intraocularly and claim 4, which depends from claim 3, requires the doses are intravitreal.

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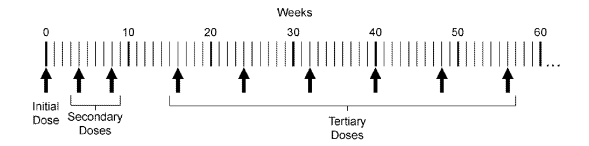
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71. Claim 5 requires all doses are within a range of 0.5 to 2.0 mg, claim 6 specifics 0.5 mg doses, and claim 7 specifies 2.0 mg.

72. Claim 8 requires the regimen treat one of a list of angiogenic eye disorders: age related macular degeneration, diabetic retinopathy, diabetic macular edema, central retinal vein occlusion, branch retinal vein occlusion, and corneal neovascularization. Claims 9-11 require one from the list: age related macular degeneration (claim 9), diabetic retinopathy, (claim 10), and diabetic macular edema (claim 11).

73. The '345 patent generally describes dosing regimens of monthly "secondary doses" followed by longer "tertiary doses." Figure 1 illustrates the dosing regimen with an 8-week tertiary dose.



74. The patent describes seven examples. Example 1 is "a Phase I study [where] 21 subjects with neovascular AMD received a single intravitreal (IVT) dose of VEGFT. Five groups of three subjects each received either 0.05, 0.15, 0.5, 2 or 4 mg of VEGFT, and a sixth group of six subjects received 1 mg." 43

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(Ex. 1001, Column 8, Lines 4-27.) This example corresponds to Regeneron's Phase 1 trials described above.

75. Example 2 describes a Phase 2 clinical trial with "doses (0.5, 2, and 4 mg) of VEGFT tested at 4-week and/or 12-week dosing intervals. There were 5 treatment arms in this study, as follows: 1) 0.5 mg every 4 weeks, 2) 0.5 mg every 12 weeks, 3) 2 mg every 4 weeks, 4) 2 mg every 12 weeks and 5) 4 mg every 12 weeks." (Ex. 1001, Column 8, Lines 29-59.) This example corresponds to Regeneron's Phase 2 AMD trials, publicly announced in May 2007.

76. Example 3 describes a Phase 1 trial studying neovascular AMD, similar to Example 1. The subjects received 4 doses of VEGF-Trap over an eightweek period, with dose levels of 0.3, 1, or 3 mg per kg. (Ex. 1001, Column 8, Line 61 – Column 9, Line 20.) Example 3, like Examples 1 and 2, included no tertiary dosing.

77. Example 4 describes two Phase III clinical trials studying neovascular AMD. Patients were given one of the following dosing regimens: "(1) 2 mg VEGFT administered every 4 weeks (2Q4); (2) 0.5 mg VEGFT administered every 4 weeks (0.5Q4); (3) 2 mg VEGFT administered every 4 weeks to week 8 and then every 8 weeks (with sham injection at the interim 4-week visits when study drug was not administered (2Q8); and (4) 0.5 mg ranibizumab administered

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every 4 weeks (RQ4)." (Ex. 1001, Column 9, Line 22 – Column 14, Line 4.) This example corresponds to the Phase 3 AMD clinical trial described in the 2009 Press Release.

78. Example 5 describes a Phase 2 clinical trial in diabetic macular edema where "[t]wo groups received 0.5 or 2 mg of VEGFT once every four weeks throughout the 12-month dosing period (0.5Q4 and 2Q4, respectively). Two groups received three initial doses of 2 mg VEGFT once every four weeks (i.e., at baseline, and weeks 4 and 8), followed through week 52 by either once every 8 weeks dosing (2Q8) or as needed dosing with very strict repeat dosing criteria (PRN)." (Ex. 1001, Column 14, Lines 6–53.) This example corresponds to the DME trial described in the 2009 Press Release.

79. Example 6 describes a Phase 3 dosing study in CRVO where "patients received 6 monthly injections of . . . 2 mg intravitreal VEG Trap," then received 2 mg as needed. (Ex. 1001, Column 14, Line 55 – Column 15, Line 35.) This example corresponds to the CRVO trial described in the 2009 Press Release.

80. Example 7 lists 20 "examples of dosing regimens within the scope of the present invention." (Ex. 1001, Column 15, Line 36 – Column 17, Line 27.) Although Example 7 discloses tertiary dosing, the dosing frequency is described as either "once every 8 weeks," "less frequent" than the secondary dosing, or PRN.

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None of the 20 exemplary dosing regimens provided in Example 7 include a 12week tertiary dose as required by claim 1 of the '345 patent.

VIII. LEVEL OF SKILL IN THE ART

81. In determining the characteristics of a hypothetical person of ordinary skill in the art of the '345 patent at the time of the claimed invention, I considered several things, including various prior art techniques relating to treatment of angiogenic eye disorders, the type of problems that such techniques gave rise to, and the rapidity with which innovations were made. I also considered the sophistication of the technologies involved, and the educational background and experience of those actively working in the field at the time. I also considered the level of education that would be necessary to understand the '345 patent. Finally, I placed myself back in the relevant period of time and considered the doctors that I have worked with and educated in the field of retinal disease treatment.

82. I came to the conclusion that a person of ordinary skill in the field of art of the '345 patent would have been a person with a medical doctorate, an internship and residency in ophthalmology, and a 1-year medical retina fellowship or 2-year vitreoretinal surgical fellowship. A person with less education but more

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relevant practical experience with retinal disease treatment may also be a person of ordinary skill in the art.

83. In the 2011-12 timeframe, a person of ordinary skill in the art would have known and had the skills necessary to administer intravitreal injections. An ophthalmologist with a 1-year medical retina or 2-year vitreoretinal surgical fellowship would have this experience. Since the introduction of bevacizumab and ranibizumab, general ophthalmologists also receive training in this area as part of their core residency and are familiar with the management of these conditions with intravitreal injections, but typically refer to retina specialists for treatment. Other knowledge and skills in 2011-12 included:

- Ability to examine a retina with dilated fundus examination,
- Ability to interpret fluorescein angiography,
- Ability to interpret optical coherence tomography, and
- Ability to perform intravitreal injections.

84. I would have qualified as a person of at least ordinary skill in the art as of the relevant timeframe. I have a sufficient level of knowledge, experience, and education to provide an expert opinion in the field of the '345 patent.

85. My opinions in this declaration are based on the perspective of a person of ordinary skill in the art as of the relevant timeframe.

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IX. CLAIM CONSTRUCTION

86. To reach my opinions in this case, I have evaluated the claims of the '992 patent using my understanding of the patent law standards described above in Section V. I have given each word its ordinary and accepted meaning in the art. I have reviewed the claims, the patent specification, and the prosecution history and did not find that the inventors gave any claim term a meaning other than that commonly understood by a person of ordinary skill in the art.

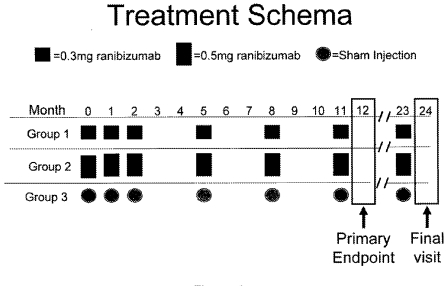
X. ANALYSES OF CHALLENGED CLAIMS OF THE '992 PATENT

A. Shams discloses claims 1-11

87. Shams, published in 2006, was publicly available at least four years before Regeneron filed the earliest patent application in the '345 patent family. (Ex. 1004, Title Page.) Filed by Genentech in 2005, Sham's "Summary of Invention" describes treatment "methods includ[ing] administering to a mammal a number of first individual doses of a VEGF antagonist to the mammal followed by administering to the mammal a number of second individual doses of the VEGF antagonist." (Ex. 1004, Page 4, Line 31 – Page 5, Line 2.) Shams' Figure 2 illustrates an exemplary treatment plan using ranibizumab as the VEGF antagonist (Ex. 1004, Page 6, Lines 8-9):

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88. Shams discloses all the limitations of claim 1. Claim 1 covers a method for treating an angiogenic eye disorder in a patient; Shams discloses a method for treating "intraocular neovascular disease" in a "mammal." (Ex. 1004 at Page 4, Line 31 – Page 5, Line 2; *see also* Page 1, Lines 5-9.) One of skill in the art would understand "intraocular neovascular disease" to include the "angiogenic eye disorders" of claim 1. For example, the '345 patent's claims 8-11 list specific disorders (e.g., age-related macular degeneration) under the "angiogenic eye disorders" umbrella that Shams includes as examples of "intraocular neovascular diseases:"

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An "intraocular neovascular disease" is a disease characterized by ocular neovascularization. Examples of intraocular neovascular diseases include, but are not limited to, e.g., proliferative retinopathies, choroidal neovascularization (CNV), age-related macular degeneration (AMD), diabetic and other ischemia-related retinopathies, diabetic macular edema. myopia, Hippel-Lindau pathological von disease. histoplasmosis of the eye, Central Retinal Vein Occlusion (CRVO), corneal neovascularization, retinal neovascularization, etc.

(Ex. 1004, Page 21, Lines 1-6; *see also* Shams "Background of Invention," Page 1, Lines 12-14 ("Angiogenesis is implicated in the pathogenesis of intraocular neovascular diseases, e.g., proliferative retinopathies, age-related macular degeneration (AMD), etc., as well as a variety of other disorders.").) Although Shams generally refers to treatment of "mammals," Shams includes "a patient" (as used in '345 patent, claim 1) within the treatment plan. For example, Shams describes a "therapeutic dose" as having "a therapeutic effect on the patient." (Ex. 1004, Page 20, Line 33; *see also* Shams Page 23, Lines 30-32 ("Another aspect of the invention is the treatment of an intraocular neovascular disease, e.g., wet form AMD, by administering to a mammal, preferably a human patient").) Thus, although Shams sometimes uses different language than the '345 patent, one of skill in the art would understand Shams to disclose claim 1's "method for treating an angiogenic eye disorder in a patient."

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89. Claim 1 includes sequentially administering doses of a VEGF antagonist to the patient. As shown in Figure 2 above, Shams discloses administering the doses at different points in time, which is the definition of "sequentially administered" given in the '345 patent. (Ex. 1001, Column 3, Lines 32-36 ("[S]equentially administering' means that each dose of VEGF antagonist is administered to the patient at a different point in time, e.g., on different days separated by a predetermined interval (e.g., hours, days, weeks or months).").) Claim 1 requires the VEGF antagonist to be "a receptor-based chimeric molecule comprising an immunoglobin-like (Ig) domain 2 of a first VEGF receptor which is Fltl and Ig domain 3 of a second VEGF receptor which is Flk1, and a multimerizing component." As is commonly known in the art. Regeneron's VEGF-Trap is such a molecule. (See Ex. 1001, col. 2, lines 41-45 ("In one embodiment, the VEGF antagonist comprises one or more VEGF receptor-based chimeric molecule(s), (also referred to herein as a 'VEGF-Trap' or 'VEGFT'). An exemplary VEGF antagonist [is] 'VEGFR1R2-Fc Δ C1(a)' or 'aflibercept.''); see also Ex. 1001, claim 2 (listing "aflibercept" as an example of the VEGF antagonist in claim 1).) Shams explicitly discloses Regeneron's VEGF trap:

> A "VEGF antagonist" refers to a molecule capable of neutralizing, blocking, inhibiting, abrogating, reducing or interfering with VEGF activities including its binding to one or more VEGF receptors. VEGF antagonists include anti-VEGF

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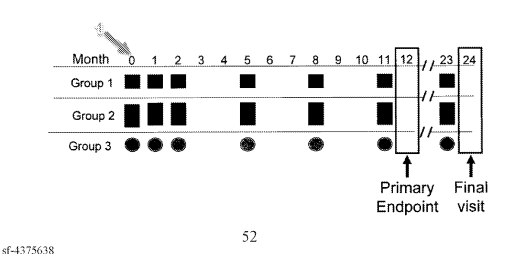
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antibodies and antigen-binding fragments thereof, receptor molecules and derivatives which bind specifically to VEGF thereby sequestering its binding to one or more receptors, anti-VEGF receptor antibodies and VEGF receptor antagonists such as small molecule inhibitors of the VEGFR tyrosine kinases, and fusions proteins, e.g., VEGF-Trap (Regeneron), VEGFi 2rgelonin (Peregrine). VEGF antagonists also include antagonist variants of VEGF, antisense molecules directed to VEGF, RNA aptamers specific to VEGF, and ribozymes against VEGF or VEGF receptors. Antagonists of VEGF act by interfering with the binding of VEGF to a cellular receptor, by incapacitating or killing cells which have been activated by VEGF, or by interfering with vascular endothelial cell activation after VEGF binding to a cellular receptor. All such points of intervention by a VEGF antagonist shall be considered equivalent for purposes of this invention.

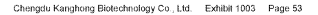
(Ex. 1004, Page 6, Line 27 – Page 7, Line 6 (emphasis added).)

90. Claim 1's treatment plan includes a single initial dose of the

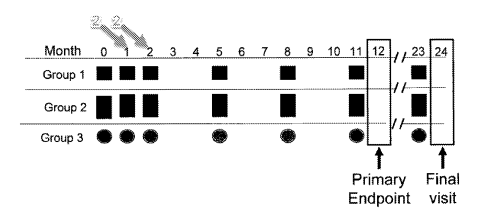
VEGF antagonist. Shams discloses a single initial dose at "day 0" (labelled with



numeral "1" below):



91. Claim 1's treatment plan continues with "secondary doses" administered 4 weeks after the immediately preceding dose. In my opinion, Shams discloses secondary doses every 4 weeks. For example, the second and third doses (labelled with numeral "2" below) in Shams' Figure 2 meet the claimed "secondary doses":



Shams does not use the term "secondary doses" for the second and third doses, instead grouping them with the initial dose as "first individual doses." (*See e.g.* Ex. 1004 at Page 5, Lines 20-21 ("In one embodiment of the invention, the first individual doses are administered at one month intervals (e.g., about 3 individual doses). Typically, there is more than one first individual dose.").) Although the '345 patent uses different language, Shams discloses the same "secondary doses." The '345 patent defines "initial dose" to be "the dose which is administered at the beginning of the treatment regimen" (Ex. 1001, Column 3, Lines 44-46); this is

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Shams' "day 0" dose. The '345 patent defines "secondary doses" to be "the doses which are administered after the initial dose" (Ex. 1001, Column 3, Lines 46-47); in Shams, the doses at month 1 and month 2 follow the single initial dose at day 0. Thus, Shams' doses at month 1 and month 2 are the '345 patent's secondary doses.

92. It is my opinion that one of skill in the art would understand claim 1's "four week" dosing and Shams "one month" dosing to be the same dosing frequency. Typically, surgeons and patients calendar follow-up treatments on a weekly basis (i.e., the same day (and time) of a following week), instead of returning on the same date in a future month. In those cases, returning "monthly" is understood to mean returning in 4 weeks on the same day of the week. By contrast, surgeons and patients avoid reusing the same calendar date for return treatments because patients need to review their weekly schedules for conflicts and calendar date appointments fail when the date falls on a weekend. Additionally, many surgeons have practices in different locations. Typically, the surgeons visit a specific office location on the same day of every week. In such instances, the surgeon and patient may arrange a one-month "follow-up" but imply meeting on the same day in a future week. Further, monthly injections of Lucentis are well known, as pointed out in the '345 patent. This monthly interval is stipulated in part by the FDA-approved dosing guidelines for ranibizumab which state that the

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medicine should be administered between every 28 days and every 3 months. (FDA, Lucentis, Initial US Approval: 2006. US BLA (BL125156) Ranibizumab Injection (Ex. 1029).) With this justification, many Medicare Administrative Contractors do not cover lucentis more frequently than every 28 days per eye (For example, CMS, Local Coverage Determination (LCD) for Ranibizumab (Lucentis) (L29266, First Coast Service Options, Inc June 14, 2011 (Ex. 1034)). More frequent use of Lucentis was deemed a "questionable billing for medical ophthalmology services" in a report by the deputy inspector general for evaluation and inspections for HHS. (Ex. 1035.) Aflibercept is similarly not allowed more frequently than every 28 days based on its FDA approval, and a similar limit is placed on bevacizumab even though it does not have FDA guidelines for ophthalmologic use. (Ex. 1036.) In the rare circumstances that more frequent dosing is needed, a retinal specialist may alternate a more costly medicine such as ranibizumab or aflibercept that will be covered by insurance with bevacizumab, which the patient can afford to pay out of pocket.

93. The '345 patent supports my opinion. The '345 patent uses "four week" dosing and "monthly" dosing interchangeably. For example, the '345 patent describes a dosing frequency of "2 mg monthly (2Q4)." (Ex. 1001, Table 1, Column 13, Lines 29-32.) "Q4" is understood in the art to be shorthand for dosing

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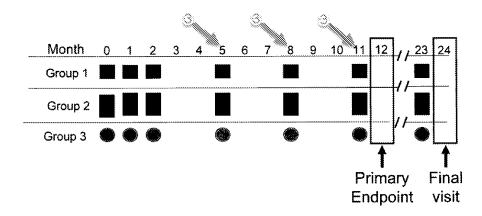
"every 4 weeks" (*see*, *e.g.*, Ex. 1001, Column 9, Lines 60-61.) Thus, the '345 patent treats "monthly" and "Q4" (i.e., every "4 weeks") as equivalent. Other disclosure in the '345 patent describes monthly as including "4 weeks." (*See*, *e.g.*, Col. 14:59-66 (describing the patients in Example 6 as receiving "6 monthly injections . . . once every four weeks"); Col. 15:40-41 (describing a dosing regimen of Example 7 as "once every 4 weeks (monthly)").) Shams too uses "monthly" broadly. For example, Shams describes the prior art methods as "administered in equal monthly (about 28 days) doses of 0.3 mg or 0.5 mg." (Ex. 1004, Page 24, Lines 2-4.)

94. Regeneron's own publications are consistent with my opinion. A 2011 Regeneron publication to Heier equated "4 week" with "monthly" dosing when describing the results of Regeneron's Phase 2 study: "During the 12-week fixed dosing phase, patients in the monthly dosing groups received 0.5 or 2 mg of VEGF Trap-Eye every 4 weeks on day 0 and at weeks 4, 8, and 12 for a total of 4 doses." (*Heier et al*, "The 1-year Results of CLEAR-IT 2, a Phase 2 Study of Vascular Endothelial Growth Factor Trap-Eye Dosed As-needed After 12-week Fixed Dosing," Ophthalmology ,Volume 118, Number 6, June 2011, (Ex. 1013) at 1110, Legend for Figure 2.)

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95. Following the secondary doses, claim 1 requires "tertiary doses" administered every 12 weeks after the immediately preceding dose. Again, Shams discloses the claimed tertiary doses. In Shams, doses at month 5, 8, and 11 follow the "secondary" doses at months 1 and 2.



96. As with the "secondary" doses, Shams does not use the same terminology for claim 1's "tertiary" doses, but again there is no difference. The '345 patent defines "tertiary doses" to be "doses which are administered after the secondary doses." (Ex. 1001, Column 3, Lines 48-49.) Shams month 5, 8, and 11 doses are administered after Shams' "secondary doses" at months 1 and 2.

97. As I described above, one of skill in the art would consider "4 week dosing" and "monthly dosing" to be equivalent. In the same way, one of skill in the art would find "12 weeks" and "three-month dosing" to be equivalent. With 12 week dosing, one of skill in the art might also use the term "quarterly"

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dosing, which is another term used in Shams to describe the "tertiary doses" in the treatment plan. (Ex. 1004, Page 23, Lines 9-11 ("For example, doses may be administered on a monthly schedule followed by subsequent quarterly or more dose schedule.").) All of these terms would be considered equivalent and used synonymously by those skilled in the art. For example, a 2011 Regeneron publication to Heier equated "12-week" with "quarterly" dosing when describing the results of Regeneron's Phase 2 study: "During the 12-week fixed dosing phase, patients . . . in the quarterly dosing groups received 0.5, 2, or 4 mg of VEGF Trap-Eye every 12 weeks on day 0 and at week 12 for a total of 2 doses." (Ex. 1013 at 1110, Legend for Figure 2.)

98. Thus, it is my opinion that Shams discloses all the limitations of claim 1.

99. Shams also discloses the limitations of claim 2. Claim 2 specifies that the VEGF antagonist is aflibercept, another name for Regeneron's VEGF Trap, i.e., the drug referenced in Shams as a VEGF antagonist. (*See, e.g.,* Ex. 1001, col. 2, lines 41-45 ("In one embodiment, the VEGF antagonist comprises one or more VEGF receptor-based chimeric molecule(s), (also referred to herein as a 'VEGF-Trap' or 'VEGFT'). An exemplary VEGF antagonist [is] 'VEGFR1R2-Fc Δ C1(a)' or 'aflibercept.''').) The '345 patent confirms that Regeneron's VEGF

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Trap for eye disorders is called "aflibercept." (*See, e.g.,* Ex. 1001, col. 2, lines 51-54 ("Aflibercept (EYLEA[™], Regeneron Pharmaceuticals, Inc) was approved by the FDA in November 2011, for the treatment of patients with neovascular (wet) age-related macular degeneration.").) As noted, Shams discloses VEGF Trap (Regeneron), and also specifies that the VEGF antagonist is used for treating eye disorders (for example, Shams' Title, "Method for Treating Intraocular Neovascular Diseases"), confirming that Shams is disclosing Regeneron's VEGF Trap treatment for eye disorders.

100. Claim 3 requires "all doses of the VEGF antagonist are administered to the patient by intraocular administration," which is also taught by Shams. Intraocular administration is disclosed by Shams at Page 25, Lines 15-16: "[t]he therapeutic compound for treatment of an intraocular neovascular disease is typically administered by ocular, intraocular, and/or intravitreal injection." (*See also* Ex. 1004, Page 5, Lines 12-13 ("In one embodiment, the administration of the VEGF antagonist is ocular. In one aspect, the administration is intraocular. In another aspect, the administration is intravitreal.").) This disclosure also includes the limitations of the '345 patent's claim 4, i.e. that "the intraocular administration is intravitreal administration".

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101. Shams discloses the '345 patent's claims 5, 6, and 7. Claim 5 requires "all doses of the VEGF antagonist comprise from about 0.5 mg to about 2 mg of the VEGF antagonist" and claim 6 requires a specific dose, 0.5 mg, within claim 5's range. Shams discloses the specific 0.5 mg dose of claim 6 that lies within claims 5's range. (Ex. 1004, Figure 2; Page 31, Lines 58-11.) For claim 7, Shams does not discuss the 2.0 mg dose as a specific dose, but does disclose the range 0.1 mg to 20 mg (Ex. 1004, Page 24, Lines 18-20) which encompasses the 2.0 mg dose of claim 5 and 0.5 mg dose of claim 6.

102. Claim 8 ("wherein the angiogenic eye disorder is selected from the group consisting of: age related macular degeneration, diabetic retinopathy, diabetic macular edema, central retinal vein occlusion, branch retinal vein occlusion, and corneal neovascularization") lists various eye disorders treated by the dosing frequency, which are further specified in claims 9 (age related macular degeneration), claim 10 (diabetic retinopathy), and claim 11 (diabetic macular edema). Shams discloses all of the specific disorders in claims 9-11: "age-related macular degeneration (AMD), diabetic and other ischemia-related retinopathies, diabetic macular edema." (Ex. 1004, Page 21, Lines 1-6.) Thus, Shams discloses each of claims 8-11.

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B. The 2009 Press Release in view of Shams renders claims 1-11 obvious

103. The 2009 Press Release was published on September 14, 2009, more than one year before the filing of the first application in the '345 patent family. As modified by Shams, the 2009 Press Release renders obvious claims 1-11.

i. The 2009 Press Release in view of Shams renders obvious claim 1

a. The 2009 Press Release

104. Regeneron issued the 2009 Press Release to announce its completion of enrollment in clinical trials evaluating the effect of "VEGF Trap-Eye" on wet AMD, CRVO, and DME. (Ex. 1005 at Title ("Enrollment Completed in Regeneron and Bayer Healthcare Phase 3 Studies of VEGF Trap-Eye in Neovascular Age-Related Macular Degeneration (Wet AMD)").) The wet AMD trial included scheduled doses of 0.5 milligram (mg) every four weeks, 2.0 mg every four weeks, or 2.0 mg every eight weeks (with one additional 2.0 mg dose at week four), as compared with intravitreal ranibizumab administered 0.5 mg every four weeks during the first year of the studies. (*Id.* at 1, First and Fourth Paragraphs.) After the first year, patients would continue to be followed and treated for another year on a flexible, criteria-based extended PRN regimen with a

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dose administered at least every 12 weeks, but not more often than every four weeks until the end of the study. (*Id.*) The 2009 Press Release also describes a "development for the treatment of [DME]" where VEGF Trap-Eye is dosed at 0.5 mg or 2.0 mg monthly, 2 mg on an as-needed basis after three monthly loading doses, or 2 mg every eight weeks after three monthly loading doses. (*Id.* at 2, Second Paragraph.)

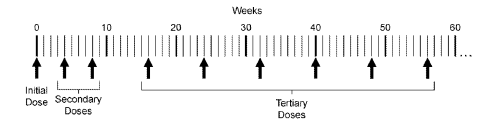
b. The 2009 Press Release and Shams teach the limitations of claim 1

105. The 2009 Press Release teaches a method for treating an angiogenic eye disorder in a patient, as required by claim 1. For example, the trials included treatment of AMD and DME, both listed by the '345 patent as examples of angiogenic eye disorders. (*See, e.g.,* '345 patent dependent claims 8, 9, and 11.) The Press Release also teaches administration of the specific VEGF trap required by claim 1. The 2009 Press Release describes studies related to "VEGF Trap-Eye," which one of skill in the art would understand includes "a receptor-based chimeric molecule comprising an immunoglobin-like (Ig) domain 2 of a first VEGF receptor which is Fltl and Ig domain 3 of a second VEGF receptor which is Flkl, and a multimerizing component." (Ex. 1005 at 1, Last Paragraph ("VEGF Trap-Eye is a fully human, soluble VEGF receptor fusion protein that binds all forms of VEGF-A along with the related placental growth factor (PIGF)."); *Dixon* 62

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et al., "VEGF Trap-Eye for the treatment of neovascular age-related macular degeneration," (2009) 18(10):1573-1580 (Ex. 1048).)

106. Claim 1 requires sequential administration of the VEGF antagonist, beginning with a single initial dose of the VEGF antagonist and followed by 4 week secondary doses. The 2009 Press Release describes two different trials that include 4 weeks doses followed by longer doses: (1) AMD treated with 2.0 mg doses "at an eight-week dosing interval following one additional 2.0 mg dose at week four;" (2) DME treated with "2 mg every eight weeks after three monthly loading doses." This is the same sequential dosing scheme as Figure 1 of the '345 patent.



107. The 2009 Press Release's first of the three monthly doses corresponds to the "initial dose" of the '345 patent's Figure 1 and claim 1. The second and third of the 2009 Press Release's three monthly doses correspond to the "secondary doses" of the '345 patent's Figure 1 and claim 1.

108. The 2009 Press Release treatment plan includes 8-week tertiary doses for both AMD and DME, and thus does not explicitly teach "each tertiary 63 sf-4375638

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dose is administered 12 weeks after the immediately preceding dose." Shams teaches 12 week tertiary doses. For example, Shams Figure 2 (reproduced below) schematically illustrates 4 week secondary doses and 12 week tertiary doses. I described Shams 12-week tertiary dosing above and incorporate that discussion here.

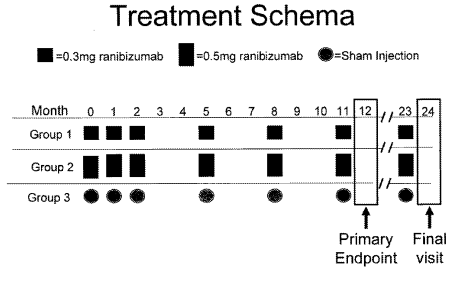


Figure 2

c. One skilled in the art would modify the 2009 Press Release's 4 + 8 treatment plans with Shams' 12 week tertiary dosing

109. By September 2009 (the date of the 2009 Press Release), the

problems associated with monthly dosing VEGF antagonists were well known in

the art. As I described earlier, VEGF antagonists revolutionized eye treatment

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when first introduced. Before then, the treatment options involved preemptively causing partial blindness to avoid total blindness with laser, to later slowing down blindness with PDT or Pegaptanib. With VEGF antagonist treatments, patients now had a treatment choice that could potentially restore their vision.

110.After the initial excitement, those skilled in the art observed drawbacks with anti-VEGF treatments. As discussed earlier in this document, while VEGF antagonists were a significant advancement, the need for serial injections of a VEGF antagonist created many new burdens. One of these was mentioned in the 2009 Press Release, in which anti-VEGF treatments included "monthly office visits and examinations that are inconvenient for these often elderly patients." (Ex. 1005 at 1, Third Paragraph.) In my experience, the "inconvenience" in this article refers to the physical discomfort of receiving an intraocular injection as well as the mobility limitations of many elderly patients and their need to rely on others for help getting to and from the office. Cost is another drawback. In September 2009, a single injection of Lucentis costs almost \$2,000 per month and so monthly injections cost \$24,000 per year; any additional spacing of injections would reduce patient costs and would be very welcomed, given the high price of Lucentis. Again, retinal surgeons were also inconvenienced. With monthly injections, retina specialists' practice could consist

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solely of providing maintenance injections to existing patients, giving few opportunities to expand a practice and treat new patients.

111. Given these market incentives, it was quite common in the art to dose VEGF antagonist at frequencies longer than monthly. This was termed "treat and extend" and, typically, included administering doses every 4 weeks initially, followed by less frequent doses. Contrary to the statements in the '345 patent, prior administration regimens for angiogenic eye disorders did not require monthly administrations throughout the entire course of treatment. (Ex. 1001, Column 2, Lines 26-30.) For example, Shams discloses non-monthly dosing, as I described above. Further, the FDA, in 2006, approved Lucentis for "treatment [of] one injection every three months after the first four injections if monthly injections are not feasible." (Ex. 1006 at 1.) Consistently, Regillo et al. reported, in 2008, that "Ranibizumab administered monthly for three months and then guarterly provided significant VA benefit to patients with AMD-related subfoveal CNV and was well tolerated. The incidence of serious ocular or nonocular adverse events was low." (Regillo et al., "Randomized, Double-Masked, Sham-Controlled Trial of Ranibizumab for Neovascular Age-related Macular Degeneration: PIER Study Year 1," (Ex. 1026) at 1, Left Column, "Conclusions.")

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112. With this backdrop, it was natural for one of skill in the art to consider extending the 2009 Press Release's 4 + 8 dosing regimens. Further, one of skill in the art would have looked to Shams when considering modifications to a VEGF antagonist dosing regimen for at least three reasons. First, Shams covers a dosing regimen for a Genentech VEGF antagonist. Genentech was an early leader in VEGF antagonist research and their research results are important to those skilled the art, especially around 2009-11. In 2009, Genentech offered one of the most popular VEGF antagonists on the market, Lucentis. Anyone considering VEGF antagonist dosing would look to Genentech's research of Lucentis (see, e.g., Ex. 1001, Column 2, Lines 30-31 (citing "prescribing information for Lucentis® [ranibizumab], Genentech, Inc." when describing "prior administration regimens")). Shams reports on Genentech research on Lucentis and, thus, would be relevant to one of skill in the art working on Regeneron's dosing frequency. Second, Regeneron's clinical trials used ranibizumab to determine efficacy of its VEGF-Trap. For example, the 2009 Press Release states that the "primary endpoint of these non-inferiority studies" included comparison with "ranibizumab patients." (Ex. 1005 at 1, First, Fourth, and Fifth Paragraphs.) Given that Regeneron was comparing its drug's efficacy to ranibizumab in the 2009 Press Release, one of skill in the art would naturally look to Genentech's research of

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ranibizumab—such as in Shams—to modify the 2009 Press Release's 4 + 8 dosing regimen. Third, Shams, in listing suitable VEGF antagonists for the 4 + 12 week dosing regimen, specifically identifies Regeneron's VEGF Trap.

113. One of skill in the art would modify the 2009 Press Release's 4 + 8 dosing regimen to 4 + 12 based on the teachings of Shams and on the 2009 Press Release teachings. First, Shams teaches that 4 + 12 dosing is possible. (*See, e.g.*, Ex. 1004, Page 23, Lines 9-11 ("For example, doses may be administered on a monthly schedule followed by subsequent quarterly or more dose schedule.").) Second, the 2009 Press Release teaches that 12-week tertiary dosing should be considered and is a potentially maximum length between tertiary doses. (Ex. 1005, ("During the second year of the study, . . . [tertiary doses] may be given as frequently as every 4 weeks, but no less frequently than every 12 weeks.").) One of skill in the art would naturally consider 12 weeks as a desirable dosing regimen because both Shams and the 2009 Press Release explicitly teach a treatment plan with a 12-week dosing component.

114. One of skill in the art would have had a reasonable expectation of success when modifying the 2009 Press Release with Shams' 12-week tertiary dosing. For instance, Regeneron told its shareholders in 2007 that 12 week dosing works: "[P]atients in the dose groups that received only a single dose, on average,

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compared to baseline, demonstrated a decrease in excess retinal thickness (p < 0.0001) and an increase in visual acuity (p = 0.012) at 12 weeks. There were no drug-related serious adverse events, and treatment with the VEGF Trap-Eye was generally well-tolerated." (Ex. 1031 at 24-25 (presenting analysis of the interim CLEAR-IT results as demonstrating that quarterly dosing (*i.e.*, dosing at week 0 and at week 12), on average, demonstrated an increase in visual acuity and a decrease in excess retinal thickness at both 8 weeks and 12 weeks into the study).) Further, Shams teaches that 12 week tertiary dosing of a VEGF antagonist is successful. As further evidence that 4-week followed by 12-week dosing would be successful, the FDA had previously approved ranibizumab for 4 + 12 dosing. (Ex. 1029; Ex. 1006.)

115. The modification of the 2009 Press Release's 8-week tertiary dose with Shams' 12-week tertiary dose would be routine to those skilled in the art. The modification merely combines prior art elements (Shams' 12-week tertiary dosing) with a known method (the 2009 Press Release's 4-week secondary dosing plus 8-week tertiary dosing) to arrive at a predictable result (a successful treatment of angiogenic eye disorders). Claim 1 is nothing more that the simple substitution of Shams' 12-week tertiary dose for the 2009 Press Release's 8-week tertiary dose.

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ii. The 2009 Press Release in view of Shams renders obvious claims 2-11

116. Claim 2 specifies that the VEGF antagonist is "aflibercept." Aflibercept is another name for Regeneron's VEGF-Trap Eye. (Ex. 1001, Col. 2:38-41 ("An exemplary VEGF antagonist that can be used in the context of the present invention is a multimeric VEGF-binding protein comprising two or more VEGF receptor-based chimeric molecules referred to herein as 'VEGFR1R2-FcllCl(a)' or 'aflibercept.'").) Thus, the 2009 Press Release teaches "the VEGF antagonist is aflibercept."

117. The '345 patent specifies "intraocular administration" and "intravitreal administration" in claims 3 and 4, respectively. The 2009 Press Release teaches intravitreal administration: "VEGF Trap-Eye is being evaluated for its effect . . . when dosed as an intervitreal injection." (Ex. 1005 at 2, First Paragraph.)

118. Claim 5 requires that all doses of the VEGF antagonist are "from about 0.5 mg to about 2 mg" and claim 7 further narrows all doses to 2 mg. In the 2009 Press Release, the 4 + 8 AMD doses are administered at 2 mg and the 4 + 8 DME doses are also administered at 2 mg. (Ex. 1005 at 1, First Paragraph, and 2, Second Paragraph.) Thus, the 2009 Press Release teaches the specific dose of claim 7 and thus also teaches the range of claim 5. As described above, Shams

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teaches the specific 0.5 mg dose of claim 6. Thus, the 2009 Press Release in view of Shams teaches each of claims 5, 6, and 7.

119. The '345 patent lists a number of angiogenic eye disorders in claim 8, including the specific eye disorders of claim 9 (AMD) and claim 11 (DME). The 2009 Press Release teaches treatment of both AMD and DME, teaching the specific eye disorders of claims 9 and 11 and, thus, also teach the elements of the list in claim 8. As described above, Shams teaches the eye disorders of claims 9-11. Thus, the 2009 Press Release in view of Shams teaches each of claims 8, 9, 10, and 11.

C. Claim 8 is not supported by an application filed before July 2013

120. Claim 8, which depends from claim 1, limits the angiogenic eye disorders treatable by the VEGF antagonist. The list includes age related macular degeneration ("AMD"), diabetic retinopathy, diabetic macular edema, central retinal vein occlusion ("CRVO"), branch retinal vein occlusion ("BRVO"), and corneal neovascularization. I understand that, to meet the "written description" requirement, a patent application must reasonably convey to those skilled in the art that the inventor had possession of each of these claimed disorders, including BRVO. It is my opinion that the pre-2013 applications in the '345 patent family do not convey that the inventor had possession of a method of treating BRVO.

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121. When Regeneron filed the first application in 2011 and then the international application in 2012, the patent application listed "examples of angiogenic eye disorders that are treatable using the methods of the present invention," but the list did not include BRVO:

The methods of the present invention can be used to treat any angiogenic eye disorder. The expression "angiogenic eye disorder," as used herein, means any disease of the eye which is caused by or associated with the growth or proliferation of blood vessels or by blood vessel leakage. Non-limiting examples of angiogenic eye disorders that are treatable using the methods of the present invention include choroidal neovascularization, age-related macular degeneration (AMD), diabetic retinopathies, diabetic macular edema (DME), central retinal vein occlusion (CRVO), corneal neovascularization, and retinal neovascularization.

(PCT Application No. PCTUS1220855, (Ex. 1016) Page 5, Paragraph 0025; see

also U.S. Provisional Application 61/432,245, (Ex. 1045) Pages 5-6, Paragraph

0024; U.S. Provisional Application 61/434,836, Pages 5-6, Paragraph 0024 (Ex.

1046); U.S. Provisional Application 61/591,657, Pages 5-6, Paragraph 0024 (Ex.

1047).)

122. In the July 2013 filing, that paragraph was changed to list additional eye disorders "treatable using the methods of the present invention," including, for the first time, BRVO:

The methods of the present invention can be used to treat any angiogenic eye disorder. The expression "angiogenic eye

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disorder," as used herein, means any disease of the eye which is caused by or associated with the growth or proliferation of blood vessels or by blood vessel leakage. Non-limiting examples of angiogenic eye disorders that are treatable using the methods of the present invention include age-related macular degeneration (e.g., wet AMD, exudative AMD, etc.), retinal vein occlusion (RVO), central retinal vein occlusion (CRVO; e.g., macular edema following CRVO), branch retinal vein occlusion (BRVO), diabetic macular edema (DME), choroidal neovascularization (CNV; e.g., myopic CNV), iris neovascularization, neovascular glaucoma, post-surgical fibrosis in glaucoma, proliferative vitreoretinopathy (PVR), optic disc neovascularization, corneal neovascularization, retinal neovascularization, vitreal neovascularization, pannus, pterygium, vascular retinopathy, and diabetic retinopathies.

(U.S. Application No. 13/940,370, (Ex. 1011) Page 5, Paragraph 0026 (emphasis

added); see also Ex. 1001, Column 5:22-39.)

123. The table below compares the eye disorders included in the '345

patent prior to (left column) and after (right column) the July 2013 patent

application. As show in the table below, no disorders were deleted from the

paragraph, but the inventor more than doubled the listed disorders.

Pre-2013 Eye Disorders	July 2013 Eye Disorders
choroidal neovascularization,	choroidal neovascularization,
AMD,	AMD,
diabetic retinopathies,	diabetic retinopathies,
DME,	DME,
CRVO,	CRVO,
corneal neovascularization,	corneal neovascularization,
retinal neovascularization	retinal neovascularization,

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iris neovascularization,
neovascular glaucoma,
post-surgical fibrosis in glaucoma,
proliferative vitreoretinopathy,
optic disc neovascularization,
vitreal neovascularization,
pannus,
pterygium,
vascular retinopathy,
retinal vein occlusion,
BRVO

As can be seen in the table above, BRVO (along with a number of other disorders) was added to the '345 patent family in 2013. Thus, the pre-2013 patent applications did not list BRVO as one of the treatable eye disorders.

124. Nor is treatment of BRVO inherent in any of the disorders listed prior to July 2013. In 2012 (the time of the international filing) one of skill in the art would consider BRVO a different disorder than those listed prior to July 2013. One of skill in the art would not recognize a disclosed treatment of any of the pre-2013 disorders to be possession of a treatment for BRVO. In 2011-12, these were different indications, each with their own standard of care. One of skill in the art would not look at successful treatment of one ocular disease (e.g., choroidal neovascularization, AMD, diabetic retinopathies, DME, CRVO, corneal neovascularization, or retinal neovascularization) and understand that another ocular disorder (e.g., BRVO) is necessarily treated.

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125. The paradigm-shifting success that anti-VEGF agents displayed with AMD made it obvious for retina specialists to consider the treatment of other vascular diseases, and the indications have risen over time. But this does not mean that a retina specialist would believe that successful treatment of one vascular disease necessarily followed from successful treatment of another. For example, cystoid macular edema (CME), which can be caused by a variety of disorders, presents similarly in an OCT image regardless of the cause. Despite the anatomical similarities, CME can respond differently to the VEGF Trap treatment, depending on the cause. We don't yet know why CME responds differently and sometimes the only way to distinguish between those underlying causes, given their anatomical similarities in an OCT image, is by testing the effects of anti-VEGF agents on a patient.

126. I would also like to provide a little more detail here regarding BRVO and CRVO, for which the pre-2013 application does disclose. Both involve impairments in the venous return system, but they are considered to be separate clinical entities for multiple reasons. Anatomically, BRVOs occur at a more distal part of the retinal venous tree, in which thickened, potentially atherosclerotic arteriole crosses a vein and impedes its flow. CRVOs occur when there is some obstruction on the other hand, occurring within the central retinal vein, and within

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the optic nerve or up to the lamina cribosa, where the vein exits the nerve to enter the eye. Because of its much more proximal location in the venous tree, the damage from a CRVO is typically much more extensive and can involve the entire retina, whereas BRVO typically involves only a sector. Although there are overlapping risk factors for CRVO and BRVO, there are differences. For instance, Asians and Hispanics appeared to have an elevated risk of BRVO compared to Caucasians, whereas no similar difference was found for CRVO. (Ex. 1037.)

127. Aside from anatomical and population distinctions between BRVO and CRVO, another reason that one of skill in the art would not assume that a treatment of CRVO would be equivalent to be a treatment of BRVO is the historical difference in response to treatments between them. Prior to the anti-VEGF era, BRVO and CRVO were considered separately in landmark ophthalmology studies. Because of this, one would not have assumed that a treatment for BRVO would work for CRVO and vice versa. Two of the most important trials for vein occlusion treatment in the pre-anti VEGF era were the Branch Vein Occlusion Study (BVOS) and the Central Vein Occlusion Study (CVOS). Macular grid laser was found to improve cystoid macular edema (CME) and vision in BRVO patients and ultimately became the standard of care of its time (and is still used in some patients today). (Ex. 1038.) However, macular grid laser

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was not found to improve vision in CRVO, and so the same treatment that was effective for BRVO was not recommended for CRVO. (Ex. 1039.)

128. The next series of landmark trials in BRVO and CRVO were the SCORE trials. BRVO and CRVO were again separated into distinct investigations. The conclusion of the SCORE BRVO trial was that there was no difference in visual outcome between standard of care to 1 mg or 4 mg triamcinolone, but the 4 mg triamcinolone arm had more side effects, so grid laser was still recommended. (Ex. 1040.) The conclusion of the SCORE CRVO trial was that triamcinolone at either dose improved visual acuity compared to standard of care, but that the 4 mg triamcinolone arm again had more side effects, making 1 mg triamcinolone a possible treatment option for CRVO. (Ex. 1041.) Thus, the SCORE CRVO and BRVO trials both considered the use of triamcinolone and reached different conclusions.

129. By this time, the VEGF antagonist ranibizumab was being tested for vein occlusions. Notably, there were separate trials performed for the study of ranibizumab's effects on BRVO and CRVO, supporting my opinion that one of skill in the art in 2011-2012 would not have assumed that treatments for one would necessarily work for the other. The BRAVO trial examined the efficacy of monthly ranibizumab for BRVO and the CRUISE trial examined the efficacy of

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monthly ranibizumab for CRVO. Both trials found that monthly ranibizumab for 6 months followed by PRN treatment resulted in anatomical and visual improvement. (Ex. 1042, Ex. 1043.) These trials ultimately did demonstrate that both conditions benefited from anti-VEGF treatment, but given the historical differences in treatment for BRVO and CRVO one would not have assumed this to necessarily be the case. Furthermore, longer term follow-up of these cohorts found that while many patients with BRVO retained their visual acuity gains despite fewer injections in the second year, the treatment was less durable for CRVO, again supporting that these two conditions are different from a clinical standpoint. (Ex. 1044.)

130. My opinion is supported by the '345 patent and its history. First, claim 8 lists BRVO as a disorder separate from AMD, diabetic retinopathy, DME, CRVO, and corneal neovascularization. If treatment of these disorders implicitly included treatment of BRVO, there would be no need for the inventor to list BRVO as a separate treatment. Second, the '345 patent family added BRVO to the disclosed embodiments in 2013 along with other newly added disorders, confirming that the inventors recognized later that BRVO (along with the other disorders) was treatable with their VEGF antagonist.

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131. Consistently, Regeneron structured their clinical trials differently for wet AMD, CRVO, and DME. In the 2009 Press Release, Regeneron reported (1) a Phase 3 clinical trial for wet AMD, (2) a Phase 2 clinical trial for CRVO, and (3) a Phase 2 clinical trial for DME. (Ex. 1005.) Each had different dosing regimens: (1) wet AMD was treated with scheduled doses of 0.5 milligram (mg) every four weeks, 2.0 mg every four weeks, or 2.0 mg every eight weeks (with one additional 2.0 mg dose at week four); (2) CRVO was treated with six monthly 2 mg doses, then on an as-needed basis; and (3) DME was treated with scheduled doses of 0.5 mg or 2.0 mg monthly, 2 mg on an as-needed basis after three monthly loading doses, or 2 mg every eight weeks after three monthly loading doses. The 2009 Press Release includes no mention of a BRVO clinical trial, much less a BRVO treatment plan with 4 week secondary doses and 12 week tertiary doses.

* * *

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I declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true, and that these statements were made with knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under section 1001 of Title 18 of the United States Code.

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Dated: 1/7/21

David Wu, M.D., Ph.D

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UNITED STATES PATENT AND TRADEMARK OFFICE

BEFORE THE PATENT TRIAL AND APPEAL BOARD

CHENGDU KANGHONG BIOTECHNOLOGY CO., LTD. Petitioner

V.

REGENERON PHARMACEUTICALS, INC. Patent Owner

Case PGR2021-00035 Patent 10,828,345

DECLARATION OF DIANA V. DO, M.D.

Regeneron Exhibit 2001 Page 01 of 35 Chengdu Kanghong Biotech. Co., Ltd. v. Regeneron Pharms., Inc., PGR2021-00035, U.S. Pat. 10,828,345, Exhibit 2001

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Regeneron Exhibit 2001 Page 02 of 35 I, Dr. Diana Do, declare as follows:

I. INTRODUCTION

1. I have been retained by counsel for Regeneron Pharmaceuticals, Inc. ("Regeneron") as a technical expert in connection with the above-captioned proceeding. I have been asked to provide my opinions and views on the materials I have reviewed in relation to the Petition for Post Grant review of U.S. Patent No. 10,828,345 (the "345 patent") (Ex. 1001), in particular the state of the art as of the earliest filing date ("priority date") of the '345 patent and responses to the opinion and views of Petitioner's declarant, David Wu, M.D., Ph.D. I submit this declaration in support of Regeneron's Patent Owner Preliminary Response ("POPR").

2. I am being paid at an hourly rate for my work on this matter. I have no personal or financial stake in the outcome of the present proceeding.

II. QUALIFICATIONS AND EXPERIENCE

3. I am a Professor of Ophthalmology and the Vice Chair for Clinical Affairs at the Byers Eye Institute at Stanford University School of Medicine and have been since 2017. I also serve as a Physician Improvement Leader at Byers Eye Institute, a position I have held since 2018. I have an active clinical and surgical practice and I work as a clinical investigator to study novel treatments for retinal diseases. In addition, I teach students, residents, and retina fellows at Stanford and am a member of the Stanford Ophthalmology Education Committee.

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Regeneron Exhibit 2001 Page 03 of 35 4. I graduated from the University of California Berkeley (summa cum laude) with a B.A. degree in Molecular and Cellular Biology in 1995 and earned my M.D. (Alpha Omega Alpha) from the University of California San Francisco School of Medicine in 1999. Following medical school, I completed an internship in internal medicine at Massachusetts General Hospital at Harvard Medical School. From 2000-2003, I completed my residency in Ophthalmology at the Wilmer Eye Institute at Johns Hopkins University School of Medicine, and then remained at the Wilmer Eye Institute for a Retina Fellowship in surgical and medical retina from 2003-2005.

5. From 2005 through 2010, I served as Assistant Professor of Ophthalmology and Assistant Head of the Retina Fellowship Training Program at the Wilmer Eye Institute. In 2011, I was promoted to Associate Professor and Head of the Retina Fellowship Training Program, positions I held through 2013.

6. In 2013, I joined the faculty at the Truhlsen Eye Institute at the University of Nebraska College of Medicine, where I became a full Professor of Ophthalmology in 2015. At the Truhlsen Eye Institute, I was Head of the Retina Fellowship Training Program and Program Director for the Ophthalmology Residency. In my leadership roles at the Truhlsen Eye Institute, I also served as Vice Chair of Education. I was recruited by Stanford University's Ophthalmology Department (the Byers Eye Institute) at Stanford in the beginning of 2017.

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7. As a physician-scientist, I am an international leader in the treatment of diabetic retinopathy and wet AMD ("wAMD"). My research has led to more than 140 peer-reviewed publications. My research interest focuses on evaluating the efficacy and safety of novel pharmacologic therapies for diabetic macular edema. diabetic retinopathy, wAMD, retinal vein occlusion, and ocular inflammation. I have led national and global clinical trials investigating intravitreal VEGF inhibitors (aflibercept and ranibizumab) for diabetic eve disease and wet AMD. Our research developed a greater understanding of how intraocular inhibition of VEGF reduces vascular permeability and angiogenesis in diabetic eye disease, thereby reducing diabetic macular edema and improving visual acuity. Before the onset of pharmacologic therapies, thermal laser photocoagulation was the only treatment option for diabetic macular edema and laser was not effective in improving vision. Our research led to new treatment paradigms and better vision outcomes for patients with diabetic macular edema, diabetic retinopathy, and wAMD. The results from the collaborative research that I led has revolutionized how ophthalmologists throughout the world think about and treat patients with VEGF-mediated retinal diseases.

8. Since 2009, I have been the lead investigator and a Steering Committee member for the evaluation of aflibercept, a fusion protein that inhibits VEGF, in diabetic macular edema. I initiated the first-in-human clinical trial of aflibercept. In

Regeneron Exhibit 2001 Page 05 of 35 addition, I also was the principal investigator on the Phase II and III clinical trials of aflibercept for diabetic macular edema to further evaluate efficacy, dosing regimens, and safety. My leadership in these global clinical trials, which enrolled over 1,000 subjects, contributed to FDA approval of aflibercept for diabetic macular edema. Aflibercept has also been approved by FDA for other angiogenic ocular diseases such as wAMD, central retinal vein occlusion, branch retinal vein occlusion, and diabetic retinopathy.

9. My research efforts have also led to a greater understanding of the role of ranibizumab, an intravitreal VEGF antibody fragment biologic, in diabetic macular edema. I was a lead investigator in the Ranibizumab for Edema of the Macula in Diabetes (READ) Study Group and was the lead author on multiple manuscripts evaluating the efficacy and safety of ranibizumab. The collaborative studies that I led contributed to understanding dosing regimens for intravitreal VEGF inhibitors, and led to the design of pivotal clinical trials involving ranibizumab for diabetic macular edema. Ranibizumab was the first FDA approved intravitreal VEGF inhibitor for diabetic macular edema, and helped to transform the management of diabetic retinopathy. I continue to lead clinical trials investigating new treatments for retinal diseases.

10. As a result of my research, I am recognized as an international thought leader on the subject of the retina and am regularly invited to lecture and teach at

Regeneron Exhibit 2001 Page 06 of 35 international and national meetings including the American Academy of Ophthalmology Retina Sub-Specialty Meeting, American Society of Retina Specialists, Asian Pacific Vitreo-Retinal Society Meeting, Canadian Ophthalmology Society Meeting, and congresses throughout Europe. I have organized and participated as a faculty member in national continuing medical education courses to teach my retina colleagues how to manage and treat diabetic macular edema, wet AMD, retinal vein occlusion, diabetic retinopathy, and other retinal disorders. Furthermore, I have held leadership positions at the American Society of Retina Specialists (Communications Committee Member to curate and develop online educational material), Women in Retina (Board Member and Secretary), Maryland Eye Society (President).

11. I am a practicing ophthalmologist with over 15 years of clinical and surgical practice in retina. I am a leader in the management of diabetic retinopathy, the leading cause of blindness in working age adults, and wAMD, the leading cause of vision loss in elderly individuals in developed countries. I have a high-volume clinical and surgical practice and spend approximately 1.5 days per week in clinic at the Byers Eye Institute and half-day per week at the Santa Clara Valley County Medical Center seeing patients in my clinical practice. In addition, I operate approximately one day per week at the Byers Eye Institute.

12. Given my extensive experience and research on diabetic retinopathy

Regeneron Exhibit 2001 Page 07 of 35 and wAMD, I have become the expert retinal specialist and surgeon in our department for evaluating these chronic eye diseases. Since joining Stanford's Ophthalmology Department, I have also become one of the highest volume retina surgeons among our faculty. Because proliferative diabetic retinopathy can lead to tractional retinal detachment and bleeding within the eye, I am referred complex cases that often require clinic-based treatments (such as intravitreal injections of medicines or pan retinal laser photocoagulation) or surgical management. Since I have clinical and research expertise using intravitreal vascular endothelial growth factor (VEGF) inhibitors in wAMD, ophthalmologists refer patients to me for consultation or co-management, particularly of chronic cases that have not responded to therapy. The majority of my patients are from the Bay Area or central California, and approximately 10% travel from more than 5 hours away to seek my expert opinion. I have been recognized as a "Top Doctor" in the Bay Area for the past three years. A current copy of my curriculum vitae is included at Ex. 2022.

III. SUMMARY OF OPINIONS

13. My opinions and views set forth in this declaration are based on my education, training, research, and clinical experience in ophthalmology, specifically in researching and treating retinal diseases, as well as the materials I reviewed in preparing this declaration and the state of scientific knowledge in the art pertaining to the subject matter of the '345 patent at the time of its earliest priority application.

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Regeneron Exhibit 2001 Page 08 of 35 14. In forming my opinions, I have reviewed the following materials: (a) the Petition for Post Grant Review of the '345 patent, PGR2021-00035, including all cited exhibits, (b) all priority applications leading to the issuance of the '345 patent, (c) all other documents and references herein, and (d) the Patent Owner's Preliminary Response to which my declaration relates.

15. For purposes of preparing this declaration in support of Patent Owner's Preliminary Response, I have been asked to apply Dr. Wu's definition of a person of ordinary skill in the art (who I also refer to as the "skilled artisan"): a person with a medical doctorate, internship and residency in ophthalmology and either a 1-year medical retina fellowship or a 2-year vitreoretinal surgical fellowship. Ex. 1003 ¶ 82. Likewise, for purposes of preparing this declaration, I have been informed and understand that the earliest filing date of the '345 Patent is January 13, 2011, based on the filing of a Provisional Application on that date.¹

16. It is my opinion that by January of 2011, a person of ordinary skill in the art would have understood that branch retinal vein occlusion ("BRVO") was an "angiogenic eye disorder" that could be treated by a VEGF inhibitor. Likewise, it is my opinion that by January of 2011, the person of ordinary skill would have

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¹ Although most of my opinions as to a skilled artisan expressed in this declaration are as of January 2011, I express some opinions as to the state of the art as of November 2011. In either case, I am applying the same definition of the skilled artisan.

understood that successful treatment of central retinal vein occlusion ("CRVO") with a VEGF inhibitor indicates that treatment of BRVO would also be successful.

17. It is also my opinion that by 2011, a skilled artisan would have understood that a fixed quarterly dosing regimen of ranibizumab, as disclosed in the Shams patent publication and corresponding PIER clinical trial, was a failure and not an effective method of treating an angiogenic eye disorder.

IV. THE '345 PATENT

A. Claim 1

18. The '345 patent has one independent claim, claim 1:

A method for treating an angiogenic eye disorder in a patient, said method comprising

sequentially administering to the patient

a single initial dose of a VEGF antagonist,

followed by one or more secondary doses of the VEGF antagonist,

followed by one or more tertiary doses of the VEGF antagonist;

wherein each secondary dose is administered 4 weeks after the immediately preceding dose; and

wherein each tertiary dose is administered 12 weeks after the immediately preceding dose;

wherein the VEGF antagonist is a receptor-based chimeric molecule comprising an immunoglobin-like (Ig) domain 2 of a first VEGF receptor which is Fltl and Ig domain 3 of a second VEGF receptor which is Flk1, and a multimerizing component.

Ex. 1001 at 21:55-22:55

19. The dosing regimen of Claim 1 is directed to the treatment of any type

of angiogenic eye disorder with a set of VEGF antagonist fusion proteins that

Regeneron Exhibit 2001 Page 10 of 35 comprise an "immunoglobin-like (Ig) domain 2 of a first VEGF receptor which is Fltl and Ig domain 3 of a second VEGF receptor which is Flk1, and a multimerizing component."

20. The dosing regimen of Claim 1 requires treatment of an angiogenic eye disorder by administration of an initial dose of the claimed VEGF antagonist followed by one or more "secondary" doses administered four weeks after the preceding dose, and then one or more "tertiary" doses that are administered at twelve week intervals following the preceding dose.

21. Claim 1 requires that "tertiary dose[s]" are "administered 12 weeks after the immediately preceding dose." Ex. 1001 at 21:59-64. As of the filing date, and even today, the term "tertiary dose" does not have a well-understood meaning to/ a skilled artisan in the fields of ophthalmology or retina medicine. In my experience, the term "tertiary dose" is not typically used by clinicians or the skilled artisan.

B. Claim 2

22. Dependent Claim 2 is directed to the method for treating angiogenic eye disorders with aflibercept, which is the unique fusion protein in Regeneron's Eylea product. Ex. 1001 at 22:56-57.

C. Claim 8

23. Dependent Claim 8 recites "wherein the angiogenic eye disorder is

Regeneron Exhibit 2001 Page 11 of 35 selected from the group consisting of: age related macular degeneration, diabetic retinopathy, diabetic macular edema, central retinal vein occlusion, branch retinal vein occlusion, and corneal neovascularization." Ex. 1001 at 23:3-8.

24. Claim 8 specifically lists types of eye disorders with pathological angiogenic characteristics, including several of the most significant "angiogenic eye disorders." A skilled artisan would recognize that these listed eye disorders are the major angiogenic eye disorders treated with VEGF antagonist pharmaceutical compounds.

25. The '345 Patent states that it is related to three provisional applications filed in 2011 — Provisional Application No. 61/432,245 (filed Jan. 13, 2011), Provisional Application No. 61/434,836 (filed Jan. 21, 2011), and Provisional Application No. 61/561,957 (filed Nov. 21, 2011). The January 13, 2011 Provisional Application, the earliest priority application, discloses methods that can be used to treat "any angiogenic eye disorder," which is defined as "any disease of the eye which is caused by or associated with the growth or proliferation of blood vessels or by blood vessel leakage." Ex. 1045 at [0024]. The January 13, 2011 Provisional Application also provides non-limiting examples of angiogenic eye disorders including "choroidal neovascularization, age-related macular degeneration (AMO), diabetic retinopathies, diabetic macular edema (DME), central retinal vein occlusion (CRVO), corneal neovascularization, and retinal neovascularization." *Id.* The

Regeneron Exhibit 2001 Page 12 of 35 provisional application also provides clinical trial data for Regeneron's aflibercept phase I, II, and III trials in wet AMD, and a phase II trial in DME. *Id.* at [0034]-[0041]. The November 21, 2011 Provisional Application adds the additional disclosure of clinical trial data for Regeneron's aflibercept phase III trial in CRVO. Ex. 1047 at [0064]-[0066].

V. STATE OF THE ART AS OF JANUARY 2011

A. Angiogenesis and VEGF Inhibition

26. Angiogenic eye disorders, also referred to as neovascular ocular diseases, are a group of diseases characterized by pathologic growth of abnormal blood vessels in the eye and vascular leakage from damaged, pre-existing blood vessels. Both events can lead to severe visual impairment. Angiogenesis is process controlled by a series of angiogenic agents such as growth factors, cytokines, and extracellular matrix components. One such agent is vascular endothelial growth factor ("VEGF"), a glycoprotein that acts as a potent proangiogenic factor. It has been well established that there is a correlation between elevated levels of VEGF and the presence of angiogenic eye disorders. Ex. 2004 at 23. Studies have demonstrated that elevated levels of VEGF are sufficient to induce ocular neovascularization and vascular leakage. *Id.*

27. VEGF plays multiple roles in the pathology of the angiogenesis in the retina. First, VEGF is a potent inducer of vascular permeability, which causes blood

Regeneron Exhibit 2001 Page 13 of 35 vessels to leak and results in macular edema (swelling in the central retina) that causes vision impairment and is a common feature of angiogenic eye disorders. Ex. 2004 at 23. VEGF expression is upregulated by hypoxia (low oxygen in the tissue); hypoxia in the retina is commonly seen with retinal vascular diseases such as diabetic retinopathy, central retinal vein occlusion, and branch retinal vein occlusion where blood vessels in the retina are damaged and thereby fail to supply adequate oxygen to the retina. *Id.* Increased levels of VEGF in turn promote vascular permeability and angiogenesis, both of which threaten vision.

28. By January of 2011, a person of ordinary skill in the art recognized that a hallmark of angiogenic eye disorders was excess levels of VEGF in the eye. Correlations between elevated ocular levels of VEGF and presentation of ocular neovascular disease had been demonstrated in conditions such as iris neovascularization, retinal vein occlusion, diabetic retinopathy, diabetic macular edema, neovascular glaucoma, wAMD, and retinopathy of prematurity. Ex. 2004 at 23. The ordinarily skilled retinal specialist in 2011 understood that the term "angiogenic eye disorders" was a well-defined group of neovascular diseases. Indeed, there were no more than a few dozen types and subtypes of diseases that would be understood as comprising the universe of angiogenic eye disorder as of 2011. Ciulla and Rosenfeld illustrate in their 2009 publication in Current Opinion in Ophthalmology that there were nine distinct categories of neovascular eye

Regeneron Exhibit 2001 Page 14 of 35 diseases, some of which exhibited neovascular characteristics that defined a subtype of that disease category. Ex. 2003 at Table 1.

29. By January of 2011, it was also well-established that inhibition of VEGF was a method for reducing this pathologic angiogenesis, and thereby treating the angiogenic eye diseases and improving vision prognosis. Ex. 2004 at 23. "The discovery of VEGF-A's role in the pathogenesis of neovascular ocular disease provided a strong rationale for the development of anti-VEGF-based therapies. There is now ample evidence that anti-VEGF therapies are viable treatment options for these [neovascular eye] diseases." Ex. 2003 at 1.

B. Methods of Treating Angiogenic Eye Disorders in the Art

30. Wet AMD is an angiogenic eye disorder characterized by abnormal growth of new blood vessels in the macula, the central portion of the retina responsible for high-resolution vision. Ex. 2025 at 2. Historically, wAMD was a devastating diagnosis that frequently led to irreversible vision loss. Early treatments with laser and photodynamic therapy would often, at best, slow inevitable vision loss. At worst, these treatments could cause further vision damage through retinal scarring. Wet AMD was the first angiogenic eye disorder where anti-VEGF agents were widely tested as a potential therapy. By 2006, however, two large Phase III clinical trials, MARINA and ANCHOR, demonstrated that intravitreal administration of an anti-VEGF antibody fragment, ranibizumab (or "Lucentis"), not

Regeneron Exhibit 2001 Page 15 of 35 only slowed vision loss associated with wAMD, but could actually improve vision. Similar efficacy was likewise demonstrated by the use of another anti-VEGF antibody bevacizumab (or "Avastin") through off-label case studies. *E.g.*, Ex. 2024.

31. The MARINA trial ran from March of 2003 to December 2005 and enrolled 716 patients with AMD with either minimally classic or occult choroidal neovascularization. Ex. 2025 at 1. Patients were randomly assigned to received 24 monthly intravitreal injections of Lucentis (either 0.3 mg or 0.5 mg) or sham injections. *Id.* The primary endpoint of the study was the proportion of patients losing fewer than 15 letters from baseline visual acuity at 12 months. *Id.*

32. The ANCHOR trial ran from May of 2003 to September of 2006 and enrolled 423 patients with predominantly classic choroidal neovascularization in wAMD. Ex. 2026 at 1. Patients were randomized on a 1:1:1 ratio to receive monthly intravitreal Lucentis (0.3 mg or 0.5 mg) plus sham photodynamic verteporfin therapy or monthly sham injections plus active verteporfin therapy. *Id.* As in MARINA, the primary endpoint of the study was also the proportion of patients losing fewer than 15 letters from baseline visual acuity at 12 months. *Id.*

33. The two-year results of the MARINA trial were published in the New England Journal of Medicine on October 5, 2006. Ex. 2025 at 1. On the same day, the one-year results of the ANCHOR trial were also published in the New England Journal of Medicine. Ex. 2026 at 1. The two-year results of ANCHOR were

Regeneron Exhibit 2001 Page 16 of 35 published in January 2009 in Ophthalmology. Ex. 2027 at 1.

34. The MARINA and ANCHOR trials demonstrated that monthly intravitreal Lucentis could not only effectively prevent vision loss, but could actually lead to mean improvements in vision that were sustained throughout the second year of the studies. In the MARINA trial, mean increases in visual acuity were +6.5 letters in the 0.3 mg group and +7.2 letters in the 0.5 mg groups, compared with a decrease of -10.3 letters in the sham-injection group. Ex. 2025 at 1. In fact, visual acuity improved by 15 or more letters in 24.8% of the 0.3-mg group and 33.8% of the 0.5-mg group. *Id.* Likewise, in the ANCHOR study, visual acuity was improved from baseline, on average, by +8.1 to +10.7 letters, versus a mean decline of -9.8 letters in the verteporfin photodynamic group. Ex. 2026 at 1.

35. Lucentis received FDA approval for the treatment of wet AMD in June 2006. As demonstrated in its label, monthly injections of Lucentis resulted in sustained gains in visual acuity as compared to baseline vision. Ex. 2028 at 7. The successes seen in the treatment of wAMD with these anti-VEGF agents were a gamechanger for the treatment of angiogenic eye disorders. As Ciulla and Rosenfeld noted in 2009, "[t]he success of anti-vascular endothelial growth factor (VEGF) therapies in neovascular age-related macular degeneration (AMD) has spurred investigation of similar treatment strategies for other exudative ocular diseases." Ex. 2003 at 1.

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C. RVO, BRVO, and CRVO

36. Retinal vein occlusions ("RVO") was a recognized category of neovascular eye disease well-before January of 2011. Ex. 2004 at 23. CRVO and BRVO are closely related angiogenic eye disorders that were both known to fall within the category of RVOs. Ex. 2003 at Table 1 (RVO includes "Central RVO, hemicentral RVO, or branch RVO" based on the specific neovascular characteristics exhibited).

37. BRVO and CRVO share numerous disease characteristics including the development of a thrombus in the retinal vein resulting in reduced blood flow, dilation and tortuosity of the affected and damaged veins, retinal hemorrhages, cotton wool spots, evidence of ischemia, up-regulation of VEGF, and subsequent macular edema. If there is extensive ischemia in the retina, retinal neovascularization develops and can lead to vitreous hemorrhage and severe vision loss. Shared risk factors for RVOs include older age, arteriosclerosis, systemic arterial hypertension, and diabetes. Ex. 2029 at 1. The principal difference with these RVO subtypes is the locus of the occlusion. In CRVO, there is an obstruction of the retinal vein at or posterior to the optic nerve head, while in BRVO there is complete or partial obstruction at a branch or tributary of the central retinal vein.

38. As with other angiogenic eye disorders, the art recognized that anti-VEGF treatments could be a viable therapeutic option for patients with BRVO (and

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Regeneron Exhibit 2001 Page 18 of 35 CRVO) well-before 2011. As noted by clinicians at this time, "[h]igh levels of VEGF have been found in the aqueous humor of patients with ME [macular edema] secondary to vein occlusion. Accordingly, patients with higher levels of VEGF often have more severe cases of ME [macular edema]. Therefore, anti-VEGF therapy would seem a reasonable treatment option in these cases." Ex. 2006 at 2.

39. As early as 2005, researchers began testing the use of anti-VEGF agents, beginning with off-label intravitreal Avastin, on patients with BRVO and CRVO. Ex. 2029 at 1 ("Since the first report of the efficacy of intravitreal bevacizumab...in a patient with macular oedema secondary to CRVO in 2005, several retrospective case series have shown the benefit of this treatment, with an improvement in visual acuity and a decrease of central retinal thickness (CRT) in patients with macular oedema associated with both BRVO and CRVO.")

40. By 2009, studies "demonstrated rapid visual improvements after VEGF inhibition with ranibizumab and bevacizumab in patients with CRVO and BRVO." Ex. 2003 at 5. Despite being an off-label use, in 2009 the Patterns and Trends Survey by the American Society of Retina Specialists showed that approximately 50% of respondents used intravitreal Avastin as a first-line therapy for CRVO and BRVO. Ex. 2030 at 2.

41. By 2010, Genentech had completed full Phase III randomized controlled trials that assessed the efficacy and safety of intravitreal ranibizumab

Regeneron Exhibit 2001 Page 19 of 35 (Lucentis®) in BRVO and CRVO. The BRAVO trial, which began recruitment in July 2007, evaluated ranibizumab injections compared with sham in patients with macular edema secondary to BRVO. In the BRAVO trial, 397 patients were randomly assigned to six monthly injections of ranibizumab, either 0.3 mg or 0.5 mg, or to sham injections. Ex. 1042 at 1. The primary efficacy outcome was mean change from baseline BCVA ("Best Corrected Visual Acuity") at 6 months. Secondary outcomes included the percentage of patients who gained 3 lines (15 letters) of BCVA at 6 months. *Id.* The mean visual acuity gain from baseline at month 6 was +16.6 letters in patients receiving 0.3 mg of ranibizumab, +18.3 letters in those receiving 0.5 mg, and +7.3 in those receiving sham injection. *Id.* at 2.

42. The CRUISE trial, which ran concurrently with the BRAVO trial and shared the same outcome measurements, evaluated ranibizumab injections compared with sham injections in patients with macular edema secondary to CRVO. Ex. 1043 at 1. The "results of CRUISE mirror those of BRAVO." Ex. 2030 at 2. In the CRUISE trial, 392 patients were randomized, the mean gain from baseline BCVA at month 6 was +12.7 letters in patients who received 0.3 mg ranibizumab, +14.9 letters in patients who received 0.5 mg ranibizumab, and +0.8 letters in those who received sham injections. Ex. 1043 at 1.

43. By June 2010, FDA had approved the use of Lucentis® (ranibizumab) on a monthly basis for the treatment of BRVO and CRVO. Ex. 2005 at 9.

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VI. CLAIM 8 IS ADEQUATELY SUPPORTED IN THE ORIGINAL PROVISIONAL FILING

A. A Skilled Artisan Would Have Understood the January 13, 2011 Provisional Application's Disclosure of "Angiogenic Eye Disorders" to Include BRVO

44. The January 13, 2011 Provisional Application explicitly states that "[t]he methods of the present invention can be used to treat any angiogenic eye disorder, including, e.g., age related macular degeneration, diabetic retinopathy, diabetic macular edema, central retinal vein occlusion, corneal neovascularization, etc." Ex. 1045 at [0024]. The same application also defines "any angiogenic eye disorder," which is defined as "any disease of the eye which is caused by or associated with the growth or proliferation of blood vessels or by blood vessel leakage." Ex. 1045 at [0024].

45. By January 2011, the set of "angiogenic eye disorders" was welldefined and, further, BRVO was widely recognized as an angiogenic eye disorder in that set. In addition, as discussed above, by January of 2011, BRVO had been successfully treated with anti-VEGF therapies, including bevacizumab and ranibizumab, which was widely reported in the literature. Further, before January of 2011, ranibizumab had been approved by FDA for treatment of BRVO by monthly intravitreal injection.

46. Given the known underlying pathology for angiogenic eye disorders and established efficacy of anti-VEGF agents in ameliorating that pathology, it is

Regeneron Exhibit 2001 Page 21 of 35 my opinion that one of ordinary skill in the art, with Regeneron's January 13, 2011 Provisional Application in hand, would have understood it to be teaching that BRVO was a type of "angiogenic eye disorder," that could be treatable with a VEGF antagonist.

47. At paragraph 124 of his declaration, Dr. Wu suggests that in 2011-2012, one of ordinary skill in the art would not look at the successful treatment of one angiogenic ocular disease (e.g., choroidal neovascularization, AMD, diabetic retinopathies, DME, CRVO, corneal neovascularization, or retinal neovascularization) and understand that another ocular disorder (e.g., BRVO) could be treated.² Ex. 1003 ¶ 124. I disagree with Dr. Wu's assertion.

48. As noted above, by 2011, anti-VEGF agents had demonstrated efficacy with respect to many types of angiogenic eye disorders. Importantly, by 2011, anti-VEGF agents bevacizumab and ranibizumab (a VEGF antibody and a VEGF antibody fragment, respectively) had been shown to effectively treat BRVO. Indeed, Dr. Wu acknowledges that Genentech's BRAVO and CRUISE trials (Phase III trials of ranibizumab in BRVO and CRVO, respectively) demonstrated that both conditions benefit from anti-VEGF treatment. Ex. 1003 ¶ 129. The results of these

² While I have reviewed the entirety of Dr. Wu's declaration and there are many points on which he and I disagree, I do not attempt to respond or rebut each of these points of difference in this declaration. Rather, I reserve the right to respond more fully to Dr. Wu's declaration at a future date if asked to do so.

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trials were presented in conferences around the world beginning in May of 2010. *See* Ex. 1042 at 9. In fact, in 2010, ranibizumab (Lucentis®) was FDA approved for "Macular Edema Following Retinal Vein Occlusion" without differentiating between BRVO or CRVO in the label. Ex. 2005 at 9. Given the successful experience in treating a variety angiogenic disorders, including BRVO and CRVO, with anti-VEGF agents before 2011, it is my opinion that a skilled artisan reading the January 13, 2011 Provisional Application, would have understood it to be teaching that "BRVO" was among the "angiogenic eye disorders" that could be treated with the disclosed treatment regimens.

49. In addition, at paragraph 125 of his declaration, Dr. Wu opines that a retina specialist would not think that anti-VEGF therapy could work for all vascular diseases, just because it worked for one such disease. In support of his assertion, Dr. Wu offers the example of the differential response of cystoid macular edema ("CME") to anti-VEGF treatment. Again, I disagree with Dr. Wu's statement and believe that his reliance on CME as a supporting example is misplaced. Dr. Wu ignores the fact that critical features of CME would directly, and predictably, impact the efficacy of an anti-VEGF agent on treatment of that disorder. Ex. 1003 ¶ 125. CME is a disease that has multiple etiologies; some, but not all, cases of CME are caused by angiogenesis. CME can also be caused by other factors, for example, inflammation after cataract surgery. Where CME is caused by inflammation after

Regeneron Exhibit 2001 Page 23 of 35 cataract surgery, elevated levels of VEGF do not play a central role in the pathology as one sees in an ocular disorder that is characterized by angiogenesis specifically, and thereby inflammation-associated CME is usually treated with topical antiinflammatory medicines. RVOs, including BRVO, by contrast, are normally associated with upregulated VEGF and, as a consequence, are susceptible to anti-VEGF therapy. Dr. Wu's CME example is thus a highly imperfect analogy to a skilled artisan's expectations with respect to BRVO.

B. A Skilled Artisan Would Have Understood the 2011 Provisional Applications To Teach that Regeneron's VEGF Antagonist Fusion Proteins Would Treat BRVO

50. The January 13, 2011 Provisional Application explicitly stated that one of the "angiogenic eye disorders" that could be treated by the disclosed methods was CRVO. This supports my opinion that a skilled artisan reviewing the provisional disclosures would have understood that an anti-VEGF agent would be an effective treatment for BRVO.

51. As noted above, BRVO and CRVO were known to be closely related types of RVOs that share numerous disease characteristics. As noted above, by January of 2011 it had become standard practice to treat both CRVO and BRVO with anti-VEGF agents and ranibizumab was tested in parallel Phase III trials in CRVO and BRVO. In fact, the retina community often described the results of BRAVO and CRUISE in tandem and did not distinguish between the subtypes when

Regeneron Exhibit 2001 Page 24 of 35 announcing that anti-VEGF agents were effective for treatment of these closely related disorders. For example, David Brown, M.D., a clinical investigator on the studies noted that the trials "showed that with intensive, monthly treatment, patients achieve very good results, superior to anything we have seen previously with other treatment modalities." Ex. 2030 at 1. Other clinicians remarked of BRAVO and CRUISE "[t]hus far, off-label use of anti-VEGF drugs has been shown to effectively target the underlying pathogenesis associated with the development of ME secondary to vein occlusion." Ex. 2006 at 2.

52. Given the successful outcomes of these Lucentis Phase III trials for both BRVO and CRVO, an ordinarily skilled artisan would have understood that positive clinical trial results seen with one anti-VEGF agent in CRVO could forecast successful outcomes in a BRVO patient with that same anti-VEGF agent.

53. An ordinarily skilled artisan with the benefit of Regeneron's January 13, 2011 Provisional Application would have understood that aflibercept, a VEGF antagonist fusion protein, had demonstrated efficacy in two Phase III pivotal trials in wAMD. The January 13, 2011 Provisional Application describes these Phase III studies (in Example 4) and reports clinical trial results at the primary endpoint of 52 weeks. Ex. 1045 at [0038]-[0060]. The Phase III clinical results show that by week 52 in Study 1, the patients receiving aflibercept 2 mg every four weeks gained +10.9 letters and patients dosed every eight weeks gained +7.9 letters. *Id.* at [0038].

Regeneron Exhibit 2001 Page 25 of 35 Similarly, in Study 2, patients who received aflibercept 2 mg every four weeks gained +7.6 letters and patients dosed every eight weeks gained +8.9 letters. *Id.*

54. In addition, Regeneron's January 13, 2011 Provisional Application disclosed that aflibercept was effective in a Phase II trial in DME. Ex. 1045 at [0061]-[0063]. Clinical trial results were reported at 24 weeks and 52 weeks. The Phase II results show that by week 52, patients who received aflibercept 2 mg every four weeks gained + 13.1 letters. *Id.* at [0061]. Likewise, patient who received aflibercept 2 mg every eight weeks gained +9.7 letters. *Id.*

November 21, 2011 Provisional 55. Furthermore, Regeneron's Application disclosed clinical trial results from the COPERNICUS trial for the treatment of CRVO with aflibercept. Ex. 1047 at [0064]-[0066]. The COPERNICUS trial was a randomized, double-masked, phase III study where patients received 6 monthly injections of either 2 mg intravitreal aflibercept or sham injections. From week 24 to week 52 of the study, all patients received 2 mg aflibercept on a PRN (as-needed) basis according to pre-defined retreatment criteria. *Id.* at [0064]. The primary endpoint for the trial was the proportion of patients who gained ≥ 15 letters from baseline at week 24. The November 21, 2011 Provisional Application disclosed that at week 24, 56.1% of patients treated with aflibercept gained ≥ 15 letters as compared to 12.3% of sham treated patients. Id. at [0065]. And, at week 52, 55.3% of patients treated with affibercept gained ≥ 15 letters as

Regeneron Exhibit 2001 Page 26 of 35 compared to 30.1% of sham treated patients; the aflibercept arm gained a mean of +16.2 letters vs. +3.8 letters for sham patients. *Id.* This disclosure clearly demonstrated to a skilled artisan that 2 mg intravitreal aflibercept produced statistically significant improvements in visual acuity that were maintained through week 52 on the PRN dosing regimen.

56. Given the positive results reported in Regeneron's 2011 Provisional Applications regarding clinical trials with aflibercept in wAMD and DME, a person of ordinary skill in the art would have viewed the disclosure of COPERNICUS results in CRVO as a clear signal that Regeneron's anti-VEGF fusion protein therapy would be successful in treating BRVO.

57. At paragraphs 126-127 of his declaration, Dr. Wu opines that given known differences in the anatomy, affected patient population, and historical treatment differences between BRVO and CRVO, a person of ordinary skill in the art would not have assumed that a treatment for CRVO would equate to a treatment of BRVO. Ex. 1003 ¶126-127.

58. I disagree. Notably, Dr. Wu relies on historical differences in response to treatment "[p]rior to the anti-VEGF era" (Ex. 1003 ¶127) but by 2011, those historical differences were no longer relevant.

59. For example, at paragraphs 127-28, Dr. Wu discusses the disparate impact of macular grid laser treatment, a historical treatment modality, on BRVO as

Regeneron Exhibit 2001 Page 27 of 35 compared to CRVO. Ex. 1003 ¶ 127-28. More specifically, macular grid laser treatment was found to improve vision for BRVO patients but not CRVO patients. However, Dr. Wu neglects to include important context about the nature of the diseases and how it relates to the particular treatment. As noted earlier, the key difference between BRVO and CRVO is the locus of the retinal vein occlusion. When using laser therapy treatment, this anatomic difference is highly relevant. BRVO responded better to laser treatment because the area of the vein occlusion is smaller, and a clinician can more easily target the specific retinal area that needs treatment. On the other hand, CRVO is posterior to the optic nerve and the diseased area is more extensive—it is impacting all four retinal quadrants – and macular laser is not effective in CRVO. These anatomic impacts on the efficacy of laser therapy have no relevance to the efficacy of anti-VEGF treatment, which seeks to arrest the cause of the vascular leakage and neovascularization in the first place by inhibiting the VEGF pathway.

60. By 2011, anti-VEGF therapy had been demonstrated to be effective for the treatment of BRVO and CRVO. In fact, Genentech's Lucentis® (ranibizumab) had been FDA-approved for the treatment of BRVO and CRVO by June of 2010. Ex. 2005 at 9. Thus, Dr. Wu's hypothetical concerns regarding differences in anatomy and historical treatment modalities for BRVO and CRVO (Ex. 1003 ¶126-127) were mooted once VEGF inhibition was demonstrated to be effective for

Regeneron Exhibit 2001 Page 28 of 35 treatment of BRVO. Notably, Dr. Wu fails to acknowledge the critical role that Avastin had played in clinical practice for both of these RVO subtypes well before the 2011-2012 time frame during which he asserts that a skilled artisan "would not have assumed" that "both conditions could benefit from anti-VEGF." Ex. 1003 ¶ 129.

61. Dr. Wu also tries to differentiate BRVO and CRVO by noting that patient populations of different ethnicities have disparate risks for BRVO, but not for CRVO. Ex. 1003 ¶ 126. This is again a difference without a distinction. By 2011, anti-VEGF therapy had been shown to be effective in large, randomized Phase III clinical trials and there were no proven studies in 2011 (nor are there as of the present day) that show a disparate response to anti-VEGF therapy in patient populations of different ethnicities for either CRVO or BRVO.

62. Simply put, given the closely related nature of CRVO and BRVO and the demonstrated efficacy of anti-VEGF agents in treating both conditions by January 2011, it is my opinion that the disclosure of CRVO in the January 13, 2011 Provisional Application would have bolstered a skilled artisan's understanding that BRVO was an angiogenic eye disorder treatable by the disclosed methods of the '345 Patent.

VII. SHAMS DISCLOSED AN UNSUCCESSFUL 12-WEEK DOSING REGIMEN

63. I understand that Petitioner has asserted that the dosing regimen of

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Regeneron Exhibit 2001 Page 29 of 35 Claim 1 of the '345 Patent is not novel in light of Shams (Ex. 1004), a Genentech patent application that was published on May 4, 2006. I have reviewed Shams' disclosure and recognize that the Trial Design (Figure 1) and Treatment Schema (Figure 2) set forth in Shams, as well as its description of a dosing regimen in Example 1, all correspond to Genentech's PIER Study, a clinical trial of ranibizumab (Lucentis®). *Compare* Ex. 1004, Figure 2, *with* Ex. 1026 at 2.

64. The PIER study, which ran from August 2004 to March 2007, was designed to compare three monthly loading doses followed by fixed quarterly dosing of 0.3 mg and 0.5 mg Lucentis® (ranibizumab) against sham control over 24 months. Ex. 1026. at 2. This same dosing regimen is outlined in Figure 2 of Shams, which illustrates the administration of the "first individual doses" at months 0, 1, and 2, followed by the "secondary doses" at months 5, 8, 11, and continuing every 3 months through 24 months. Ex. 1004, Figure 2.

65. During the first year of the PIER Study, while the treatment arms gained visual acuity during the three monthly loading doses, those visual acuity gains were lost when patients transitioned into the quarterly fixed dosing period of the treatment regimen. Ex. 1026, Figure 1. Worse yet, by the end of month 12, both treatment arms had on average lost letters as compared to baseline. *Id.* In addition to the visual acuity losses reported in PIER, *post-hoc* analyses of the study data showed that patients in the treatment arm of PIER saw no benefit in the incidence of macular

Regeneron Exhibit 2001 Page 30 of 35 hemorrhage as compared to sham control and, in fact incidence rates were numerically higher. Ex. 2020 at 3, 7. By the time PIER year one results were first presented, in September of 2006, Genentech had run two phase III ranibizumab trials —MARINA and ANCHOR — that demonstrated the efficacy of monthly intravitreal injections of Lucentis. Both clinical trials showed that Lucentis could improve visual acuity, and maintain those vision improvements over the course of treatment when monthly therapy was administered. Ex. 2025; Ex. 2026. Given the historical challenges in effectively treating wAMD and the significant risk of permanent vision loss if treatment was delayed, the disclosure of these positive results swiftly impacted the community standard of care for wAMD.

66. In view of the results of the MARINA and ANCHOR trials, the PIER study sponsor recognized that a sham control arm was no longer acceptable and the study protocol was amended in February 2006 to allow control subjects to cross over to 0.5 mg ranibizumab for the remainder of the treatment period. Ex. 1026 at 2. In addition, in light of the highly disappointing first year results of the treatment arms, the PIER study organizers amended the treatment protocol in August of 2006 to allow all patients in the quarterly treatment arms to rollover and receive monthly injections of 0.5mg ranibizumab through the remainder of the study. Ex. 1026 at 2. In my experience as a clinical investigator, protocol amendments on this scale, in the middle of a study, are typically only implemented when there are serious safety

Regeneron Exhibit 2001 Page 31 of 35 or efficacy concerns with drug or dosing regimen.

67. I note that in paragraph 111 of Dr. Wu's declaration, he suggests that fixed q4/q12 dosing of Lucentis (the Shams disclosure) was effective based on (1) the Lucentis 2006 label; and (2) the Regillo publication reporting first year results of the PIER Study. Ex. 1003, ¶ 111. I disagree with Dr. Wu's suggestion.

68. The Lucentis label was first approved by FDA for the treatment of wet AMD in 2006. The Dosage and Administration section of the label recommends monthly dosing of Lucentis: "LUCENTIS 0.5 mg (0.05 mL) is recommended to be administered by intravitreal injection once a month." Ex. 2028 at 2. Lucentis was not approved based on the Shams protocol or PIER data. Indeed, the Genentech Press Release that Dr. Wu cites in his declaration makes clear that "FDA approval of LUCENTIS is based on data from two large Phase III clinical trials (MARINA and ANCHOR)," which tested monthly injections of ranibizumab. Ex. 1006 at 2. The press release also notes: "In addition to data from the two pivotal studies, data from the Phase I/II FOCUS and Phase IIIb PIER studies were included in the FDA review." Id. The inclusion of PIER data in the Lucentis label does not suggest that q4/q12 Lucentis was an effective method of treating wet AMD. To the contrary, Lucentis' FDA-approved label reflects the concerns raised by both FDA and the study sponsor based on the results of the PIER trial. The label states "Although less effective, treatment may be reduced to one injection every three months after the

Regeneron Exhibit 2001 Page 32 of 35 first four injections if monthly injections are not feasible. Compared to continued monthly dosing, dosing every 3 months will lead to an approximate 5-letter (1-line) loss of visual acuity benefit, on average, over the following 9 months. Patients should be evaluated regularly." Ex. 2028 at 2. In addition, the clinical studies section of the Lucentis label (at §§ 14.1 and 14.2) shows the dramatic difference between the trial results in studies 1 and 2 (ANCHOR and MARINA), where patients gained visual acuity in the treatment arm during the study (see Figure 1) versus study 3 (PIER), where patients lost visual acuity in the treatment arm (see Figure 2). *Id.* at 7. An ordinarily skilled retinal specialist would not read the Lucentis label language as an FDA endorsement for fixed quarterly dosing of Lucentis. Rather, this language would be viewed as a warning to retina practitioners that this dosing regimen carriers a high risk for vision loss.

69. Dr. Wu selectively relies on a single sentence in the conclusion of the paper, that "Ranibizumab administered monthly for three months and then quarterly provided significant VA benefit to patients with AMD-related subfoveal CNV and was well tolerated." Ex. 1003 ¶ 111; Ex. 1026 at 1. But Dr. Wu's reliance on a single sentence in Regillo is misplaced. Regillo reports that both the treatment and control groups lost vision ("[t]he differences between the ranibizumab dose groups and the sham group in mean change in VA [visual acuity] from month three to month 12 were not statistically significant"). Ex. 1026 at 7. Regillo's comparison to sham

Regeneron Exhibit 2001 Page 33 of 35 control ignores the fact that, by this point in time, losing less vision than sham control (no intervention), even if statistically significant, was not considered an effective treatment when MARINA and ANCHOR demonstrated average visual acuity gains with a monthly intravitreal dosing of ranibizumab. Genentech's amendment of the PIER study protocol to allow cross-over from sham, discussed above, reflects the community view that sham (or no intervention) was not an appropriate or ethical comparator by this point in time. Furthermore, Genentech's subsequent protocol amendment to allow all patients in the quarterly dosing arm to roll-over to monthly dosing of ranibizumab, as reported in the PIER Two Year Results, reflects the recognition that PIER quarterly dosing was ineffective as a method of treating an angiogenic eye disorder. Ex. 2016 at 2. Contrary to Dr. Wu's suggestion, an ordinarily skilled retinal specialist would not have understood that quarterly maintenance dosing of ranibizumab reported in Regillo to be an effective treatment regimen. Indeed, Regillo later concludes that "observations from the MARINA and ANCHOR trials suggest that the PIER regimen of dosing every three months after three monthly doses provides less benefit in VA on average than continued monthly dosing." Ex. 1026 at 9.

70. In light of Lucentis's FDA approval and the fact that retina practitioners could now maintain or even improve vision in their wAMD patients, fixed quarterly dosing that produced vision loss was not viewed as an acceptable or

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effective treatment option. In my role as a key opinion leader, academic educator and expert clinician, I am very familiar with how retina specialists are trained and how they practice, particularly as it relates to intravitreal injections of VEGF inhibitors. I am not aware of any retinal specialists who have treated or presently treat their wAMD patients with fixed quarterly dosing of ranibizumab. In other words, I am not aware of any of my peers implementing the PIER regimen (Q4 followed by fixed Q12 dosing of ranibizumab) as a course of treatment for a patient with wet AMD.

I declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believe to be true, and that these statements were made with knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under section 1001 of Title 18 of the United States Code.

Dated: April 14, 2021

Diana Do MD

California

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> **Celltrion Exhibit 1014** Page 1195

UNITED STATES PATENT AND TRADEMARK OFFICE

BEFORE THE PATENT TRIAL AND APPEAL BOARD

CHENGDU KANGHONG BIOTECHNOLOGY CO., LTD. Petitioner

V.

REGENERON PHARMACEUTICALS, INC. Patent Owner

Case PGR2021-00035 Patent 10,828,345

DECLARATION OF DAVID M. BROWN, M.D.

Chengdu Kanghong Biotech. Co., Ltd. v. Regeneron Pharms., Inc., PGR2021-00035, U.S. Pat. 10,828,345, Exhibit 2002

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I, Dr. David Brown, declare as follows:

I. INTRODUCTION

1. I have been retained by counsel for Regeneron Pharmaceuticals, Inc. ("Regeneron") as a technical expert in connection with the above-captioned proceeding. I have been asked to provide my opinions and views on the materials I have reviewed in relation to the Petition for Post Grant review of U.S. Patent No. 10,828,345 (the "345 patent") (Ex. 1001), in particular the state of the art as of the earliest filing date ("priority date") of the '345 patent and responses to the opinion and views of Petitioner's declarant, David Wu, M.D., Ph.D. I submit this declaration in support of Regeneron's Patent Owner Preliminary Response ("POPR").

2. I am being paid at an hourly rate for my work on this matter. I have no personal or financial stake in the outcome of the present proceeding.

II. QUALIFICATIONS AND EXPERIENCE

3. I am the Director of the Greater Houston Retina Research Center, where I have been a Physician Partner and Researcher since 1995. I also have a series of academic appointments: Clinical Professor of Ophthalmology, Cullen Eye Institute at Baylor College of Medicine; Vice-Chair of Ophthalmology for Research and Associate Clinical Professor of Ophthalmology at the Methodist Hospital, Weill Cornell College of Medicine in Houston, Texas; and the NASA-Research and

Regeneron Exhibit 2002 Page 03 of 22 Clinical Advisory Panel-Space Associated Neuro-Ophthalmic Syndrome at NASA Johnson Space Center in Houston, Texas.

4. I graduated from Baylor College of Medicine with highest honors in 1988. I completed a medical/surgical internship at Baylor College of Medicine from 1989-1990. From 1990-1995, I completed ophthalmology and retina training at the University of Iowa where I was a Thomas Heed Fellow, a Hermann Knapp Fellow, and was awarded the Ron Michels Fellowship award presented to the top retinal surgery fellow in the United States in 1994.

5. I have served on the Board of Directors of the American Society of Retina Specialists since 2014; the Macula Society Credentials Committee since 2013; and the Retina Society Finance Committee since 2018. I have served in numerous additional leadership roles in professional organizations and societies in the retina and ophthalmology field over the past three decades. I have also been a peer reviewer for the journals in these fields, including OPHTHALMOLOGY, RETINA, and the New England Journal of Medicine,

6. I maintain an active medical and surgical practice focused on treatment of retinal diseases and have continuously been elected as one of the "Best Doctors in America" 2007-2021 and "Texas Super Docs" from 2009-2021. I am also an elected member of the Macula Society, the Retina Society, and the Club Jules Gonin. My honors include the American Academy of Ophthalmology Honor Award (2000),

Regeneron Exhibit 2002 Page 04 of 22 the American Society of Retina Specialists Honor Award (2008), the ASRS Senior Honor Award (2010), the AAO Senior Honor Award (2014), and Retina Hall of Fame inaugural inductee (2017).

7. My research and clinical trial experience has led to my recognition as an international thought leader on treatments and current standards of care for age related macular degeneration, retinal vein occlusion, and diabetic retinopathy. I have written and published over 400 national meeting presentations, abstracts, and scientific papers including many of the primary papers establishing the safety and efficacy of use of anti-VEGF agents for wet AMD ("wAMD"), retinal vein occlusion, and diabetic retinopathy.

8. I have served as a key investigator on the seminal Phase III clinical trials establishing the efficacy of anti-VEGF agents ranibizumab (Genentech's Lucentis) and aflibercept (Regeneron's Eylea) in wAMD, diabetic macular edema and diabetic retinopathy, and retinal vein occlusions. For example, I was a lead investigator on Genentech's Anti-VEGF Antibody for the Treatment of Predominantly Classic Choroidal Neovascularization in Age-related Macular Degeneration (ANCHOR) Study, the Minimally classic/occult trial of the anti-VEGF antibody ranibizumab in the treatment of neovascular AMD (MARINA) Study, and Regeneron's VEGF Trap-Eye: Investigation of Efficacy and Safety in Wet AMD (VIEW1) Study. My research efforts have contributed to a

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Regeneron Exhibit 2002 Page 05 of 22 transformation in the nature of treatments and outcomes for angiogenic eye disorders. A current copy of my curriculum vitae is filed herewith as Ex. 2023.

III. SUMMARY OF OPINIONS

9. My opinions and views set forth in this declaration are based on my education, training, research, and clinical experience in ophthalmology, specifically in researching and treating retinal diseases, as well as the materials I reviewed in preparing this declaration and the state of scientific knowledge in the art pertaining to the subject matter of the '345 patent at the time of its earliest priority application.

10. In forming my opinions, I have reviewed the following materials: (a) the Petition for Post Grant Review of the '345 patent, PGR2021-00035, including all cited exhibits, (b) all other documents and references herein, and (c) the Patent Owner's Preliminary Response to which my declaration relates.

11. For purposes of preparing this declaration in support of Patent Owner's Preliminary Response, I have been asked to apply Dr. Wu's definition of a person of ordinary skill in the art: a person with a medical doctorate, internship and residency in ophthalmology and either a 1-year medical retina fellowship or a 2-year vitreoretinal surgical fellowship. Ex. 1003 ¶ 82. Likewise, for purposes of preparing this declaration, I have been informed and understand that the earliest filing date of the '345 Patent is January 13, 2011, based on the filing of a Provisional Application on that date.

Regeneron Exhibit 2002 Page 06 of 22 12. It is my opinion that by 2011, the person of ordinary skill in the art, would have understood that a fixed quarterly dosing regimen of ranibizumab, as disclosed in the Shams patent publication and corresponding PIER clinical trial, was a failure and not an effective method of treating an angiogenic eye disorder.

IV. STATE OF THE ART AS OF JANUARY 2011

A. Anti-VEGF Therapies for Angiogenic Eye Disorders

13. Angiogenic eye disorders, characterized by pathologic growth of abnormal blood vessels and vascular leakage from damaged blood vessels in the retina, present significant risks to patients' vision absent treatment. Angiogenic eye disorders, or neovascular eye diseases, include conditions such as iris neovascularization, retinal vein occlusion, diabetic retinopathy, diabetic macular edema, neovascular glaucoma, wAMD, and retinopathy of prematurity. Ex. 2004 at 23. A shared feature of angiogenic eye disorders is the presence of elevated ocular levels of VEGF, a molecule that plays a critical role in the pathology of angiogenesis and vascular permeability.

14. By January 2011, it was recognized by skilled retinal specialists, or the ordinarily skilled artisan, that treating patients with certain anti-VEGF agents, which reduce ocular VEGF levels, could reduce the incidence of pathologic angiogenesis and vascular permeability, and prevent loss of vision and even, in many cases, improve vision. Before the use of anti-VEGF agents, treatments for angiogenic eye

Regeneron Exhibit 2002 Page 07 of 22 disorders included methods such as laser ablation and photodynamic therapy (PDT). These treatments generally did not improve vision in a clinically significant manner and often carried risks of further vision loss through, for example, scarring around the area of the choroidal neovascularization site targeted for treatment.

15. Of the angiogenic eye disorders, wAMD had a particularly poor prognosis, as vision loss in these patients could be sudden, severe, and irreversible. Laser and PDT treatments generally could only slow eventual vision loss. Unlike certain other angiogenic eye disorders such as diabetic macular edema and retinal vein occlusion, wAMD is less forgiving if patients wait too long to receive initial treatment or are not treated at sufficiently regular intervals. With wAMD, irreversible vision loss stems from a combination of retinal pigment epithelium ("RPE") rips, subretinal hemorrhages, and atrophy of the photoreceptors overlying the RPE, as well as fibrosis secondary to long-standing retinal and subretinal edema.

16. Early investigation of anti-VEGF agents to treat wAMD included the use of pegaptanib (Macugen) and investigation of the use of off-label injections of Genentech's VEGF antibody drug bevacizumab (Avastin). The major clinical experimentation that established for the retinal community the efficacy of anti-VEGF therapy, however, came with Genentech's drug, ranibizumab (Lucentis), a VEGF antibody fragment designed to be injected intravitreally into the patient's eye at regular intervals.

Regeneron Exhibit 2002 Page 08 of 22 17. In 2003, Genentech began two large-scale, randomized Phase III clinical trials to test monthly ranibizumab injections in patients with wAMD — MARINA and ANCHOR. I served as a principal investigator for both of these studies and was the first author on the NEJM primary manuscript for ANCHOR, (Brown DM, et al., *Ranibizumab versus verteporfin for neovascular age-related macular degeneration*. N Engl J Med. 2006 Oct 5;355(14):1432-44) (Ex. 2026) and second author on the NEJM primary manuscript for MARINA (Rosenfeld PJ, Brown DM, et al., *Ranibizumab for neovascular age-related macular degeneration*. N Engl J Med. 2006 Oct 5;355(14):1432-44) (Ex. 2026) and second author on the NEJM primary manuscript for MARINA (Rosenfeld PJ, Brown DM, et al., *Ranibizumab for neovascular age-related macular degeneration*. N Engl J Med. 2006 Oct 5;355(14):1419-31) (Ex. 2025).

18. The MARINA trial ran from March of 2003 to December 2005 and enrolled 716 patients with wAMD with either minimally classic or occult choroidal neovascularization. Ex. 2025 at 1. Patients were randomly assigned to received 24 monthly intravitreal injections of Lucentis (either 0.3 mg or 0.5 mg) or sham injections. *Id.* The primary endpoint of the study was the proportion of patients losing fewer than 15 letters from baseline visual acuity at 12 months. *Id.*

19. One year results of the MARINA trial were presented by Genentech in July of 2005. Ex. 2031. The results showed that nearly ninety-five percent of patients treated with Lucentis maintained or improved vision at 12 months. *Id.* The two-year results of the MARINA trial were then published in the New England Journal of Medicine on October 5, 2006. Ex. 2025 at 1. The mean improvements

Regeneron Exhibit 2002 Page 09 of 22 in vision demonstrated in the first 12 months of the study were sustained through the second years of study. *Id.* Mean increases in visual acuity were +6.5 letters in the 0.3 mg group and +7.2 letters in the 0.5 mg groups, compared with a decrease of -10.3 letters in the sham-injection group. *Id.* Impressively, visual acuity improved by 15 or more letters in 24.8% of the 0.3 mg group and 33.8% of the 0.5 mg group. *Id.*

20. The unmasking of the one-year results of the MARINA study prompted discussion with the data and safety monitoring committee, and it was determined in October 2005, 2 months before the end of the patient's final study visit at 24 months, that all patients still in the sham arm could be offered ranibizumab injections. Monthly ranibizumab injections were determined by this point to be a critical tool to not only arrest vision loss in wAMD patients, but to offer the hope for sustained improvements in visual acuity.

21. The MARINA results were supplemented by the outcomes from the ANCHOR trial. The ANCHOR trial ran from May of 2003 to September of 2006 and enrolled 423 patients with predominantly classic choroidal neovascularization in wAMD. Ex. 2026 at 1. Patients were randomized to receive monthly intravitreal Lucentis (0.3 mg or 0.5 mg) plus sham photodynamic verteporfin therapy or monthly sham injections plus active verteporfin therapy. *Id.* As in MARINA, the primary endpoint of the study was also the proportion of patients losing fewer than 15 letters

Regeneron Exhibit 2002 Page 10 of 22 from baseline visual acuity at 12 months. Id.

22. One year results of the ANCHOR trial were presented by Genentech in November of 2005. Ex. 2031. Preliminary one-year data showed that approximately 94 percent of patients treated with 0.3 mg of Lucentis and 96 percent of those treated with 0.5 mg of Lucentis maintained or improved vision compared to approximately 64 percent of those treated with PDT alone. *Id.* The one-year results were published in the New England Journal of Medicine on October 5, 2006, a paper on which I served as the first author. Ex. 2026 at 1. The two-year results of ANCHOR were then published in January 2009 in Ophthalmology. Ex. 2027 at 1. In the ANCHOR study, visual acuity improved from baseline, on average, by +8.1 to +10.7 letters, versus a mean decline of -9.8 letters in the verteporfin photodynamic group. Ex. 2027 at 1. Impressively, visual acuity improved by 15 or more letters in 35.7% of the 0.3-mg group and 40.3% of the 0.5-mg group. *Id.*

23. ANCHOR, while confirming the effectiveness of a monthly intravitreal ranibizumab treatment regimen, also represented a "major breakthrough in the treatment of predominantly classic CNV secondary to AMD" by showing this treatment was "superior to verteporfin PDT" treatment. Ex. 2027 at 7. "The VA benefit from ranibizumab was both rapid and sustained: The superiority of ranibizumab to PDT was evident by 1 month after starting treatment, increased to a plateau by the end of the first year, and then persisted through month 24." *Id.* Like

Regeneron Exhibit 2002 Page 11 of 22 the MARINA study, the positive results demonstrated in the ranibizumab treatment arms resulted in a protocol amendment that allowed patients in the PDT-alone arm of the study to cross over to ranibizumab injections during the latter part of the study. *Id.* at 4.

24. Based on this data from MARINA and ANCHOR, Lucentis received FDA approval for the treatment of wAMD in June 2006. Ex. 1006 at 2. By this point in time, it was well established in the retinal community that standard of care had moved beyond observation and monitoring for wAMD (which was utilized as a sham control) to continuous intravitreal injections of ranibizumab (or off-label Avastin), which were effective methods for improving patients' vision compared to baseline, and often maintained those gains over the course of treatment.

B. Extended Dosing Goals

25. While MARINA and ANCHOR demonstrated powerful breakthroughs in the treatment of wAMD, a persistent goal of the retinal community was to find an effective treatment regimen that required less than monthly visits to an ophthalmologist to treat and/or monitor the progression of wAMD. Intravitreal injections, while generally safe, present the risk of rare but serious adverse events such as endophthalmitis, severe intraocular inflammation, and retinal detachment. Further, monthly visits for injections are costly and burdensome to patients. Even simple monthly monitoring, while reducing risk from IVT injections themselves, is

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burdensome, as patients with wAMD are typically elderly and in-person visits present a challenge for the patient and their caretakers.

26. By 2011, the field continued to investigate extended dosing regimens to treat angiogenic eye disorders. As Jeffrey Heier, M.D., a colleague and coinvestigator in the Regeneron Phase III wAMD trials noted: "Because of the large treatment burden, extensive efforts have been devoted toward developing an optimized treatment paradigm that avoids the need for monthly injections or monitoring visits." Ex. 2021 at 9. Despite these efforts, before 2011, studies showed "fixed quarterly or 'as needed' (*pro re nata* [PRN]) dosing regimens, without requiring monthly monitoring visits, were not effective at maintaining vision." *Id.* at 1.

27. One such study was Genentech's PIER study. The PIER Study ran from August 2004 to March 2007, and was designed to compare three monthly loading doses followed by fixed quarterly dosing of 0.3 mg and 0.5 mg Lucentis against sham control over 24 months in 184 patients. Ex. 1026. at 2. I participated as a clinical investigator and was part of the PIER Study Group, and was involved in the presentation and publication of the Year One data from PIER. As explained below, the PIER Study revealed that fixed quarterly intravitreal injections of ranibizumab over an extended treatment period was not an effective method of treatment.

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28.It was not until the clinical trial results from VIEW I and VIEW II. Regeneron's Phase III trials of aflibercept in wAMD, were released that an anti-VEGF inhibitor demonstrated the ability to provide a safe and highly effective treatment for wAMD on an extended fixed dosing regimen. The VIEW trials compared intravitreal aflibercept 0.5 mg monthly, 2 mg monthly, 2 mg every 2 months after 3 initial monthly doses, and ranibizumab 0.5 mg monthly. Ex. 2021 at 1. The primary endpoint of the VIEW trials was noninferiority (margin of 10%) of the aflibercept regimens to ranibizumab in the proportion of patients maintaining vision at week 52 (losing <15 letters). Id. The one-year results from the VIEW trials demonstrated that intravitreal affibercept dosed either monthly or every two months after the three initial loading doses produced similar efficacy outcomes on average to monthly ranibizumab. Id. This finding of the VIEW trials was viewed with excitement across the retina community. As Heier 2012 noted: "[T]he finding that remarkably similar improvement in vision and anatomic measures can be achieved with less than monthly intravitreal aflibercept injections and without requiring monthly monitoring visits provides an important advance for both patients and their treating physicians." Id. at 10.

V. SHAMS DISCLOSED AN UNSUCCESSFUL 12-WEEK DOSING REGIMEN

29. I have reviewed Shams' disclosure (Ex. 1004) and recognize that the Trial Design (Figure 1) and Treatment Schema (Figure 2) set forth in Shams, as well

Regeneron Exhibit 2002 Page 14 of 22 as its description of a dosing regimen in Example 1, all correspond to Genentech's PIER Study, a clinical trial of ranibizumab (Lucentis®). *Compare* Ex. 1004, Figure 2, *with* Ex. 1026 at 2.

30. Figure 2 of Shams illustrates the administration of the "first individual doses" at months 0, 1, and 2, followed by the "secondary doses" at months 5, 8, 11, and continuing every 3 months through 24 months. Ex. 1004, Figure 2. This corresponds precisely to the study arm in PIER: "The ranibizumab groups received their assigned dose by intravitreal injection every month for three doses (day zero, months one and two), followed by doses every three months (months five, eight, 11, 14, 17, 20, and 23)." Ex. 1026 at 2.

31. As one of the lead clinical investigators on the PIER trial, I received the first read-out of the one-year data from the study from Genentech in early 2006, and was the first to present on this data at the Retinal Physician Symposium held in the Bahamas from May 31-June 3, 2006. The data was highly disappointing to say the least. While patients saw an initial gain in visual acuity during the three monthly loading doses, these gains were entirely lost once quarterly dosing began. Ex. 1026 at 7 ("On average, there was 4.5-letter decline in VA between month three and month 12 for both ranibizumab dose groups."). By month 12, the 0.3 mg study arm saw a -1.6 letter difference from baseline visual acuity, and the 0.5 mg study arm saw a -0.2 letter difference from baseline. Ex. 1026 at 7 (Figure 2).

Regeneron Exhibit 2002 Page 15 of 22 32. OCT-assessed anatomic outcomes in the Year One data also confirmed ranibizumab's failure to maintain efficacy over the quarterly dosing period. The maximal decrease in foveal center point thickness was seen at months two and three for both ranibizumab groups. Ex. 1026 at 7. During assessments made at months five and eight, the foveal center point thickness was on average greater than at months two and three, and was also greater than at month 12, which had followed a ranibizumab dose at month 11. *Id.* This suggested that, on average, ranibizumab's therapeutic effectiveness in a patient would wane between injections, pointing to recurrent neovascular activity and associated exudation occurring between injections. Ex. 2018 at 4.

33. Over the course of the PIER study, the study sponsor (Genentech) implemented two key protocol amendments. First, the protocol was amended on February 27, 2006 to provide sham injection patients the opportunity to cross over to receive 0.5 mg ranibizumab quarterly after completing the month-12 visit (*i.e.*, the assessment time point for the primary analysis). Ex. 2016 at 2. As explained *supra*, the 12-month data from MARINA and ANCHOR had established to the retinal community that it would be in the best interest of the sham groups of patients to be treated with ranibizumab, rather than be put further at risk for severe, irreversible vision loss under an observation-only scheme.

34. The second protocol amendment was the direct result of the review of

Regeneron Exhibit 2002 Page 16 of 22 the 12-month PIER data. On August 21, 2006, the study was amended to provide all patients remaining in the study the opportunity to roll over to receive 0.5 mg ranibizumab monthly for the remainder of the study. Ex. 2016 at 2. The second year of the PIER study was functionally designed to be confirmatory of fixed quarterly dosing's efficacy, and given the lack of efficacy observed in the PIER quarterly treatment regimen, the study sponsor initiated a rollover at this point to mitigate against future visual acuity losses, which would be expected on continued quarterly dosing.

35. The year-two results from PIER confirmed the lack of efficacy of the dosing regimen. The 0.3 mg ranibizumab group saw a mean loss of -2.2 letters as compared to baseline, and the 0.5 mg ranibizumab arm saw a mean loss of -2.3 letters as compared to baseline. Ex. 2016 at 2. This stood in stark contrast to MARINA and ANCHOR. *Id.* at 8 ("In those studies, patients who received monthly injections of ranibizumab experienced a gain of 5 to 11 letters from baseline at month 24 compared to a loss of approximately 2 letters with the PIER dosing regimen."). After approval of Lucentis, and certainly by 2011, the goal of any treatment regimen for age-related macular degeneration was to improve vision and prevent blindness.

36. Post-hoc analyses of the study data from MARINA, ANCHOR, and PIER also demonstrated that while patients on monthly ranibizumab were significantly less likely to develop macular hemorrhages as compared to sham control, patients in the treatment arm of PIER saw no benefit in the incidence of macular hemorrhage as compared to sham control and, indeed, incidence rates were numerically higher. Ex. 2020 at 3, 7. Macular hemorrhages are a hallmark of wAMD and are considered to be a definitive sign of disease progression. "[W]hen occupying larger areas or located in the subfoveal region, they are usually associated with a poor visual prognosis in a majority of cases." *Id.* at 1. It was a serious concern, therefore, that quarterly dosing did not even decrease the incidence of macular hemorrhage as compared to sham and it was recognized that "switching from monthly to quarterly injection intervals may not have the same beneficial effect and could put the patient at an increased risk for vision threatening complications." *Id.* at 9.

37. As a result, my conclusion from the PIER Study results was that "we cannot just mandatorily treat on a quarterly basis and maintain the visual gains seen with the first three monthly injections." Ex. 2017 at 1; *id.* at 2 ("You can't just do mandatory quarterly injections.") My expressed concerns with fixed quarterly injections were shared across the retina community at the time. *E.g.*, Ex. 2018 at 5 ("A recent analysis of the ANCHOR, MARINA, and PIER data demonstrated that monthly intravitreal ranibizumab dosing significantly reduced the frequency of macular hemorrhages compared with the sham controls or photodynamic therapy-treated patients regardless of lesion type. The effect was lost when patients were

switched from monthly to quarterly dosing in the PIER study. Reducing the frequency of injections should, therefore, be done with caution."); Ex. 2015 ("The PIER data have led Genentech to recommend that patients receive either monthly injections of ranibizumab, or have their retreatment schedules determined through individualized testing.")

38. These concerns were also reflected in the FDA's labeling when Lucentis was approved for wAMD treatment in June 2006, as PIER's year-one results were included in the FDA's review. Ex. 1006 at 2. The label explains: "Although less effective, treatment may be reduced to one injection every three months after the first four injections if monthly injections are not feasible. Compared to continued monthly dosing, dosing every 3 months will lead to an approximate 5-letter (1-line) loss of visual acuity benefit, on average, over the following 9 months. Patients should be evaluated regularly." Ex. 2028.

39. In my opinion, FDA's review and consideration of the PIER results, and this language in the label, would not suggest to the person of ordinary skill that quarterly maintenance dosing was an effective or acceptable option for a treatment regimen for a patient with an angiogenic eye disorder, such as wAMD.

40. I disagree with Dr. Wu's suggestion in paragraph 111 that this language in the FDA label would indicate that the Shams/PIER regimen would have been a longer than monthly dosing regimen utilized by those in the art. Ex. 1003, ¶ 111.

Indeed, Dr. Wu does not address the label's explicit note that this is a "less effective" regimen than fixed monthly, and that it "will lead" to a "loss of visual acuity benefit." Ex. 2028 at 2. Rather, it is my understanding that FDA would present a comprehensive look at the clinical data it is presented with when issuing approval guidelines, and indeed the clinical studies section of the Lucentis label (at §§ 14.1 and 14.2) plainly shows the drastic disparity between the study results from ANCHOR and MARINA and those of PIER. *Id.* at 7. Figure 1 shows that Lucentis 0.5 mg arm in Study 1 (MARINA) had mean changes in visual acuity of +6.6 letters at 24 months, and Lucentis 0.5 mg arm in Study 2 (ANCHOR) had +11.3 letters mean changes at 24 months, while Figure 2 shows Lucentis 0.5 mg arm in Study 3 (PIER) had a mean change of -0.2 letters at 12 months. *Id.* at 7. The import of these Figures would be clear to retina practitioners: the fixed quarterly dosing regimen of PIER carriers a high risk for permanent vision loss.

41. I also disagree with Dr. Wu's suggestion in paragraph 63 that the EXCITE study "was consistent with the findings in PIER in that both monthly and quarterly dosing of ranibizumab was able to improve vision of wet AMD patients." Ex. 1003 ¶ 63. As an initial matter, it is incorrect to suggest that the PIER study found quarterly dosing to "improve vision of wet AMD patients." A finding of superiority to sham is not equated with improved vision; in the context of the PIER Study, it simply meant that patients in the ranibizumab treatment arms lost less

vision at the 12 and 24 month endpoints as compared to baseline vision than sham. True vision improvement for wAMD patients occurred in ANCHOR and MARINA, where there were significant visual acuity gains compared to baseline at 12 and 24 months.

42. Further, Dr. Wu fails to address the fact that the objective of the EXCITE study, conducted from December 2005 to January 2008, was to demonstrate the "noninferiority of a quarterly treatment regimen to a monthly regimen of ranibizumab in patients" with subfoveal CNV secondary to wAMD; and that "noninferiority of a quarterly regimen was not achieved with reference to 5.0 letters." Ex. 1027 at 1. In other words, this study was a failure by its own terms. Like PIER, the treatment arms of EXCITE were administered as three monthly loading doses prior to fixed quarterly doses of ranibizumab, and also like PIER, EXCITE demonstrated losses of initial visual acuity gains after patients moved to quarterly injections. *Id.* at 5. The study group concluded: "The direct comparative analysis between monthly and quarterly treatment regimens of the EXCITE study is consistent with the clinical guidance on ranibizumab treatment, which recommends rigorous monthly monitoring with timely retreatment of patients." *Id.* at 8.

43. In my role as a key opinion leader in the treatments for angiogenic eye disorders, as well as an active clinician and clinical instructor, I am very familiar

Regeneron Exhibit 2002 Page 21 of 22 with the standards for how retina specialists are trained, best clinical practices, and how in fact retina doctors treat patients. As of 2011, following the disclosure of the PIER results and the post hoc analyses presented in peer reviewed publications, due to the dismal one-year visual acuity results and the unacceptable risks of permanent vision loss, retinal specialists would not have implemented the PIER dosing regimen (Q4 followed by fixed Q12 dosing of ranibizumab) as a course of treatment for a patient with an angiogenic eye disorder. After the results of PIER were presented and published, I am not aware of any retina specialist who recommends or has treated patients on a fixed q4/q12 quarterly dosing regimen of ranibizumab, and I believe from a medical malpractice standpoint, that such treatment would be inconsistent with the community standard of care.

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements are made with the knowledge that willful false statements and the like are punishable by fine, imprisonment, or both under Section 1001 of Title 18 of the United States Code.

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Dated <u>April 14</u>, 25 2.1

David M. Brown, M.D. , ľ cxas

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VEGF-TRAP_{R1R2} Suppresses Choroidal Neovascularization and VEGF-Induced Breakdown of the Blood-Retinal Barrier

YOSHITSUGU SAISHIN,¹ YUMIKO SAISHIN,¹ KYOICHI TAKAHASHI,¹ RAQUEL LIMA E SILVA,¹ DONNA HYLTON,² JOHN S. RUDGE,² STANLEY J. WIEGAND,² AND PETER A. CAMPOCHIARO¹*

¹ The Departments of Ophthalmology and Neuroscience, The Johns Hopkins University School of Medicine, Maumenee, Baltimore, Maryland ²Regeneron Pharmaceuticals, Tarrytown, New York, New York

Vascular endothelial growth factor (VEGF) plays a central role in the development of retinal neovascularization and diabetic macular edema. There is also evidence suggesting that VEGF is an important stimulator for choroidal neovascularization. In this study, we investigated the effect of a specific inhibitor of VEGF, VEGF-TRAP_{R1R2}, in models for these disease processes. VEGF-TRAP_{R1R2} is a fusion protein, which combines ligand binding elements taken from the extracellular domains of VEGF receptors 1 and 2 fused to the Fc portion of IgG1. Subcutaneous injections or a single intravitreous injection of VEGF-TRAP_{R1R2} strongly suppressed choroidal neovascularization in mice with laser-induced rupture of Bruch's membrane. Subcutaneous injection of VEGF-TRAP_{R1R2} also significantly inhibited subretinal neovascularization in transgenic mice that express VEGF in photoreceptors. In two models of VEGF-induced breakdown of the blood-retinal barrier (BRB), one in which recombinant VEGF is injected into the vitreous cavity and one in which VEGF expression is induced in the retina in transgenic mice, VEGF-TRAP_{R1R2} significantly reduced breakdown of the BRB. These data confirm that VEGF is a critical stimulus for the development of choroidal neovascularization and indicate that VEGF-TRAP_{R1R2} may provide a new agent for consideration for treatment of patients with choroidal neovascularization and diabetic macular edema. J. Cell. Physiol. 195: 241-248, 2003. © 2003 Wiley-Liss, Inc.

Ocular neovascularization, consisting of retinal and choroidal neovascularization, is an enormous public health problem. Retinal neovascularization occurs in ischemic retinopathies, the most prevalent of which is diabetic retinopathy, the most common cause of severe vision loss in young people in developed countries (Klein et al., 1984). Choroidal neovascularization complicates several diseases in which there are abnormalities of the Bruch's membrane/retinal pigmented epithelial (RPE) cell complex, such as age-related macular degeneration (AMD), the most common cause of severe vision loss in the elderly (The Macular Photocoagulation Study Group, 1991). While retinal and choroidal neovascularization are responsible for the vast majority of severe vision loss in Americans, diabetic macular edema is the major cause of moderate vision loss (Klein et al., 1984).

Multiple stimulatory factors may contribute to the development of retinal neovascularization, but vascular endothelial growth factor (VEGF) plays a critical role. Signaling through VEGF receptors is both necessary and sufficient for development of retinal neovascularization (Okamoto et al., 1997; Seo et al., 1999; Ozaki et al., 2000). VEGF also causes breakdown of the bloodretinal barrier (BRB) (Ozaki et al., 1997), and has been implicated in the early breakdown of the BRB that occurs in diabetes (Qaum et al., 2001). In addition,

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VEGF is also an important stimulus for choroidal neovascularization (Kwak et al., 2000). Therefore, antagonizing VEGF is a potentially useful strategy for several ocular diseases.

Many approaches for antagonizing VEGF are being considered. One strategy is to inject relatively large inhibitors, such as aptamers or FAb fragments of anti-VEGF antibodies directly into the eye. Phase I clinical

PAC is the George S. and Dolores Dore Eccles Professor of Ophthalmology and Neuroscience.

Contract grant sponsor: Public Health Service; Contract grant numbers: EY05951, EY12609, P30EY1765; Contract grant sponsor: Foundation Fighting Blindness; Contract grant sponsor: Research to Prevent Blindness (Lew R. Wasserman Merit Awards and unrestricted grant); Contract grant sponsor: Dr. and Mrs. William Lake.

*Correspondence to: Peter A. Campochiaro, Maumenee 719, The Johns Hopkins University School of Medicine, 600 N. Wolfe Street, Baltimore, MD 21287-9277. E-mail: pcampo@jhmi.edu

Received 30 August 2002; Accepted 20 November 2002 DOI: 10.1002/jcp.10246

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trials testing the safety and tolerability of this approach have been completed and phase II and III trials are planned or in progress. Preliminary reports suggest that inflammation may occur following intraocular injection of antibodies or aptamers, but this has not been a severe enough problem to discontinue evaluation of these approaches (Guyer et al., 2001; Schwartz et al., 2001). This approach has some concerns, because repeated intraocular injections carry risks of retinal detachment and endophthalmitis, and may not be feasible depending upon the frequency of injections required. Another strategy is to avoid repeated intraocular injections by systemic administration of small molecule VEGF antagonists (Seo et al., 1999; Kwak et al., 2000; Ozaki et al., 2000). There is a theoretical concern that some beneficial types of angiogenesis, such as collateral formation in ischemic myocardium, may be inhibited. But there are no data to support this concern and it is equally plausible that systemic inhibition of VEGF could have many additional benefits, since angiogenesis has been implicated in tumor growth, atherosclerosis, and arthritis (for review, see Folkman, 1995). Oral administration of VEGF receptor kinase inhibitors results in dramatic suppression of retinal and choroidal neovascularization and is a very promising approach (Seo et al., 1999; Kwak et al., 2000; Ozaki et al., 2000). These agents are selective, but not specific VEGF antagonists, be-cause it is difficult to inhibit VEGF receptor kinases without inhibiting homologous kinases such as plateletderived growth factor (PDGF) receptor kinase and c-kit, the receptor for stem cell factor (Fabbro et al., 1999; Bold et al., 2000; Drevs et al., 2000; Wood et al., 2000). The effects of these additional activities are unknown and while they are being investigated, it is prudent to consider and test more selective VEGF inhibitors

Soluble VEGF receptors provide a very specific way to antagonize VEGF, and several studies have demonstrated that the extracellular domain of VEGF receptor 1 (VEGF-R1) has antiangiogenic activity (Goldman et al., 1998; Kong et al., 1998; Honda et al., 2000; Shiose et al., 2000; Takayama et al., 2000; Lai et al., 2001; Mahasreshti et al., 2001; Bainbridge et al., 2002; Lai et al., 2002). A disadvantage of soluble VEGF-R1 is that it is cleared fairly rapidly. Pharmacokinetic properties can be improved by linking the ligand binding domains of VEGF receptors to the Fc portion of IgG, which slows clearance by conferring the long circulating half-life of an antibody to the chimeric molecule. A potential trade off is that the relatively large size of such constructs could limit tissue penetration from the systemic circulation, which is a particularly important consideration for treatment of ocular diseases. In this study, we have evaluated both local and systemic administration of a novel chimeric molecule, VEGF- $TRAP_{R1R2}$, which comprises portions of the extracellular domain of VEGFR-1 (flt-1) and VEGFR-2 (KDR), in models of ocular neovascularization and breakdown of the BRB.

$\begin{array}{c} \text{MATERIALS AND METHODS} \\ \text{VEGF-TRAP}_{\text{R1R2}} \end{array}$

 $VEGF\text{-}TRAP_{R1R2}$ (Regeneron Pharmaceuticals, Tarrytown, NY) is a recombinant fusion protein that contains Ig domain 2 of VEGF-R1 and Ig domain 3 of VEGF-R2 fused to the Fc portion of human IgG1

(Wulff et al., 2002). VEGF-TRAP_{R1R2} binds VEGF with high affinity (kD \approx 1 pM) and subcutaneous injection of 25 mg/kg of VEGF-TRAP_{R1R2} has been shown to effectively neutralize VEGF in mice with VEGF-secreting tumors (Wong et al., 2001). Recombinant human Fc was used as a control protein.

Treatment of mice with laser-induced choroidal neovascularization

Choroidal neovascularization was generated by modification of a previously described technique (Tobe et al., 1998b). Briefly, 4-5-week-old female C57BL/6J mice were anesthetized with ketamine hydrochloride (100 mg/kg body weight) and the pupils were dilated with 1% tropicamide. Three burns of 532 nm diode laser photocoagulation (75 µm spot size, 0.1 sec duration, 120 mW) were delivered to each retina using the slit lamp delivery system of an OcuLight GL Photocoagulator (Iridex, Mountain View, CA) and a hand held cover slide as a contact lens. Burns were performed in the 9, 12, and 3 o'clock positions of the posterior pole of the retina. Production of a bubble at the time of laser, which indicates rupture of Bruch's membrane, is an important factor in obtaining CNV (Tobe et al., 1998b), so only burns in which a bubble was produced were included in the study. Mice were treated with subcutaneous injections of 25 mg/kg of VEGF-TRAP_{R1R2} or Fc fragment 1 day prior to laser and on days 2, 5, 8, and 11 after laser. At 14 days after laser, the mice were euthanized, serum was collected and stored, and eyes were rapidly dissected for choroidal flat mounts or frozen in optimum cutting temperature embedding compound (OCT; Miles Diagnostics, Elkhart, IN).

Some mice were given intraocular injection of $4.92~\mu g$ of VEGF-TRAP_{R1R2} in one eye and $4.92~\mu g$ Fc fragment in the other eye. Two weeks later, mice were perfused with fluorescein-labeled dextran and choroidal neovas-cularization was measured.

Quantitative analysis of the amount of choroidal neovascularization

The sizes of CNV lesions were measured in choroidal flat mounts (Edelman and Castro, 2000) by an investigator masked with respect to treatment group. Mice used for the flat mount technique were anesthetized and perfused with 1 ml of phosphate-buffered saline containing 50 mg/ml of fluorescein-labeled dextran $(2\times 10^6$ average mw, Sigma, St. Louis, MO) as previously described (Tobe et al., 1998a). The eyes were removed and fixed for 1 h in 10% phosphate-buffered formalin. The cornea and lens were removed and the entire retina was carefully dissected from the eyecup. Radial cuts (4-7), average 5) were made from the edge to the equator and the eyecup was flat mounted in Aquamount with the sclera facing down. Flat mounts were examined by fluorescence microscopy on an Axioskop microscope (Zeiss, Thornwood, NY) and images were digitized using a 3 color CCD video camera (IK-TU40A, Toshiba, Tokyo, Japan) and a frame grabber. Image-Pro Plus software (Media Cybernetics, Silver Spring, MD) was used to measure the total area of choroidal neovascularization associated with each burn with the operator masked with respect to treatment group. Statistical comparisons were made between the size of lesions in mice treated with VEGF-TRAP_{R1R2} versus those in mice treated with Fc fragment by two-tailed *t*-test. In addition, the average size of choroidal neovascularization in each mouse was calculated and plotted against the serum level of VEGF-TRAP_{R1R2} obtained by ELISA.

In some mice, the eyes were rapidly removed and frozen in optimum cutting temperature embedding compound (OCT; Miles Diagnostics). Ten µm frozen sections were cut through entire lesions and the sections were histochemically stained with biotinylated Griffonia simplicifolia lectin B4 (GSA, Vector Laboratories, Burlingame, CA), which selectively binds to vascular cells. Slides were incubated in methanol/ H_2O_2 for 10 min at 4°C, washed with 0.05 M Tris-buffered saline, pH 7.6 (TBS), and incubated for 30 min in 10% normal porcine serum. Slides were incubated 2 h at room temperature with biotinylated GSA and after rinsing with 0.05 M TBS, they were incubated with avidin coupled to peroxidase (Vector Laboratories) for 45 min at room temperature. The slides were developed with Histo-Mark Red (Kirkegaard and Perry, Cabin John, MD) to give a red reaction product and counter stained with Contrast Blue (Kirkegaard and Perry).

Transgenic mice with increased expression of VEGF in photoreceptors

Transgenic mice with VEGF driven by the rhodopsin promoter develop subretinal neovascularization due to expression of VEGF in photoreceptors beginning at about P7 (Okamoto et al., 1997). Hemizygous transgenepositive mice were given a subcutaneous injection of 25 mg/kg of VEGF-TRAP_{R1R2} or Fc fragment at P7, P10, P13, P16, and P19. At P21, the mice were sacrificed and the amount of subretinal neovascularization was quantified as previously described (Tobe et al., 1998a). Briefly, mice were anesthetized and perfused with 1 ml of phosphate-buffered saline containing 50 mg/ml of fluorescein-labeled dextran $(2 \times 10^6$ average mw, Sigma). The eyes were removed and fixed for 1 h in 10% phosphate-buffered formalin. The cornea and lens were removed and the entire retina was carefully dissected from the eyecup, radially cut from the edge of the retina to the equator in all 4 quadrants, and flat-mounted in Aquamount with photoreceptors facing upward. The retinas were examined by fluorescence microscopy at 200x magnification, which provides a narrow depth of field so that when focusing on neovascularization on the outer surface of the retina, the remainder of the retinal vessels are out-of-focus allowing easy delineation of the neovascularization. The outer edge of the retina, which corresponds to the subretinal space in vivo, is easily identified and therefore there is standardization of focal plane from slide to slide. Images were digitized using a 3 CCD color video camera and a frame grabber. Using Image-Pro Plus software, an investigator masked with respect to treatment group delineated each of the lesions and calculated the total area of neovascularization per retina as previously described (Tobe et al., 1998a).

VEGF-induced breakdown of the BRB

Adult C57BL/6 mice were given a subcutaneous injection of 25 mg/kg of VEGF-TRAP_{\rm R1R2} or Fc and on the following day VEGF-induced breakdown of the BRB

was quantified as previously reported (Derevjanik et al., 2002). Mice were anesthetized with 25 mg/kg of ketamine and 4 mg/kg of xylazine, pupils were dilated with 1% tropicamide. Intraocular injections were performed with a Harvard pump microinjection apparatus and pulled glass micropipets (Mori et al., 2001). Each micropipet was calibrated to deliver 1 µl of fluid upon depression of a foot switch. Under a dissecting microscope, the sharpened tip of a micropipet was passed through the sclera just behind the limbus into the vitreous cavity, and the foot switch was depressed injecting 1 µl of 10^{-6} M human vascular endothelial growth factor (VEGF; R&D Systems, Minneapolis, MN). Six hours later, retinal vascular permeability was measured using [³H]mannitol as a tracer.

Double transgenic rho/rtTA-TRE/VEGF mice with doxycycline-inducible expression of VEGF in photo-receptors (Ohno-Matsui et al., 2002) were also used. Double transgenics were given a subcutaneous injection of 25 mg/kg of VEGF-TRAP_{R1R2} or Fc fragment of IgG and on the following day they were started on 2 mg/ml of doxycycline in their drinking water. The next day they were given a second subcutaneous injection of 25 mg/kg of VEGF-TRAP_{R1R2} or Fc fragment and after two days, retinal vascular permeability was measured.

Measurement of BRB breakdown using [³H]mannitol as tracer

Six hours after intraocular injection of VEGF in wild type mice or 2 days after rho/rtTA-TRE/VEGF were started on doxycycline, mice were given an intraperitoneal injection of 1 µCi/gram body weight of [³H]mannitol (New England Nuclear, Boston, MA). After 1 h, mice were sacrificed and eyes were removed. The cornea and lens were removed and the entire retina was carefully dissected from the eyecup and placed within pre-weighed scintillation vials. The thoracic cavity was opened and the left superior lobe of the lung was removed and placed in another pre-weighed scintillation vial. All liquid was removed from the vials and remaining droplets were allowed to evaporate over 20 min. The vials were weighed and the tissue weights were recorded. One ml of NCSII solubilizing solution (Amersham, Chicago, IL) was added to each vial and the vials were incubated overnight in a 50°C water bath. The solubilized tissue was brought to room temperature and decolorized with 20% benzoyl peroxide in toluene in a 50°C water bath. The vials were brought to room temperature and 5 ml of Cytoscint ES (ICN, Aurora, OH) and 30 µl of glacial acetic acid were added. The vials were stored for several hours in darkness at 4°C to eliminate chemoluminescence. Radioactivity was counted with a Wallac 1409 Liquid Scintillation Counter (Gaithersburg, MD).

RESULTS Subcutaneous injection of VEGF-TRAP_{R1R2} inhibits choroidal neovascularization

Bruch's membrane was ruptured at 3 locations in each eye by laser photocoagulation in C57BL/6 mice. One day prior to laser and on days 2, 5, 8, and 11 after laser, mice received subcutaneous injection of 25 mg/kg of VEGF-TRAP_{R1R2} or Fc fragment. Retinal whole mounts from fluorescein dextran-perfused mice treated with VEGF- 244

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 $TRAP_{R1R2}$ (Fig. 1A,B) had areas of neovascularization that were much smaller than those seen in mice treated with Fc fragment (Fig. 1C,D). Sections through Bruch's membrane rupture sites in other mice treated with VEGF-TRAP_{R1R2} showed complete or near-complete inhibition of choroidal neovascularization (Fig. 1E,F). Mice treated with Fc fragment (Fig. 1G,H) had choroidal neovascularization similar to that seen in mice treated with vehicle in several other studies (Seo et al., 1999; Kwak et al., 2000). Measurement of the area of choroidal neovascularization by image analysis confirmed that there was significantly less neovascularization in eyes treated with VEGF-TRAP_{R1R2} compared to those treated with Fc fragment (Fig. 1I). The level of VEGF-

 $\rm TRAP_{R1R2}$ was measured in plasma obtained from each of the mice at the time of sacrifice. Each of the mice that had been injected with Fc fragment had no detectable VEGF-TRAP_{R1R2} in its plasma, while mice that had been injected with VEGF-TRAP_{R1R2} had plasma levels ranging from 57 to 205 $\mu g/ml$. All of the plasma levels of VEGF-TRAP_{R1R2} between 57 and 205 $\mu g/ml$ were associated with strong inhibition of choroidal neovascularization (Fig. 1J).

Immediately after laser, some mice were given intraocular injection of VEGF-TRAP_{R1R2} or Fc fragment of IgG. Two weeks later, mice were perfused with fluorescein-labeled dextran and choroidal neovascularization was measured. Mice that received intraocular

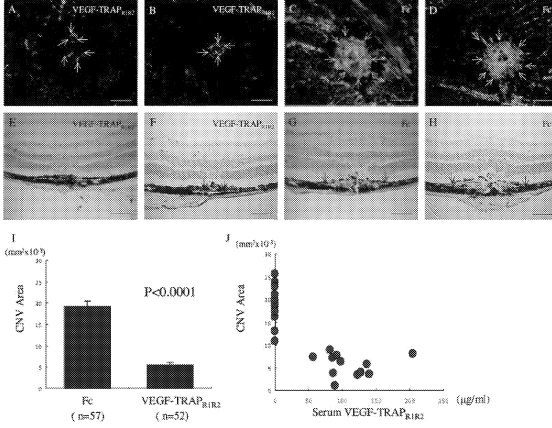


Fig. 1. Subcutaneous VEGF-TRAP_{R1R2} suppresses choroidal neovascularization at sites of rupture of Bruch's membrane. Adult C57BL/6 mice were had rupture of Bruch's membrane by laser photocoagulation in 3 locations in each eye. Prior to laser and on days 2, 5, 8, and 11 after laser, mice received subcutaneous injection of 25 mg/kg of VEGF-TRAP_{R1R2} or Fc fragment of IgG. Parts **A** and **B** show small areas of neovascularization (surrounded by arrows) in retinal whole mounts from two fluorescein dextran-perfused mice treated with VEGF-TRAP_{R1R2}. *Griffonia simplicifolia* (GSA) lectin-stained sections in two other mice treated with VEGF-TRAP_{R1R2} show minimal choroidal neovascularization (**E**- none visible and **F**- between arrows). Parts **C** and **D** show large areas of neovascularization (surrounded by

arrows) in choroidal flat mounts from two Fc fragment-treated mice and GSA-stained sections from two other mice treated with Fc fragment show prominent areas of neovascularization (G and H, between arrows). Measurement by image analysis of the area of neovascularization on choroidal flat mounts (I) showed an average area that was significantly smaller (P < 0.0001 by Student's two-tailed t-test) in VEGF-TRAP_RIR2-treated mice (20 eyes, 52 rupture sites) compared to Fc-treated mice (20 eyes, 57 rupture sites). Plasma levels of VEGF at the time of sacrifice determined by ELISA plotted against the average area of choroidal neovascularization per mouse showed marked suppression of neovascularization at all plasma levels between 50 and 200 $\mu g/ml$ (J). Bar = 100 μm .

VEGF-TRAP AND OCULAR NV

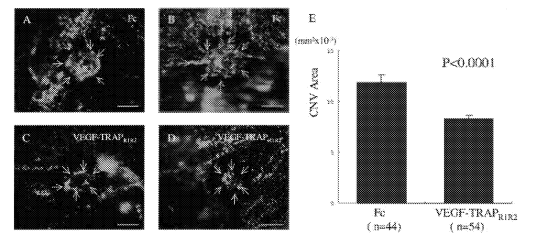


Fig. 2. A single intravitreous injection of VEGF-TRAP_{R1R2} suppresses choroidal neovascularization at Bruch's membrane rupture sites. Immediately after laser, C57BL/6 mice were given intraocular injection of 4.92 μ g of VEGF-TRAP_{R1R2} in one eye and 4.92 μ g of Fc fragment in the other eye. Two weeks later, mice were perfused with fluorescein-labeled dextran and choroidal neovascularization (surrounded by arrows) are seen in flat mounts from two separate mice treated with

intravitreous injection of Fc fragment. C and D: Small areas of neovascularization (surrounded by arrows) are seen in two separate mice given a single intravitreous injection of VEGF-TRAP_{R1R2}. E: The area of choroidal neovascularization measured by image analysis was significantly less (P < 0.0001; Student's two-tailed *t*-test) in VEGF-TRAP_{R1R2}-treated eyes (19 eyes, 54 rupture sites) compared to Fc-treated eyes (19 eyes, 44 rupture sites). Bar = 100 μ m

injection of Fc fragment had larger areas of choroidal neovascularization (Fig. 2A,B) than those seen in mice that received a single intraocular injection of VEGF-TRAP_{R1R2} (Fig. 2C,D). There was a statistically significant reduction in the mean area of neovascularization in VEGF-TRAP_{R1R2}-injected eyes compared to Fc fragment-injected eyes (Fig. 2E).

VEGF-TRAP_{R1R2} inhibits subretinal neovascularization in Rho/VEGF transgenic mice

Rho/VEGF transgenic mice express VEGF in photoreceptors starting about postnatal day (P) 7 resulting in extensive subretinal neovascularization by P21

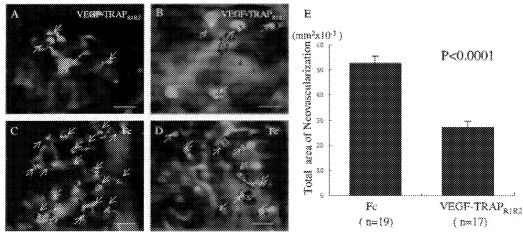


Fig. 3. Subcutaneous VEGF-TRAP_{R1R2} inhibits subretinal neovascularization in rho/VEGF transgenic mice. Rho/VEGF transgenic mice begin to express VEGF in photoreceptors about postnatal day (P) 7. At P7, mice were divided into two groups and treated with 25 mg/kg of VEGF-TRAP_{R1R2} (9 mice, 17 eyes) or Fc fragment (10 mice, 19 eyes) on P7, P10, P13, P16, and P19, and on P21, the mice were anesthetized and perfused with fluorescein-labeled dextran. Retinal whole mounts

from mice treated with VEGF-TRAP_{R1R2} showed few areas of neovascularization (**A** and **B**, arrows), while there were numerous clumps of new vessels in the subretinal space of mice that had been treated with Fc fragment (**C** and **D**, arrows). Measurement of the total area of neovascularization per retina by image analysis showed significantly less neovascularization in VEGF-TRAP_{R1R2}-treated mice, compared to those treated with Fc fragment (**E**). Bar = 100 μ m.

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(Okamoto et al., 1997; Tobe et al., 1998a). Rho/VEGF mice received subcutaneous injection of 25 mg/kg of VEGF-TRAP_{R1R2} or Fc fragment of IgG on P7, P10, P13, P16, and P19, and on P21, they were perfused with fluorescein-labeled dextran. Mice treated with VEGF-TRAP_{R1R2} had very few clumps of neovascularization (Fig. 3A,B, arrows), while there were numerous clumps of new vessels in the subretinal space of mice that had been treated with Fc fragment of IgG (Fig. 3C,D, arrows). Image analysis showed that mice treated with VEGF-TRAP_{R1R2} had an average area of neovascularization per retina that was significantly smaller total area than mice treated with Fc fragment (Fig. 3E).

VEGF-TRAP_{R1R2} inhibits VEGF-induced breakdown of the BRB

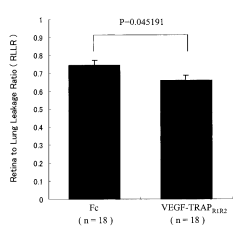
Adult C57BL/6 mice were given a subcutaneous injection of 25 mg/kg of VEGF-TRAP_{R1R2} or Fc fragment and on the following day received an intravitreous injection of 1 µg of 10^{-6} M VEGF. Six hours later, retinal vascular permeability was measured using [³H]mannitol as a tracer. Mice treated with VEGF-TRAP_{R1R2} had a significantly smaller retina to lung leakage ratio than mice treated with Fc fragment of IgG indicating less breakdown of the BRB (Fig. 4A).

We have previously produced and characterized double transgenic mice with doxycycline-inducible expression of VEGF in the retina (Ohno-Matsui et al., 2002). Double transgenics were given a subcutaneous injection of 25 mg/kg of VEGF-TRAP_{R1R2} or Fc fragment and on the following day they were started on 2 mg/ml of doxycycline in their drinking water. Two days later, they were given a second subcutaneous injection of 25 mg/kg of VEGF-TRAP_{R1R2} or Fc fragment and then the next day retinal vascular permeability was measured with [³H]mannitol. Double transgenic mice treated with VEGF-TRAP_{R1R2} had a significant reduction in the retina to lung leakage ratio compared to mice treated with Fc fragment (Fig. 4B).

DISCUSSION

Retinal ischemia is the underlying cause of retinal neovascularization. Since VEGF and VEGFR1 are upregulated in ischemic tissue (Forsythe et al., 1996; Gerber et al., 1997; Iyer et al., 1998), it is not surprising that VEGF plays a central role in the pathogenesis of retinal neovascularization. The pathogenesis of choroidal neovascularization is poorly understood. Choroidal blood flow is decreased in patients with AMD (Grunwald et al., 1998; Ross and Barofsky, 1998), but it is not known if this is sufficient to cause hypoxia. Also, it is unlikely that hypoxia is present in other disease processes, such as ocular histoplasmosis or degenerative myopia, in which choroidal neovascularization occurs in young patients. Since ischemia has not been implicated in the pathogenesis of choroidal neovascularization, this piece of evidence that made VEGF a prime suspect for retinal neovascularization is lacking for choroidal neovascularization. On the other hand, surgically removed choroidal neovascular membranes show immunohistochemical staining for VEGF (Amin et al., 1994; Frank et al., 1996; Kvanta et al., 1996; Lopez et al., 1996) and there is increased VEGF mRNA in experimentally induced choroidal neovascularization (Ogata

A. Exogenous VEGF-Induced BRB Breakdown



B. Endogenous VEGF-Induced BRB Breakdown

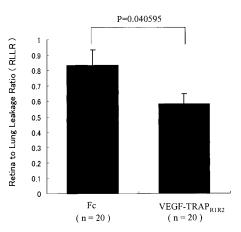


Fig. 4. Subcutaneous injections of VEGF-TRAP_{R1R2} suppress VEGF-induced breakdown of the BRB. Adult C57BL/6 mice were given a subcutaneous injection of 25 mg/kg of VEGF-TRAP_{R1R2} or Fc fragment and on the following day received an intravitreous injection of 1 µg of 10^{-6} M VEGF. Six hours later, retinal vascular permeability was measured using [³H]mannitol as a tracer. Mice treated with VEGF-TRAP_{R1R2} (9 mice, 18 eyes) had a significantly smaller retina to lung leakage ratio (RLLR) than mice treated with Fc fragment (9 mice, 18 eyes) indicating less breakdown of the BRB (A). Double transgenic rtTA/rho-TRE/VEGF mice with doxycycline-inducible expression of VEGF in the retina were given a subcutaneous injection of 25 mg/kg of VEGF-TRAP_{R1R2} (10 mice, 20 eyes) or Fc fragment (10 mice, 20 eyes) and on the following day they were started on 2 mg/ml of doxycycline in their drinking water. Two days later, they were given a second subcutaneous injection of 25 mg/kg of VEGF-TRAP_{R1R2} or Fc fragment was under the next day retinal vascular permeability was measured with [³H]mannitol as described in Materials and Methods. Double transgenic mice treated with VEGF-TRAP_{R1R2} had a significant reduction in the retina to lung leakage ratio compared to mice treated with Fc fragment (B).

et al., 1996; Yi et al., 1997). Using a combination of kinase inhibitors, we previously demonstrated that VEGF signaling is necessary for development of choroidal neovascularization after laser-induced rupture of Bruch's membrane (Kwak et al., 2000). In the present study, using VEGF-TRAP_{R1R2}, a completely different type of VEGF inhibitor that is highly specific, we have confirmed that VEGF plays a prominent role in the development of choroidal neovascularization.

Systemic administration of VEGF-TRAP_{R1R2} also markedly decreased neovascularization in rho/VEGF transgenic mice and reduced VEGF-induced breakdown of the BRB. Systemic administration of an earlier version of the VEGF-Trap also has been shown to reduce elevated ICAM-1 and eNOS levels, inhibit leukostasis, and normalize vascular permeability in the retinas of diabetic rodents (Qaumet al., 2001; Joussen et al., 2002; Poulaki et al., 2002). Thus, in model disease settings similar to diabetic retinopathy in humans, circulating VEGF-Traps penetrate into the retina and exert a strong therapeutic effect. The angiogenic stimulus is sustained in rho/VEGF mice, and subcutaneous injec-tions of VEGF-TRAP_{R1R2} every third day provided intraocular levels sufficient to neutralize this sustained stimulus. These data suggest that VEGF-TRAP_{R1R2} deserves consideration as a potential treatment for two complications of diabetic retinopathy, retinal neovascularization and macular edema.

The effects of long-term systemic inhibition of VEGF are unknown. While there are theoretical reasons why this could be problematic, VEGF inhibitors have been tested as adjuncts to chemotherapy in cancer trials, and there have not been reports of severe problems clearly linked to blockade of VEGF. Should systemic inhibition of VEGF prove problematic, there is an alternative, because we have shown that, as is the case for other anti-VEGF approaches (EyeTech Study Group, 2002; Kryzstolik et al., 2002), local administration of $VEGF-TRAP_{R1R2}$ by intravitreous injection is a viable alternative. A single intravitreous injection of VEGF- ${\rm TRAP}_{\rm R1R2}$ markedly suppressed the development of choroidal neovascularization over the course of two weeks.

This study suggests that VEGF-TRAP_{R1R2} has potential as a therapeutic agent for several VEGF-related retinal and choroidal diseases. Clinical trials are needed to assess the effect of subcutaneously administered VEGF-TRAP_{R1R2} in patients with retinal neovascularization and/or macular edema due to ischemic retinopathies including diabetic retinopathy and retinal vein occlusions, and in patients with choroidal neovascularization. Concurrently, additional preclinical studies should explore modes of local delivery to the eye that can be used adjunctively or as an alternative to systemic administration.

ACKNOWLEDGMENTS

The Research to Prevent Blindness grant was awarded to P.A.C. as Lew R. Wasserman Merit Awards.

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Anti-Angiogenic Properties of a New VEGF Antagonist, VEGF Trap, in a Mouse Model of Retinal Neovascularization

Q Wang; R Renard; J Cao; D Yancopouls; SJ Wiegand

+ Author Affiliations & Notes

Investigative Ophthalmology & Visual Science December 2002, Vol.43, 3714. doi:

Abstract

Abstract: : Purpose: Excessive upregulation of VEGF expression appears to be responsible for pathologic neovascularization in many retinal diseases. We have developed a new VEGF antagonist, VEGF Trap, that binds VEGF with high affinity thereby neutralizing its action. The current study investigates the anti-angiogenic properties of VEGF Trap in a mouse model of oxygen-induced retinopathy (OIR). Method: OIR mice were produced following the method developed by Smith et al (IOVS 1994, 35:101-111). VEGF Trap (25mg/kg body weight) was administered by intraperitoneal (ip) injection every other day from PN13 (12-24 hours after returning the mice from hyperoxia to room air) to PN17. Littermates exposed to the same regimen of hyperoxia, received ip injections of 50 µl of PBS upon to room air and served as controls. Eyes were taken on PN19, and one retina was flat mounted and stained with fluorescent Griffonia simplicifolia lectin B4 to visualize the retinal vasculature. The contralateral eye was embedded, sectioned and stained with hematoxylin and eosin. Results: One week following return to room air (PN19), the retinas of all control mice exposed to hyperoxia exhibited marked pathologic angiogenesis, characterized by the presence of vascular tufts penetrating the inner limiting membrane and chaotic sprouting of vessels on the surface of the retina. Administration of VEGFTrap almost completely blocked the development of these vascular abnormalities. Although pathologic angiogenesis was dramatically inhibited, administration of the VEGF trap did not block all retinal angiogenesis. Remarkably, by PN 19 much of the central retina was appropriately revascularized in animals treated with VEGF Trap, as evidenced by the regrowth of normal appearing vessels in the superficial, intermediate and deep layers. Conclusion: Systemic administration of VEGF Trap can efficiently suppress pathologic retinal angiogenesis

without blocking the appropriate revascularization of the previously ischemic retina. This finding distinguishes the anti-angiogenic properties of VEGF Trap from many other angiogenesis inhibitors studied in this model, which appear to be either substantially less effective in blocking pathologic angiogenesis (Aiello LP et al. PNAS 1995, 92:10457-10461), or which also compromise the appropriate revascularization of the retina (Ozaki et al. Am J Pathol 1997, 156:697-707).

Keywords: 566 retinal neovascularization • 423 growth factors/growth factor receptors

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EFS ID:	43680483			
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Confirmation Number:	5070			
Title of Invention:	USE OF A VEGF ANTAGONIST TO TREAT ANGIOGENIC EYE DISORDERS			
First Named Inventor/Applicant Name:	George YANCOPOULOS			
Customer Number:	96387			
Filer:	Karl Bozicevic/Kimberly Zuehlke			
Filer Authorized By:	Karl Bozicevic			
Attorney Docket Number:	REGN-008CIPCON10			
Receipt Date:	03-SEP-2021			
Filing Date:	21-JUN-2021			
Time Stamp:	11:35:25			
Application Type:	Utility under 35 USC 111(a)			

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6	Non Patent Literature	Cursiefen_2004_Inhibition_of_ hema.pdf	c20e10253dcf5992806ef217c653c91af9a5 6cfd	no	8
Warnings:					1
Information:					
		Cursiefen_2004_VEGF-	4515931		
7	Non Patent Literature	A_stimulates_lymphangiogene sis.pdf	d5cb6606895155d33ba1c3d3c5de790490 47e84c	no	12
Warnings:					
Information:					
			1454636		
8	Non Patent Literature	IPR2021-00880-2021-05-05_01 _Petition_for_Review_of_US96 69069_880.pdf	3eb7d60c61a299ac49257bbaa2f37a6efb84 0229	no	91
Warnings:					
Information:					

			767215		
9	Non Patent Literature	IPR2021-00880-2021-08-16_10 _POPR_Mylan_069_Patent.pdf	62b60957cb794ccdcc7ab77457f5d4c185c 699af	no	70
Warnings:					
Information:					
			1100357		
10	Non Patent Literature	IPR2021-00880- Ex_1002_Albini_Decl_880.pdf	4c924cf26f34134fb38f62959de15dc3aff6a 641	no	119
Warnings:			I		
Information:					
		IPR2021-00880-	6081340		
11	Non Patent Literature	Ex_1003_Gerritsen_Decl_880. pdf	ae519137a3b230b16a8818485f1e3c6284c e8a00	no	59
Warnings:			ł		
Information:					
		IPR2021-00881-2021-05-05_01	1239630		
12	Non Patent Literature	_Petition_for_IPR_of_9254338_ 881.pdf	6977365612a1c5c91994d47caf135b11523 38c64	no	89
Warnings:					
Information:					
			801029		
13	Non Patent Literature	IPR2021-00881-2021-08-16_10 _POPR_Mylan_338_Patent.pdf	3c8b306e28ac79c6dc86d9d327013014abf a8136	no	76
Warnings:		-	I		
Information:					
			23233259		
14	Non Patent Literature	IPR2021-00881- Ex_1002_Albini_Decl_881.pdf	309bc9a75b5ef775299bfa0ff7de17df378af 1a3	no	152
Warnings:					I
Information:					
		IPR2021-00881-	5491742		
15	Non Patent Literature	Ex_1003_Gerritsen_Decl_881. pdf	61cd4ed4ce04aa2bf5a9e9147c0ffeee6a69 585d	no	53
Warnings:					
Information:					

			306073		
16	Non Patent Literature	IPR2021-00881- Ex2001_Do_Declaration.pdf	9f39940c757a122c755015fb7978e3caf54af 627	no	19
Warnings:					
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			6781952		
17	Non Patent Literature	Mitchell_2018.pdf	dbf07aca9b56716a91c66dedab29d1b8e2b 03cde	no	9
Warnings:		1			
Information:					
			2073112		
18	Non Patent Literature	Nork_2011.pdf	0422baa35069c8a843f2285a00830fdd7464 6fe9	no	11
Warnings:			I		
Information:					
		PGR2021-00035-2021-01-07_02	2787300		
19	Non Patent Literature	_Petition_for_PGR_of_US10828 345.pdf	723df9b9c0e3571dc24ad99e395633fee94 380c5	no	92
Warnings:		+			
Information:					
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20	Non Patent Literature	PGR2021-00035-2021-04-15_06 _POPR.pdf	3e0210cd63d3078dd2aa8dfc142c599d4e8 222b6	no	96
Warnings:		•			
Information:					
			2852812		
21	Non Patent Literature	PGR2021-00035- Ex_1003_Wu_Declaration.pdf	674123a21173ad450aac4a919ea02c7fdc99 587d	no	81
Warnings:		+			
Information:					
			389537		
22	Non Patent Literature	PGR2021-00035- Ex_2001_Do_Declaration.pdf	aa9eefe29251b69326733d5b6e23289687d 9eddd	no	35
Warnings:		1			
Information:					

22		PGR2021-00035-	2704458		22			
23	Non Patent Literature	Ex_2002_D_Brown_Declaration .pdf	ffb2b452babea1bd305e93c4d4257b7124a 4c2dd	no	22			
Warnings:								
Information:								
			582298					
24	Non Patent Literature	Saishin_2003.pdf	2194d1ea1d4df56a20264398ec00939c017 df130	no	8			
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			41117					
25	Non Patent Literature	Wang_2002.pdf	ba3f92115f8ff1f74d748cf92929101bf40fc9 bc	no	2			
Warnings:								
Information:								
		Total Files Size (in bytes)	66	069892				
This Acknowledgement Receipt evidences receipt on the noted date by the USPTO of the indicated documents, characterized by the applicant, and including page counts, where applicable. It serves as evidence of receipt similar to a Post Card, as described in MPEP 503. New Applications Under 35 U.S.C. 111 If a new application is being filed and the application includes the necessary components for a filing date (see 37 CFR 1.53(b)-(d) and MPEP 506), a Filing Receipt (37 CFR 1.54) will be issued in due course and the date shown on this Acknowledgement Receipt will establish the filing date of the application. National Stage of an International Application under 35 U.S.C. 371 If a timely submission to enter the national stage of an international application is compliant with the conditions of 35 U.S.C. 371 and other applicable requirements a Form PCT/DO/EO/903 indicating acceptance of the application as a national stage submission under 35 U.S.C. 371 will be issued in addition to the Filing Receipt, in due course. New International Application is being filed and the international application includes the necessary components for an international filing date (see PCT Article 11 and MPEP 1810), a Notification of the International Application Number and of the International Filing Date (Form PCT/RO/105) will be issued in due course, subject to prescriptions concerning national security, and the date shown on this Acknowledgement Receipt will establish the international filing date of the application.								

Electronically Filed				
	Attorney Docket No.	REGN-008CIPCON10		
	Confirmation No.	5070		
SUPPLEMENTAL INFORMATION DISCLOSURE STATEMENT	First Named Inventor	George D. Yancopoulos		
	Application Number	17/352,892		
	Filing Date	June 21, 2021		
	Group Art Unit	To Be Assigned		
Address to:	Examiner Name	To Be Assigned		
Commissioner for Patents P.O. Box 1450 Alexandria, VA 22313-1450	Title: "Use of a VEGF Eye Disorders"	Antagonist to Treat Angiogenic		

Electronically Filed

Sir:

Applicant submits herewith documents which may be material to the examination of this application and in respect of which there may be a duty to disclose in accordance with 37 C.F.R. § 1.56. This submission is not intended to constitute an admission that any document referred to therein is "prior art" for this invention unless specifically designated as such. A listing of the documents is shown on enclosed Form PTO/SB/08A and copies of the foreign patents and non-patent literature are also enclosed.

The publications discussed herein are provided to comply with the duty to disclose in accordance with 37 C.F.R. § 1.56. However, nothing herein is to be construed as an admission that the present invention is not entitled to antedate such publication by virtue of prior invention. Further, the dates of publication provided may be different from the actual publication dates which may need to be independently confirmed.

The Examiner is requested to make the documents listed on the enclosed PTO/SB/08A of record in this application. Applicants would appreciate the Examiner initialing and returning the initialed copy of form PTO/SB/08A, indicating the documents cited therein have been considered and made of record herein.

Statements

No statement

PTA Statement under 37 CFR § 1.704(d)(1): Each item of information contained in the information disclosure statement filed herewith:

(i) Was first cited in any communication from a patent office in a counterpart foreign or international application or from the Office, and this communication was not received by any individual designated in § 1.56(c) more than thirty days prior to the filing of the information disclosure statement; or

(ii) Is a communication that was issued by a patent office in a counterpart foreign or international application or by the Office, and this communication was not received by any individual designated in § 1.56(c) more than thirty days prior to the filing of the information disclosure statement.

- **IDS Statement under 37 CFR § 1.97(e)(1):** Each item of information contained in the information disclosure statement was first cited in any communication from a foreign patent office in a counterpart foreign application not more than three months prior to the filing of the information disclosure statement; or
 - **IDS Statement under 37 CFR § 1.97(e)(2):** No item of information contained in the information disclosure statement was cited in a communication from a foreign patent office in a counterpart foreign application, and, to the knowledge of the person signing the certification after making reasonable inquiry, no item of information contained in the information disclosure statement was known to any individual designated in § 1.56(c) more than three months prior to the filing of the information disclosure statement.

<u>Fees</u>

 No fee is believed to be due.

The appropriate fee set forth in 37 C.F.R. §1.17(p) accompanies this information disclosure statement.

The Commissioner is hereby authorized to charge any underpayment of fees up to a strict limit of \$3,000.00 beyond that authorized on the credit card, but not more than \$3,000.00 in additional fees due with any communication for the above-referenced patent application, including but not limited to any necessary fees for extensions of time, or credit any overpayment of any amount to Deposit Account No. 50-0815, order number REGN-008CIPCON10.

Respectfully submitted, BOZICEVIC, FIELD & FRANCIS LLP

Date: <u>3 September 2021</u>

BOZICEVIC, FIELD & FRANCIS LLP 201 Redwood Shores Parkway, Suite 200 Redwood City, CA 94065 Telephone: (650) 327-3400 Facsimile: (650) 327-3231 By: <u>/Karl Bozicevic, Reg. No. 28,807/</u> Karl Bozicevic Reg. No. 28,807

UNITED STATES PATENT AND TRADEMARK OF			UNITED STATES DEPARTMENT United States Patent and Trade Address: COMMISSIONER FOR P. P.O. Box 1450 Alexandria, Virginia 22313-1450 www.uspto.gov	mark Office ATENTS
APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.
17/352,892	06/21/2021	George YANCOPOULOS	REGN-008CIPCON10	5070
	7590 10/01/2021		EXAM	INER
201 REDWOO	zicevic, Field & Francis D SHORES PARKWAY		CENTRAL,	DOCKET
SUITE 200 REDWOOD CI	ITY, CA 94065		ART UNIT	PAPER NUMBER
			OPAP	
			NOTIFICATION DATE	DELIVERY MODE
			10/01/2021	ELECTRONIC

Please find below and/or attached an Office communication concerning this application or proceeding.

The time period for reply, if any, is set in the attached communication.

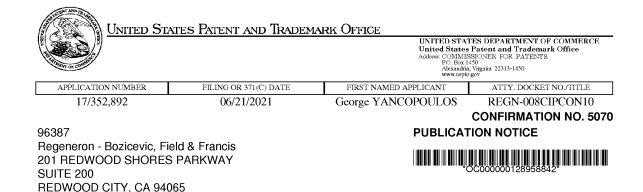
Notice of the Office communication was sent electronically on above-indicated "Notification Date" to the following e-mail address(es):

docket@bozpat.com

PTOL-90A (Rev. 04/07)

	Decisio	n Granting Request for	Application No. 17/352,892	Applicant(s) YANCOPOU	LOS, George
		ed Examination (Track I)	Examiner CHERYL P GIBSON BAYLOR	Art Unit OPET	AIA (FITF) Status No
1. T	HE REC	QUEST FILED <u>21 June 2021</u> IS <u>(</u>	GRANTED .		
Т	he abov A. B.	re-identified application has met t ✓ for an original nonprovision □ for an application undergoin	al application (Track I).		tion
		ve-identified application will ur special status throughout its ent			
	Α.	filing a petition for extension	of time to extend the ti	me period for fili	ng a reply;
	В.	filing an amendment to amen independent claims, more th			
	C.	filing a request for continued	examination;		
	D.	filing a notice of appeal;			
	E.	filing a request for suspension	of action;		
	F.	mailing of a notice of allowance	e;		
	G.	mailing of a final Office action	• •		
	Η.	completion of examination as	defined in 37 CFR 41.1	02; or	
	I.	abandonment of the application	n.		
Т	elephon	e inquiries with regard to this de	cision should be directe	d to CHERYL G	IBSON BAYLOR at
(571)272	-3213. In his/her absence, calls r	nay be directed to Petiti	on Help Desk a	t (571) 272-3282.
	CHERY	L GIBSON BAYLOR/			

U.S. Patent and Trademark Office PTO-2298 (Rev. 02-2012)



Title:USE OF A VEGF ANTAGONIST TO TREAT ANGIOGENIC EYE DISORDERS

Publication No.US-2021-0308217-A1 Publication Date:10/07/2021

NOTICE OF PUBLICATION OF APPLICATION

The above-identified application will be electronically published as a patent application publication pursuant to 37 CFR 1.211, et seq. The patent application publication number and publication date are set forth above.

The publication may be accessed through the USPTO's publically available Searchable Databases via the Internet at www.uspto.gov. The direct link to access the publication is currently http://www.uspto.gov/patft/.

The publication process established by the Office does not provide for mailing a copy of the publication to applicant. A copy of the publication may be obtained from the Office upon payment of the appropriate fee set forth in 37 CFR 1.19(a)(1). Orders for copies of patent application publications are handled by the USPTO's Public Records Division. The Public Records Division can be reached by telephone at (571) 272-3150 or (800) 972-6382, by facsimile at (571) 273-3250, by mail addressed to the United States Patent and Trademark Office, Public Records Division, Alexandria, VA 22313-1450 or via the Internet.

In addition, information on the status of the application, including the mailing date of Office actions and the dates of receipt of correspondence filed in the Office, may also be accessed via the Internet through the Patent Electronic Business Center at www.uspto.gov using the public side of the Patent Application Information and Retrieval (PAIR) system. The direct link to access this status information is currently https://portal.uspto.gov/pair/PublicPair. Prior to publication, such status information is confidential and may only be obtained by applicant using the private side of PAIR.

Further assistance in electronically accessing the publication, or about PAIR, is available by calling the Patent Electronic Business Center at 1-866-217-9197.

Office of Data Managment, Application Assistance Unit (571) 272-4000, or (571) 272-4200, or 1-888-786-0101

page 1 of 1

To:	docket@bozpat.com,,
From:	PAIR_eOfficeAction@uspto.gov
Cc:	PAIR_eOfficeAction@uspto.gov
Subject:	Private PAIR Correspondence Notification for Customer Number 96387

Oct 08, 2021 04:59:58 AM

Dear PAIR Customer:

Regeneron - Bozicevic, Field & Francis 201 REDWOOD SHORES PARKWAY SUITE 200 REDWOOD CITY, CA 94065 UNITED STATES

The following USPTO patent application(s) associated with your Customer Number, 96387, have new outgoing correspondence. This correspondence is now available for viewing in Private PAIR.

The official date of notification of the outgoing correspondence will be indicated on the form PTOL-90 accompanying the correspondence.

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The list of documents shown below is provided as a courtesy and is not part of the official file wrapper. The content of the images shown in PAIR is the official record.

Application	Document	Mailroom Date	Attorney Docket No.
17352892	NTC.PUB	10/07/2021	REGN-008CIPCON10

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Thank you for prompt attention to this notice,

UNITED STATES PATENT AND TRADEMARK OFFICE PATENT APPLICATION INFORMATION RETRIEVAL SYSTEM

UNITED STATES PATENT AND TRADEMARK OFFICE			UNITED STATES DEPARTMENT United States Patent and Trade Address: COMMISSIONER FOR P. P.O. Box 1450 Alexandria, Virginia 22313-1450 www.uspto.gov	mark Office ATENTS
APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.
17/352,892	06/21/2021	George YANCOPOULOS	REGN-008CIPCON10	5070
	7590 10/28/2021		EXAM	INER
201 REDWOO	ozicevic, Field & Francis D SHORES PARKWAY		LOCKARD, JON	MCCLELLAND
SUITE 200 REDWOOD CI	ITY, CA 94065		ART UNIT	PAPER NUMBER
			1647	
			NOTIFICATION DATE	DELIVERY MODE
			10/28/2021	ELECTRONIC

Please find below and/or attached an Office communication concerning this application or proceeding.

The time period for reply, if any, is set in the attached communication.

Notice of the Office communication was sent electronically on above-indicated "Notification Date" to the following e-mail address(es):

docket@bozpat.com

PTOL-90A (Rev. 04/07)

	Application No. 17/352,892	Applicant(s)) LOS, George
Office Action Summary	Examiner	Art Unit	AIA (FITF) Status
	JON M LOCKARD	1647	No
The MAILING DATE of this communication app Period for Reply	ears on the cover sheet with the c	corresponden	ce address
A SHORTENED STATUTORY PERIOD FOR REPLY IS SET TO EXPIRE <u>3</u> MONTHS FROM THE MAILING DATE OF THIS COMMUNICATION. - Extensions of time may be available under the provisions of 37 CFR 1.136(a). In no event, however, may a reply be timely filed after SIX (6) MONTHS from the mailing date of this communication. - If NO period for reply is specified above, the maximum statutory period will apply and will expire SIX (6) MONTHS from the mailing date of this communication. - Failure to reply within the set or extended period for reply will, by statute, cause the application to become ABANDONED (35 U.S.C. § 133). Any reply received by the Office later than three months after the mailing date of this communication, even if timely filed, may reduce any earned patent term adjustment. See 37 CFR 1.704(b).			
Status			
 1) Responsive to communication(s) filed on <u>21 June 2021</u>. □ A declaration(s)/affidavit(s) under 37 CFR 1.130(b) was/were filed on 2a) □ This action is FINAL. 2b) This action is non-final. 3) □ An election was made by the applicant in response to a restriction requirement set forth during the interview on; the restriction requirement and election have been incorporated into this action. 4) □ Since this application is in condition for allowance except for formal matters, prosecution as to the merits is closed in accordance with the practice under <i>Ex parte Quayle</i>, 1935 C.D. 11, 453 O.G. 213. 			
 Disposition of Claims* 5) Claim(s) 21-50 is/are pending in the application. 5a) Of the above claim(s) is/are withdrawn from consideration. 6) Claim(s) is/are allowed. 7) Claim(s) 21-50 is/are rejected. 8) Claim(s) is/are objected to. 9) Claim(s) is/are objected to. 9) Claim(s) are subject to restriction and/or election requirement * If any claims have been determined allowable, you may be eligible to benefit from the Patent Prosecution Highway program at a participating intellectual property office for the corresponding application. For more information, please see http://www.uspto.gov/patents/init_events/pph/index.jsp or send an inquiry to PPHfeedback@uspto.gov. 			
 Application Papers 10) ☐ The specification is objected to by the Examiner. 11) ☑ The drawing(s) filed on <u>21 June 2021</u> is/are: a) ☑ accepted or b) ☐ objected to by the Examiner. Applicant may not request that any objection to the drawing(s) be held in abeyance. See 37 CFR 1.85(a). Replacement drawing sheet(s) including the correction is required if the drawing(s) is objected to. See 37 CFR 1.121(d). Priority under 35 U.S.C. § 119 			
 12) Acknowledgment is made of a claim for foreign priority under 35 U.S.C. § 119(a)-(d) or (f). Certified copies: a) All b) Some** c) None of the: Certified copies of the priority documents have been received. Certified copies of the priority documents have been received in Application No Copies of the certified copies of the priority documents have been received in this National Stage application from the International Bureau (PCT Rule 17.2(a)). ** See the attached detailed Office action for a list of the certified copies not received. 			
Attachment(s) 1) Notice of References Cited (PTO-892)	3) 🗖 Interview Summer	((PTO 419)	
 2) ✓ Information Disclosure Statement(s) (PTO/SB/08a and/or PTO/S Paper No(s)/Mail Date U.S. Patent and Trademark Office 			

Office Action Summary

Notice of Pre-AIA or AIA Status

1. The present application is being examined under the pre-AIA first to invent provisions.

DETAILED ACTION

Status of Application, Amendments, and/or Claims

2. The Preliminary Amendment filed on 21 June 2021 has been entered in full. Claims 1-20 have been cancelled, and claims 21-50 have been added. Therefore, claims 21-50 are pending and the subject of this Office Action.

Information Disclosure Statement

3. The information disclosure statements (IDS) filed 21 June 2021, 09 July 2021 and 03 September 2021 have been considered by the examiner.

Double Patenting

4. The nonstatutory double patenting rejection is based on a judicially created doctrine grounded in public policy (a policy reflected in the statute) so as to prevent the unjustified or improper timewise extension of the "right to exclude" granted by a patent and to prevent possible harassment by multiple assignees. A nonstatutory double patenting rejection is appropriate where the claims at issue are not identical, but at least one examined application claim is not patentably distinct from the reference claim(s) because the examined application claim is either anticipated by, or would have been obvious over, the reference claim(s). See, e.g., *In re Berg*, 140 F.3d 1428, 46 USPQ2d 1226 (Fed. Cir. 1998); *In re Goodman*, 11 F.3d 1046, 29 USPQ2d 2010 (Fed. Cir. 1993); *In re Longi*, 759 F.2d 887, 225 USPQ 645 (Fed. Cir. 1985); *In re Van*

Ornum, 686 F.2d 937, 214 USPQ 761 (CCPA 1982); *In re Vogel*, 422 F.2d 438, 164 USPQ 619 (CCPA 1970); and *In re Thorington*, 418 F.2d 528, 163 USPQ 644 (CCPA 1969).

5. A timely filed terminal disclaimer in compliance with 37 CFR 1.321(c) or 1.321(d) may be used to overcome an actual or provisional rejection based on a nonstatutory double patenting ground provided the reference application or patent either is shown to be commonly owned with this application, or claims an invention made as a result of activities undertaken within the scope of a joint research agreement. A terminal disclaimer must be signed in compliance with 37 CFR 1.321(b).

6. The USPTO internet Web site contains terminal disclaimer forms which may be used. Please visit http://www.uspto.gov/forms/. The filing date of the application will determine what form should be used. A web-based eTerminal Disclaimer may be filled out completely online using web-screens. An eTerminal Disclaimer that meets all requirements is auto-processed and approved immediately upon submission. For more information about eTerminal Disclaimers, refer to http://www.uspto.gov/patents/process/file/efs/guidance/eTD-info-I.jsp.

7. Claims 21-50 are rejected on the ground of nonstatutory obviousness-type double patenting as being unpatentable over claims 1-26 of U.S. Patent No. 9,254,338. Although the conflicting claims are not identical, as they recite different dosing schedules, they are not patentably distinct from each other because claims 1-26 of the '338 patent are drawn to a method for treating an angiogenic eye disorder, including age-related macular degeneration, diabetic retinopathy, choroidal neovascularization, vascular leak, and/or retinal edema, comprising administering a fusion polypeptide having the amino acid sequence set forth in SEQ ID NO:2, which comprises an immunoglobin-like (Ig) domain 2 of a first VEGF receptor (VEGFR1) and Ig domain 3 of a second VEGF receptor (VEGFR2) and a multimerizing component, which is what aflibercept

comprises. While the '338 patent does not disclose the dosing schedules set forth in the instant claims, it is routine experimentation to optimize dosages and dosage schedules. The courts have determined that:

"[W]here the general conditions of a claim are disclosed in the prior art, it is not inventive to discover the optimum or workable ranges by routine experimentation." *In re Aller*, 220 F.2d 454, 454, 105 USPQ 223,235, (CCPA 1955).
Therefore, the claims are overlapping in scope.

8. Claims 21-50 are rejected on the ground of nonstatutory obviousness-type double patenting as being unpatentable over claims 1-12 of U.S. Patent No. 9,669,069. Although the conflicting claims are not identical, as they recite different dosing schedules, they are not patentably distinct from each other because claims 1-12 of the '069 patent are drawn to a method for treating an angiogenic eye disorder, including age-related macular degeneration, diabetic retinopathy, diabetic macular edema, central retinal vein occlusion, branch retinal vein occlusion, and corneal neovascularization, comprising administering a fusion polypeptide having the amino acid sequence set forth in SEQ ID NO:2, which comprises an immunoglobin-like (Ig) domain 2 of a first VEGF receptor (VEGFR1) and Ig domain 3 of a second VEGF receptor (VEGFR2) and a multimerizing component, which is what aflibercept comprises. While the '069 patent does not disclose the dosing schedules set forth in the instant claims, it is routine experimentation to optimize dosages and dosage schedules. The courts have determined that:

"[W]here the general conditions of a claim are disclosed in the prior art, it is not inventive to discover the optimum or workable ranges by routine experimentation." *In re Aller*, 220 F.2d 454, 454, 105 USPQ 223,235, (CCPA 1955).

Therefore, the claims are overlapping in scope.

9. Claims 21-50 are rejected on the ground of nonstatutory obviousness-type double patenting as being unpatentable over claims 1-12 of U.S. Patent No. 10,130,681. Although the conflicting claims are not identical, as they recite different dosing schedules, they are not patentably distinct from each other because claims 1-12 of the '681 patent are drawn to a method for treating an angiogenic eye disorder, including age-related macular degeneration, diabetic retinopathy, diabetic macular edema, central retinal vein occlusion, branch retinal vein occlusion, and corneal neovascularization, comprising administering a fusion polypeptide having the amino acid sequence set forth in SEQ ID NO:2, which comprises an immunoglobin-like (Ig) domain 2 of a first VEGF receptor (VEGFR1) and Ig domain 3 of a second VEGF receptor (VEGFR2) and a multimerizing component, which is what aflibercept comprises. While the '681 patent does not disclose the dosing schedules set forth in the instant claims, it is routine experimentation to optimize dosages and dosage schedules. The courts have determined that:

"[W]here the general conditions of a claim are disclosed in the prior art, it is not inventive to discover the optimum or workable ranges by routine experimentation." *In re Aller*, 220 F.2d 454, 454, 105 USPQ 223,235, (CCPA 1955).

Therefore, the claims are overlapping in scope.

10. Claims 21-50 are rejected on the ground of nonstatutory obviousness-type double patenting as being unpatentable over claims 1-11 of U.S. Patent No. 10,828,345. Although the conflicting claims are not identical, as they recite different dosing schedules, they are not patentably distinct from each other because claims 1-11 of the '345 patent are drawn to a method for treating an angiogenic eye disorder, including age-related macular degeneration, diabetic retinopathy, diabetic macular edema, central retinal vein occlusion, branch retinal vein occlusion, and corneal neovascularization, comprising administering a VEGF antagonist, wherein the VEGF comprises

an immunoglobin-like (Ig) domain 2 of Flt1 and Ig domain 3 of Flk1 and a multimerizing component, or aflibercept. While the '345 patent does not disclose the dosing schedules set forth in the instant claims, it is routine experimentation to optimize dosages and dosage schedules. The courts have determined that:

"[W]here the general conditions of a claim are disclosed in the prior art, it is not inventive to discover the optimum or workable ranges by routine experimentation." *In re Aller*, 220 F.2d 454, 454, 105 USPQ 223,235, (CCPA 1955).
Therefore, the claims are overlapping in scope.

11. Claims 21-50 are rejected on the ground of nonstatutory obviousness-type double patenting as being unpatentable over claims 1-47 of U.S. Patent No. 10,888,601. Although the conflicting claims are not identical, as they recite different dosing schedules, they are not patentably distinct from each other because claims 1-47 of the '601 patent are drawn to a method for treating an angiogenic eye disorder, including age-related macular degeneration, diabetic retinopathy, diabetic macular edema, central retinal vein occlusion, branch retinal vein occlusion, and corneal neovascularization, comprising administering aflibercept. While the '601 patent does not disclose the dosing schedules set forth in the instant claims, it is routine experimentation to optimize dosages and dosage schedules. The courts have determined that:

"[W]here the general conditions of a claim are disclosed in the prior art, it is not inventive to discover the optimum or workable ranges by routine experimentation." *In re Aller*, 220 F.2d 454, 454, 105 USPQ 223,235, (CCPA 1955).

Therefore, the claims are overlapping in scope.

Summary

12. No claim is allowed.

Advisory Information

Any inquiry concerning this communication or earlier communications from the examiner should be directed to Jon M. Lockard whose telephone number is (571) 272-2717. The examiner can normally be reached on Monday through Friday, 8:00 AM to 4:30 PM.

If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, Joanne Hama, can be reached on (571) 272-2911. The fax number for the organization where this application or proceeding is assigned is 571-273-8300.

Information regarding the status of an application may be obtained from the Patent Application Information Retrieval (PAIR) system. Status information for published applications may be obtained from either Private PAIR or Public PAIR. Status information for unpublished applications is available through Private PAIR only. For more information about the PAIR system, see http://pair-direct.uspto.gov. Should you have questions on access to the Private PAIR system, contact the Electronic Business Center (EBC) at **866-217-9197** (toll-free). If you would like assistance from a USPTO Customer Service Representative or access to the automated information system, call 800-786-9199 (IN USA OR CANADA) or 571-272-1000.

/JON M LOCKARD/ Examiner, Art Unit 1647 October 22, 2021

	Application/Control No.	Applicant(s)/Patent Under Reexamination
Search Notes	17/352,892	YANCOPOULOS, George
	Examiner	Art Unit
	JON M LOCKARD	1647

CPC - Searched*		
Symbol	Date	Examiner

CPC Combination Sets - Searched*		
Symbol	Date	Examiner

US Classification - Searched*			
Class	Subclass	Date	Examiner
NONE		10/23/2021	JML

* See search history printout included with this form or the SEARCH NOTES box below to determine the scope of the search.

Search Notes		
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EAST (USPAT, US-PGPUB, EPO, DERWENT): See attached search history.	10/23/2021	JML
STN (MEDLINE, SCISEARCH, EMBASE, BIOSIS): See attached search history.	10/23/2021	JML
PALM: Inventor search.	10/23/2021	JML

Interference Sea	arch		
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U.S. Patent and Trademark Office	Part of Paper No.: 20211021

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EAST Search History

EAST Search History (Prior Art)

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L1	9,181	(flt1 or vegfr1 or (vegf adj r1)) same ((flk1 or kdr or vegfr2 or (vegf adj r2)) or (Flt4 vegfr3 or (vegf adj r3)))	US-PGPUB; USPAT; EPO; DERWENT	OR	ON	2021/10/23 15:56
L2	1,053	l1 same ((chimer\$ or fusion) same vegf)	US-PGPUB; USPAT; EPO; DERWENT	OR	ON	2021/10/23 15:56
L3	8,958	(flt1 or vegfr1 or (vegf adj r1)) with ((flk1 or kdr or vegfr2 or (vegf adj r2)) or (Flt4 vegfr3 or (vegf adj r3)))	US-PGPUB; USPAT; EPO; DERWENT	OR	ON	2021/10/23 15:56
L4	536	l3 with ((chimer\$ or fusion) with vegf)	US-PGPUB; USPAT; EPO; DERWENT	OR	ON	2021/10/23 15:56
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L6	292	(l2 l4 l5) same ((eye or ocular or retina\$ or macular) with disorder)	US-PGPUB; USPAT; EPO; DERWENT	OR	ON	2021/10/23 16:00
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L9	534	yancopoulos-g\$.in.	US-PGPUB; USPAT; EPO; DERWENT	OR	ON	2021/10/23 16:03
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| INVENTORS
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Inventor Information for 17/352892

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| YANCOPOULOS, GEORGE | YORK TOWN HEIGHTS | NEW YORK | | | | |
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| INFORMATION DISCLOSURE STATEMENT | Regeneron Pharmaceuticals, Inc. | |
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| From: | PAIR_eOfficeAction@uspto.gov |
| Cc: | PAIR_eOfficeAction@uspto.gov |
| Subject: | Private PAIR Correspondence Notification for Customer Number 96387 |

Oct 28, 2021 03:59:18 AM

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Regeneron - Bozicevic, Field & Francis 201 REDWOOD SHORES PARKWAY SUITE 200 REDWOOD CITY, CA 94065 UNITED STATES

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| Application | Document | Mailroom Date | Attorney Docket No. |
|-------------|----------|---------------|---------------------|
| 17352892 | CTNF | 10/28/2021 | REGN-008CIPCON10 |
| | 1449 | 10/28/2021 | REGN-008CIPCON10 |
| | 1449 | 10/28/2021 | REGN-008CIPCON10 |
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| EXAMINER | DATE CONSIDERED |
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| EXAMINER: Initial if reference considered, whether or not citation is in conformance with MPEP 609. Draw line three considered. Include copy of this form with next communication to Applicant. | ough citation if not in conformance and not |
| *Copies of the listed references are either submitted herewith or were previously cited by or submitted to, the Office in a 1.97(d) and MPEP §609, the indicated reference may have been previously cited by or submitted to, the Office in a prior identified by its U.S. Application Number in this Information Disclosure Statement. | |

| | | Page 12 of 12 |
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| SUBSTITUTE 1449
INFORMATION DISCLOSURE STATEMENT | ATTY. DOCKET NO. | APPLICATION NO. |
| | REGN-008CIPCON10 | 17/352,892 |
| | APPLICANT | |
| | REGENERON PHARMACEUTICALS | , INC. |
| | FILING DATE | GROUP |
| | June 21, 2021 | 1647 |

| | NON-PATENT LITERATURE DOCUMENTS - UPDATES TO PREVIOUS IDS CITATI | ONS |
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| | DOCUMENT (Including Author, Title, Date, Pertinent Pages, etc.) | REFERENCE PROVIDED* |
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| EXAMINER | DATE CONSIDERED |
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| EXAMINER: Initial if reference considered, whether or not citation is in conformance with MPEP 609. Draw line the considered. Include copy of this form with next communication to Applicant. | ough citation if not in conformance and not |
| *Copies of the listed references are either submitted herewith or were previously cited by or submitted to, the Office in 1.97(d) and MPEP §609, the indicated reference may have been previously cited by or submitted to, the Office in a prior identified by its U.S. Application Number in this Information Disclosure Statement. | |

| Electronic Patent Application Fee Transmittal | | | | | | |
|---|--|----------------------|-----------|-------------|-------------------------|--|
| Application Number: | 17 | 17352892 | | | | |
| Filing Date: | 21 | -Jun-2021 | | | | |
| Title of Invention: | USE OF A VEGF ANTAGONIST TO TREAT ANGIOGENIC EYE DISORDERS | | | E DISORDERS | | |
| First Named Inventor/Applicant Name: | Ge | orge YANCOPOULC |)S | | | |
| Filer: | Ка | rl Bozicevic/Kimberl | y Zuehlke | | | |
| Attorney Docket Number: | RE | GN-008CIPCON10 | | | | |
| Filed as Large Entity | | | | | | |
| Filing Fees for Utility under 35 USC 111(a) | | | | | | |
| Description | | Fee Code | Quantity | Amount | Sub-Total in
USD(\$) | |
| Basic Filing: | | | | | | |
| Pages: | | | | | | |
| Claims: | | | | | | |
| Miscellaneous-Filing: | | | | | | |
| Petition: | | | | | | |
| Patent-Appeals-and-Interference: | | | | | | |
| Post-Allowance-and-Post-Issuance: | | | | | | |
| Extension-of-Time: | | | | | | |

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| Electronic Acknowledgement Receipt | | | |
|--------------------------------------|--|--|--|
| EFS ID: | 44366548 | | |
| Application Number: | 17352892 | | |
| International Application Number: | | | |
| Confirmation Number: | 5070 | | |
| Title of Invention: | USE OF A VEGF ANTAGONIST TO TREAT ANGIOGENIC EYE DISORDERS | | |
| First Named Inventor/Applicant Name: | George YANCOPOULOS | | |
| Customer Number: | 96387 | | |
| Filer: | Karl Bozicevic/Kimberly Zuehlke | | |
| Filer Authorized By: | Karl Bozicevic | | |
| Attorney Docket Number: | REGN-008CIPCON10 | | |
| Receipt Date: | 24-NOV-2021 | | |
| Filing Date: | 21-JUN-2021 | | |
| Time Stamp: | 14:56:40 | | |
| Application Type: | Utility under 35 USC 111(a) | | |

Payment information:

| Submitted with Payment | yes | | |
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| Payment Type | CARD | | |
| Payment was successfully received in RAM | \$260 | | |
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| Document
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| 1 | Transmittal Letter | REGN-008CIPCON10_2021-11-2
4_SuppIDS_Trans.pdf | 162b8919da83cc173675eac4fd63bd0dac9
04188 | no | 3 |
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| 2 | Information Disclosure Statement (IDS)
Form (SB08) | REGN-008CIPCON10_2021-11-2
4_SuppIDS_1449.pdf | bbe2331eadef194d51c7ba0712d7eac5a71
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| | Attorney Docket No. | REGN-008CIPCON10 | |
| | Confirmation No. | 5070 | |
| SUPPLEMENTAL INFORMATION | First Named Inventor | George D. Yancopoulos | |
| DISCLOSURE STATEMENT | Application Number | 17/352,892 | |
| | Filing Date | June 21, 2021 | |
| | Group Art Unit | 1647 | |
| Address to: | Examiner Name | Jon McClelland Lockard | |
| Commissioner for Patents
P.O. Box 1450
Alexandria, VA 22313-1450 | Title: <i>"Use of a VEGF Antagonist to Treat Angiogenic Eye Disorders"</i> | | |

Electronically Filed

Sir:

The attention of the Examiner is invited to the documents listed on the attached Substitute 1449. Copies of the U.S. patents and published applications listed on the attached Substitute 1449 are not submitted herewith, in accordance with the Strategic Plan Final Rule, 69 Fed. Reg. 56481-56547 (September 21, 2004), effective October 21, 2004.

Copies of the foreign publications and non-patent literature documents listed on the attached Substitute 1449 are submitted in parent U.S. Application No. 17/072,417. Applicant respectfully submits that a subset of references submitted herein were previously submitted in this or a priority application. Nonetheless, Applicant is submitting these previously submitted references to provide an accurate reference citation or to provide a clearer copy of the reference.

Applicant notes that the transmittal letter accompanying the Information Disclosure Statement submitted for this application on July 9, 2021, incorrectly recited that "[a]ll of the references identified herein were disclosed in parent application serial number 17/350,958." Accordingly, the citations previously submitted in the July 9, 2021 Information Disclosure Statement are resubmitted here as Ref. Nos. 75 to 143 in order to correct the record. Applicant notes that this group of resubmitted citations accounts for part of the citations provided herein.

Applicant would also like to bring to the Examiner's attention that the PTAB has instituted *inter partes* reviews for related U.S. Patent Nos. 9,254,338 and 9,669,069.

It is respectfully requested that the information above be expressly considered during the prosecution of this application, and that the documents be made of record therein and appear among the "References Cited" on any patent to issue therefrom.

No aspect of these submissions constitute admission of prior art status or a disclaimer of claim scope.

Statements

No statement

PTA Statement under 37 CFR § 1.704(d)(1): Each item of information contained in the information disclosure statement filed herewith:

(i) Was first cited in any communication from a patent office in a counterpart foreign or international application or from the Office, and this communication was not received by any individual designated in § 1.56(c) more than thirty days prior to the filing of the information disclosure statement; or

(ii) Is a communication that was issued by a patent office in a counterpart foreign or international application or by the Office, and this communication was not received by any individual designated in § 1.56(c) more than thirty days prior to the filing of the information disclosure statement.

- **IDS Statement under 37 CFR § 1.97(e)(1):** Each item of information contained in the information disclosure statement was first cited in any communication from a foreign patent office in a counterpart foreign application not more than three months prior to the filing of the information disclosure statement; or
- **IDS Statement under 37 CFR § 1.97(e)(2):** No item of information contained in the information disclosure statement was cited in a communication from a foreign patent office in a counterpart foreign application, and, to the knowledge of the person signing the certification after making reasonable inquiry, no item of information contained in the information disclosure statement was known to any individual designated in § 1.56(c) more than three months prior to the filing of the information disclosure statement.

Fees

No fee is believed to be due.

The appropriate fee set forth in 37 C.F.R. §1.17(p) accompanies this information disclosure statement.

Atty Docket No.: REGN-008CIPCON10 USSN: 17/352,892

The Commissioner is hereby authorized to charge any underpayment of fees up to a strict limit of \$3,000.00 beyond that authorized on the credit card, but not more than \$3,000.00 in additional fees due with any communication for the above-referenced patent application, including but not limited to any necessary fees for extensions of time, or credit any overpayment of any amount to Deposit Account No. 50-0815, order number REGN-008CIPCON10.

Respectfully submitted, BOZICEVIC, FIELD & FRANCIS LLP

Date: 24 November 2021

By: <u>/Karl Bozicevic, Reg. No. 28,807/</u> Karl Bozicevic Reg. No. 28,807

BOZICEVIC, FIELD & FRANCIS LLP 201 Redwood Shores Parkway, Suite 200 Redwood City, CA 94065 Telephone: (650) 327-3400 Facsimile: (650) 327-3231 Under the Paperwork Reduction Act of 1995, no persons are required to respond to a collection of information unless it displays a valid OMB control number

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| the boxes below. | | | | | | | |
| | Applic | cation Number | | Filing Dat | e | | |
| | | 17/352,892 | | June | 21, 2021 | | |
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| Assignee or F | Person t | to Whom the Inventor is Under | r an Obligation f | o Assign (p | rovide signer's ti | tle if appli | cant is a juristic entity) |
| Person Who Otherwise Shows Sufficient Proprietary Interest (e.g., a petition under 37 CFR 1.46(b)(2) was granted in the application or is concurrently being filed with this document) (provide signer's title if applicant is a juristic entity) | | | | | | | |
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| Total of 1 | | ms are submitted. | | | | | |
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| STATEMENT UNDER 37 CFR 3.73(c) | |
| Applicant/Patent Owner: Regeneron Pharmaceuticals, Inc. | |
| Application No./Patent No.: 17/352,892 Filed/Issue Date: June 21, 2021 | _ |
| Titled: Use of a VEGF Antagonist to Treat Angiogenic Eye Disorders | _ |
| Regeneron Pharmaceuticals, Inc. , a Corporation | _ |
| (Name of Assignee) (Type of Assignee, e.g., corporation, partnership, university, government agency, etc.) | |
| states that, for the patent application/patent identified above, it is (choose one of options 1, 2, 3 or 4 below): | |
| 1. 🕑 The assignee of the entire right, title, and interest. | |
| 2. An assignee of less than the entire right, title, and interest (check applicable box): | |
| The extent (by percentage) of its ownership interest is%. Additional Statement(s) by the owners holding the balance of the interest <u>must be submitted</u> to account for 100% of the ownership interest. | |
| There are unspecified percentages of ownership. The other parties, including inventors, who together own the entiripht, title and interest are: | re |
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| 3. The assignee of an undivided interest in the entirety (a complete assignment from one of the joint inventors was made | 1 |
| The other parties, including inventors, who together own the entire right, title, and interest are: | ,. |
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| Additional Statement(s) by the owner(s) holding the balance of the interest <u>must be submitted</u> to account for the entiright, title, and interest. | re |
| 4. The recipient, via a court proceeding or the like (<i>e.g.</i> , bankruptcy, probate), of an undivided interest in the entirety (a complete transfer of ownership interest was made). The certified document(s) showing the transfer is attached. | |
| The interest identified in option 1, 2 or 3 above (not option 4) is evidenced by either (choose one of options A or B below): | |
| | |
| A. [-] An assignment from the inventor(s) of the patent application/patent identified above. The assignment was recorded in the United States Patent and Trademark Office at Reel <u>057417</u> , Frame <u>0322</u> , or for which a copy thereof is attached. | |
| B. 🗌 A chain of title from the inventor(s), of the patent application/patent identified above, to the current assignee as follows | : |
| 1. From: To: | |
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| [Page 1 of 2] | |

[Page 1 of 2] This collection of information is required by 37 CFR 3.73(b). The information is required to obtain or retain a benefit by the public which is to file (and by the USPTO to process) an application. Confidentiality is governed by 35 U.S.C. 122 and 37 CFR 1.11 and 1.14. This collection is estimated to take 12 minutes to complete, including gathering, preparing, and submitting the completed application form to the USPTO. Time will vary depending upon the individual case. Any comments on the amount of time you require to complete this form and/or suggestions for reducing this burden, should be sent to the Chief Information Officer, U.S. Patent and Trademark Office, U.S. Department of Commerce, P.O. Box 1450, Alexandria, VA 22313-1450. DO NOT SEND FEES OR COMPLETED FORMS TO THIS ADDRESS. **SEND TO: Commissioner for Patents, P.O. Box 1450, Alexandria, VA 22313-1450**.

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ted for recordation pursuant | n of title from the original owner to the to 37 CFR 3.11. | |
| [NOTE: A separate copy (i.e., a true copy of the original assignment document(s)) must be submitted to Assignment
Division in accordance with 37 CFR Part 3, to record the assignment in the records of the USPTO. See MPEP 302.08] | | | | | |
| The undersigned (whose title is supplied below) is authorized to act on behalf of the assignee. | | | | | |
| /Karl Bozi | icevic, Reg. N | 0. 28,807/ | | 29 November 2021 | |
| Signature | | | | Date | |
| Karl Bozicevic 28,807 | | | | | |
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[Page 2 of 2]

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- 2. A record from this system of records may be disclosed, as a routine use, in the course of presenting evidence to a court, magistrate, or administrative tribunal, including disclosures to opposing counsel in the course of settlement negotiations.
- 3. A record in this system of records may be disclosed, as a routine use, to a Member of Congress submitting a request involving an individual, to whom the record pertains, when the individual has requested assistance from the Member with respect to the subject matter of the record.
- 4. A record in this system of records may be disclosed, as a routine use, to a contractor of the Agency having need for the information in order to perform a contract. Recipients of information shall be required to comply with the requirements of the Privacy Act of 1974, as amended, pursuant to 5 U.S.C. 552a(m).
- A record related to an International Application filed under the Patent Cooperation Treaty in this system of records may be disclosed, as a routine use, to the International Bureau of the World Intellectual Property Organization, pursuant to the Patent Cooperation Treaty.
- 6. A record in this system of records may be disclosed, as a routine use, to another federal agency for purposes of National Security review (35 U.S.C. 181) and for review pursuant to the Atomic Energy Act (42 U.S.C. 218(c)).
- 7. A record from this system of records may be disclosed, as a routine use, to the Administrator, General Services, or his/her designee, during an inspection of records conducted by GSA as part of that agency's responsibility to recommend improvements in records management practices and programs, under authority of 44 U.S.C. 2904 and 2906. Such disclosure shall be made in accordance with the GSA regulations governing inspection of records for this purpose, and any other relevant (*i.e.*, GSA or Commerce) directive. Such disclosure shall not be used to make determinations about individuals.
- 8. A record from this system of records may be disclosed, as a routine use, to the public after either publication of the application pursuant to 35 U.S.C. 122(b) or issuance of a patent pursuant to 35 U.S.C. 151. Further, a record may be disclosed, subject to the limitations of 37 CFR 1.14, as a routine use, to the public if the record was filed in an application which became abandoned or in which the proceedings were terminated and which application is referenced by either a published application, an application open to public inspection or an issued patent.
- 9. A record from this system of records may be disclosed, as a routine use, to a Federal, State, or local law enforcement agency, if the USPTO becomes aware of a violation or potential violation of law or regulation.

| Electronic Acknowledgement Receipt | | | |
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| EFS ID: | 44381316 | | |
| Application Number: | 17352892 | | |
| International Application Number: | | | |
| Confirmation Number: | 5070 | | |
| Title of Invention: | USE OF A VEGF ANTAGONIST TO TREAT ANGIOGENIC EYE DISORDERS | | |
| First Named Inventor/Applicant Name: | George YANCOPOULOS | | |
| Customer Number: | 96387 | | |
| Filer: | Karl Bozicevic/Kimberly Zuehlke | | |
| Filer Authorized By: | Karl Bozicevic | | |
| Attorney Docket Number: | REGN-008CIPCON10 | | |
| Receipt Date: | 29-NOV-2021 | | |
| Filing Date: | 21-JUN-2021 | | |
| Time Stamp: | 14:43:18 | | |
| Application Type: | Utility under 35 USC 111(a) | | |

Payment information:

| Submitted with Payment | | | no | | | |
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| APPLICATION NUMBER | FILING OR 371(C) DATE | FIRST NAMED APPLICANT | ATTY. DOCKET NO./TITLE |
| 17/352,892 | 06/21/2021 | George YANCOPOULOS | REGN-008CIPCON10 |
| | | | CONFIRMATION NO. 5070 |
| 96387 | | POA ACC | EPTANCE LETTER |
| Regeneron - Bozicevic, Fiel
201 REDWOOD SHORES
SUITE 200
REDWOOD CITY, CA 9406 | PARKWAY | | OC000000130128563* |
| | | | Date Mailed: 12/01/2021 |

NOTICE OF ACCEPTANCE OF POWER OF ATTORNEY

This is in response to the Power of Attorney filed 11/29/2021.

The Power of Attorney in this application is accepted. Correspondence in this application will be mailed to the above address as provided by 37 CFR 1.33.

Questions about the contents of this notice and the requirements it sets forth should be directed to the Office of Data Management, Application Assistance Unit, at (571) 272-4000 or (571) 272-4200 or 1-888-786-0101.

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page 1 of 1

| To: | docket@bozpat.com,, |
|----------|--|
| From: | PAIR_eOfficeAction@uspto.gov |
| Cc: | PAIR_eOfficeAction@uspto.gov |
| Subject: | Private PAIR Correspondence Notification for Customer Number 96387 |

Dec 01, 2021 04:06:38 AM

Dear PAIR Customer:

Regeneron - Bozicevic, Field & Francis 201 REDWOOD SHORES PARKWAY SUITE 200 REDWOOD CITY, CA 94065 UNITED STATES

The following USPTO patent application(s) associated with your Customer Number, 96387, have new outgoing correspondence. This correspondence is now available for viewing in Private PAIR.

The official date of notification of the outgoing correspondence will be indicated on the form PTOL-90 accompanying the correspondence.

Disclaimer:

The list of documents shown below is provided as a courtesy and is not part of the official file wrapper. The content of the images shown in PAIR is the official record.

| Application | Document | Mailroom Date | Attorney Docket No. |
|-------------|----------|---------------|---------------------|
| 17352892 | N570 | 12/01/2021 | REGN-008CIPCON10 |

To view your correspondence online or update your email addresses, please visit us anytime at https://sportal.uspto.gov/secure/myportal/privatepair.

If you have any questions, please email the Electronic Business Center (EBC) at EBC@uspto.gov with 'e-Office Action' on the subject line or call 1-866-217-9197 during the following hours:

Monday - Friday 6:00 a.m. to 12:00 a.m.

Thank you for prompt attention to this notice,

UNITED STATES PATENT AND TRADEMARK OFFICE PATENT APPLICATION INFORMATION RETRIEVAL SYSTEM

| Electronically filed | | | |
|--|--|-------------------------|--|
| REPLY UNDER | Attorney Docket No. | REGN-008CIPCON10 | |
| 37 C.F.R. §1.111 | Confirmation No. | 5070 | |
| | First Named Inventor | George D. Yancopoulos | |
| | Application Number 17/352,892 | | |
| | Filing Date | June 21, 2021 | |
| Address to: | Group Art Unit | 1647 | |
| Mail Stop AMENDMENT | Examiner Name | Lockard, Jon McClelland | |
| Commissioner for Patents
P.O. Box 1450
Alexandria, VA 22313-1450 | Title: "Use of a VEGF Antagonist to Treat Angiogenic
Eye Disorders" | | |

Sir:

This reply is responsive to the Office Action dated October 28, 2021, for which a three-month period for response was given, making this response due on January 28, 2022. Accordingly, this response is timely filed.

In view of the attached Terminal Disclaimer and the remarks put forth below, reconsideration and allowance are respectfully requested.

Claims begin on page 2 of this paper.

Remarks/Arguments begin on page 7 of this paper.

CLAIMS

No amendment is being sought in this response. Claims are presented for the Examiner's convenience only.

1. - 20. (Canceled)

21. (Previously Presented) A method of treating an angiogenic eye disorder in a patient in need thereof comprising sequentially administering to the patient by intravitreal injection a single initial dose of 2 mg of aflibercept, followed by one or more secondary doses of 2 mg of aflibercept;

wherein each secondary dose is administered approximately 4 weeks following the immediately preceding dose; and

wherein each tertiary dose is administered approximately 8 weeks following the immediately preceding dose;

wherein the patient achieves a gain in visual acuity within 52 weeks following the initial dose.

22. (Previously Presented) The method of claim 21 wherein the patient achieves a gain in Best Corrected Visual Acuity (BCVA) according to Early Treatment Diabetic Retinopathy Study (ETDRS) letter score.

23. (Previously Presented) The method of claim 22 wherein the patient gains at least 7 letters Best Corrected Visual Acuity (BCVA) according to Early Treatment Diabetic Retinopathy Study (ETDRS) letter score.

24. (Previously Presented) The method of claim 23 wherein the patient achieves the gain in visual acuity within 24 weeks following the initial dose.

25. (Previously Presented) The method of claim 23 wherein only two secondary doses are administered to the patient.

26. (Previously Presented) The method of claim 23 wherein the aflibercept is formulated as an isotonic solution.

27. (Previously Presented) The method of claim 23 wherein the aflibercept is formulated with a nonionic surfactant.

28. (Previously Presented) The method of claim 22 wherein the patient gains at least 8 letters Best Corrected Visual Acuity (BCVA) according to Early Treatment Diabetic Retinopathy Study (ETDRS) letter score.

29. (Previously Presented) The method of claim 28 wherein the patient achieves the gain in visual acuity within 24 weeks following the initial dose.

30. (Previously Presented) The method of claim 22 wherein the patient gains at least 9 letters Best Corrected Visual Acuity (BCVA) according to Early Treatment Diabetic Retinopathy Study (ETDRS) letter score.

31. (Previously Presented) The method of claim 30 wherein only two secondary doses are administered to the patient.

32. (Previously Presented) The method of claim 30 wherein the aflibercept is formulated as an isotonic solution.

33. (Previously Presented) The method of claim 30 wherein the aflibercept is formulated with a nonionic surfactant.

34. (Previously Presented) The method of claim 21 wherein exclusion criteria for the patient include both of:

(1) active ocular inflammation; and

(2) active ocular or periocular infection.

35. (Previously Presented) A method of treating diabetic macular edema in a patient in need thereof comprising sequentially administering to the patient a single initial dose of 2 mg of aflibercept, followed by one or more secondary doses of 2 mg of aflibercept, followed by one or more tertiary doses of 2 mg of aflibercept;

wherein each secondary dose is administered to the patient by intravitreal injection approximately 4 weeks following the immediately preceding dose; and wherein each tertiary dose is administered to the patient by intravitreal injection approximately 8 weeks following the immediately preceding dose.

36. (Previously Presented) The method of claim 35 wherein the patient achieves a gain in visual acuity within 52 weeks following the initial dose.

37. (Previously Presented) The method of claim 36 wherein the patient gains at least 9 letters Best Corrected Visual Acuity (BCVA) according to Early Treatment Diabetic Retinopathy Study (ETDRS) letter score.

38. (Previously Presented) The method of claim 37 wherein the aflibercept is formulated as an isotonic solution.

39. (Previously Presented) The method of claim 37 wherein the aflibercept is formulated with a non- ionic surfactant.

40. (Previously Presented) The method of claim 37 wherein the patient achieves a gain in visual acuity within 24 weeks following the initial dose.

41. (Previously Presented) The method of claim 36 wherein the patient gains at least 8 letters Best Corrected Visual Acuity (BCVA) according to Early Treatment Diabetic Retinopathy Study (ETDRS) letter score.

42. (Previously Presented) The method of claim 41 wherein the aflibercept is formulated as an isotonic solution.

43. (Previously Presented) The method of claim 41 wherein the aflibercept is formulated with a nonionic surfactant.

44. (Previously Presented) The method of claim 35 wherein only two secondary doses are administered to the patient.

45. (Previously Presented) The method of claim 35 wherein four secondary doses are administered to the patient.

46. (Previously Presented) A method of treating age related macular degeneration in a patient in need thereof comprising sequentially administering to the patient a single initial dose of 2 mg of aflibercept, followed by one or more secondary doses of 2 mg of aflibercept, followed by one or more tertiary doses of 2 mg of aflibercept;

wherein each secondary dose is administered to the patient by intravitreal injection approximately 4 weeks following the immediately preceding dose; and wherein each tertiary dose is administered to the patient by intravitreal injection approximately 8 weeks following the immediately preceding dose;

wherein the method is as effective in achieving a gain in visual acuity as monthly administration of 0.5 mg of ranibizumab by intravitreal injection in human subjects with age-related macular degeneration at 52 weeks following the initial dose.

47. (Previously Presented) The method of claim 46 wherein only two secondary doses are administered to the patient.

48. (Previously Presented) The method of claim 46 wherein the gain in visual acuity is measured using the Early Treatment Diabetic Retinopathy Study (ETDRS) letter score.

49. (Previously Presented) A method of treating age-related macular degeneration in a patient in need thereof comprising sequentially administering to the patient a single initial dose of 2 mg of aflibercept, followed by one or more secondary doses of 2 mg of aflibercept, followed by one or more tertiary doses of 2 mg of aflibercept;

Atty Dkt. No.: REGN-008CIPCON10 USSN: 17/352,892

wherein each secondary dose is administered to the patient by intravitreal injection approximately 4 weeks following the immediately preceding dose; and wherein each tertiary dose is administered to the patient by intravitreal injection approximately 8 weeks following the immediately preceding dose;

wherein the method is as effective in maintaining visual acuity as monthly administration of 0.5 mg of ranibizumab by intravitreal injection in human subjects with age-related macular degeneration at 52 weeks following the initial dose.

50. (Previously Presented) The method of claim 49 wherein maintenance of visual acuity means loss of less than 15 letters Best Corrected Visual Acuity (BCVA) as measured by using the Early Treatment Diabetic Retinopathy Study (ETDRS) letter score.

Atty Dkt. No.: REGN-008CIPCON10 USSN: 17/352,892

REMARKS

Formal Matters

Claims 21-50 are pending. Original claims 1-20 were canceled without prejudice. No claims are amended. No claims are added. No New Matter is added.

Statement under 37 C.F.R. §§1.56 and 1.2

Applicants hereby advise the Examiner of the status of a co-pending application in compliance with the Applicant's duty to disclose under 37 C.F.R. §§1.56 and 1.2 (see also MPEP §2001.06(b)) as discussed in *McKesson Info. Soln. Inc., v. Bridge Medical Inc.*, 487 F.3d 897; 82 USPQ2d 1865 (Fed. Cir. 2007).

The Applicant wishes to bring to the Examiner's attention U.S. Patent Application No. 13/940,370, filed July 12, 2013 which issued on February 9, 2016 as U.S. Patent 9,254,338, for which *Inter Partes* Review No. IPR2021-00881 was filed on May 5, 2021, in which a trial was instituted on November 10, 2021.

The Applicant wishes to bring to the Examiner's attention U.S. Patent Application No. 14/972,560, filed December 17, 2015 which issued on June 6, 2017 as U.S. Patent No. 9,669,069, for which *Inter Partes* Review No. IPR2021-00880 was filed on May 5, 2021, in which a trial was instituted on November 10, 2021.

The Applicant wishes to bring to the Examiner's attention U.S. Patent Application No.

15/471,506, filed March 28, 2017 which issued on November 20, 2018 as U.S. Patent No. 10,130,681.

The Applicant wishes to bring to the Examiner's attention U.S. Patent Application No. 16/055,847, filed August 6, 2018 which will issue on December 8, 2020 as U.S. Patent No. 10,857,205.

The Applicant wishes to bring to the Examiner's attention U.S. Patent Application No. 16/159,282, filed October 12, 2018 which issued on November 10, 2020 as U.S. Patent No. 10,828,345, for which Post-Grant Review No. PGR2021-00035 was filed on January 7, 2021, which is now terminated.

The Applicant wishes to bring to the Examiner's attention co-pending U.S. Patent Application No. 16/397,267, filed April 29, 2019, which issued on January 12, 2021 as U.S. Patent No. 10,888,601.

The Applicant wishes to bring to the Examiner's attention co-pending U.S. Patent Application No. 17/072,417, filed October 16, 2020 for which no actions have been mailed.

The Applicant wishes to bring to the Examiner's attention co-pending U.S. Patent Application 17/112,063, filed December 4, 2020 for which no actions have been mailed.

The Applicant wishes to bring to the Examiner's attention co-pending U.S. Patent Application No. 17/112,404 filed December 4, 2020 for which no actions have been mailed.

The Applicant wishes to bring to the Examiner's attention co-pending U.S. Patent Application No. 17/350,958 filed June 17, 2021 for which no actions have been mailed.

These documents are available on PAIR, and thus are not provided with this communication. Please inform the undersigned if there is any difficulty in obtaining the documents from PAIR.

Non-statutory double patenting Rejections

Claims 21-50 were rejected on the ground of nonstatutory obviousness-type double patenting as allegedly unpatentable over claims 1-26 of U.S. Patent No. 9,254,338.

Claims 21-50 were rejected on the ground of nonstatutory obviousness-type double patenting as allegedly unpatentable over claims 1-12 of U.S. Patent No. 9,669,069.

Claims 21-50 were rejected on the ground of nonstatutory obviousness-type double patenting as allegedly unpatentable over claims 1-12 of U.S. Patent No. 10,130,681.

Claims 21-50 were rejected on the ground of nonstatutory obviousness-type double patenting as allegedly unpatentable over claims 1-11 of U.S. Patent No. 10,828,345.

Claims 21-50 were rejected on the ground of nonstatutory obviousness-type double patenting as allegedly unpatentable over claims 1-47 of U.S. Patent No. 10,888,601.

Response

Solely to expedite prosecution and not in acquiescence to the Examiner's rejections, Applicant submits a Terminal Disclaimer herewith with regard to U.S. Patent Nos. 9,254,338; 9,669,069; 10,130,681; 10,828,345; and 10,888,601. Additionally, it is noted that the filing of a terminal disclaimer to obviate a rejection based on non-statutory double patenting is not an admission of the propriety of the rejection. *See, e.g., Quad Environmental Technologies Corp. v. Union Sanitary District*, 946 F.2d 870, (Fed. Cir. 1991) (filing of a terminal disclaimer simply serves the statutory function of removing the rejection of double patenting, and raises neither a presumption nor estoppel on the merits of the rejection); *Motionless Keyboard Co. v. Microsoft Corp.*, 486 F.3d 1376, 1385 (Fed. Cir. 2007) ("A terminal disclaimer is simply not an admission that a later-filed invention is obvious."); *Ortho Pharm. Corp. v. Smith*, 959 F.2d 936, 941 (Fed. Cir. 1992) (holding that filing of terminal disclaimer did not serve as admission of obviousness-type double patenting). Because there are no other rejections outstanding, the application is believed to be in condition for allowance and an indication of such is respectfully requested.

CONCLUSION

Applicants submit that all the claims are in condition for allowance, which action is requested. If the Examiner finds that a telephone conference would expedite the prosecution of this application, the Examiner is invited to telephone the undersigned at the number provided.

The Commissioner is hereby authorized to charge any underpayment of fees up to a strict limit of \$3,000.00 beyond that authorized on the credit card, but not more than \$3,000.00 in additional fees due with any communication for the above referenced patent application, including but not limited to any necessary fees for extensions of time, or credit any overpayment of any amount to Deposit Account No. 50-0815, order number REGN-008CIPCON10.

Respectfully submitted, BOZICEVIC, FIELD & FRANCIS LLP

Date: 7 December 2021

By: <u>/Karl Bozicevic, Reg. No. 28,807/</u> Karl Bozicevic Reg. No. 28,807

BOZICEVIC, FIELD & FRANCIS LLP 201 Redwood Shores Parkway, Suite 200 Redwood City, CA 94065 Telephone: (650) 327-3400 Facsimile: (650) 327-3231

Attached: Terminal Disclaimer

PTO/SB/26 (08-11) Approved for use through 07/31/2012. OMB 0651-0031 U.S. Patent and Trademark Office; U.S. DEPARTMENT OF COMMERCE

| TERMINAL DISCLAIMER TO OBVIATE A DOUBLE PATENTING | Docket Number (Optional) |
|--|---|
| REJECTION OVER A "PRIOR" PATENT | REGN-008CIPCON10 |
| In re Application of: Yancopoulos, George D. | |
| Application No.: 17/352,892 | |
| Filed: June 21, 2021 | |
| For: Use of a VEGF Antagonist to Treat Angiogenic Eye Disorders | |
| The owner, <u>Regeneron Pharmaceuticals, Inc.</u> , of <u>100%</u> percent interest in the instant application here below, the terminal part of the statutory term of any patent granted on the instant application which wou date of the full statutory term of prior patent Nos. <u>9,254,338; 9,669,069; 10,130,681; 10,828,345; and prior patent is presently shortened by any terminal disclaimer. The owner hereby agrees that any pate application shall be enforceable only for and during such period that it and the prior patents are comm with any patent granted on the instant application and is binding upon the grantee, its successors or as</u> | Id extend beyond the expiration
10.888.601 ; as the term of said
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only owned. This agreement runs |
| In making the above disclaimer, the owner does not disclaim the terminal part of the term of any patent that would extend to the expiration date of the full statutory term of the prior patents , "as the term of sa shortened by any terminal disclaimer," in the event that said prior patents later:
expires for failure to pay a maintenance fee;
is held unenforceable; | |
| is found invalid by a court of competent jurisdiction;
is statutorily disclaimed in whole or terminally disclaimed under 37 CFR 1.321;
has all claims canceled by a reexamination certificate;
is reissued; or | |
| is in any manner terminated prior to the expiration of its full statutory term as presently shorte | ened by any terminal disclaimer. |
| Check either box 1 or 2 below, if appropriate. | |
| For submissions on behalf of a business/organization (e.g., corporation, partnership, university
etc.), the undersigned is empowered to act on behalf of the business/organization. | , government agency, |
| I hereby declare that all statements made herein of my own knowledge are true and that all statements mede herein of my own knowledge are true; and further that these statements were made with the knowledge that will made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United Statements may jeopardize the validity of the application or any patent issued thereon. | ful false statements and the like so |
| 2. X The undersigned is an attorney or agent of record. Reg. No. 28,807 | |
| | |
| /Karl Bozicevic, Reg. No. 28,807/ | 7 December 2021 |
| Signature | Date |
| Karl Bozicevic, Reg. No. 28,807 | |
| Typed or printed name | |
| | 650-833-7735 |
| | Telephone Number |
| Terminal disclaimer fee under 37 CFR 1.20(d) included. | |
| WARNING: Information on this form may become public. Credit card infor be included on this form. Provide credit card information and authorization | |
| *Statement under 37 CFR 3.73(b) is required if terminal disclaimer is signed by the assignee (own
Form PTO/SB/96 may be used for making this certification. See MPEP § 324. | ner). |
| is collection of information is required by 37 CFR 1.321. The information is required to obtain or retain a benefit by the
process) an application. Confidentiality is governed by 35 U.S.C. 122 and 37 CFR 1.11 and 1.14. This collection is es
cluding gathering, preparing, and submitting the completed application form to the USPTO. Time will vary depending u | timated to take 12 minutes to complete, |

to process) an application. Confidentiality is governed by 35 U.S.C. 122 and 37 CFR 1.11 and 1.14. This collection is estimated to take 12 minutes to complete, including gathering, preparing, and submitting the completed application form to the USPTO. Time will vary depending upon the individual case. Any comments on the amount of time you require to complete this form and/or suggestions for reducing this burden, should be sent to the Chief Information Officer, U.S. Patent and Trademark Office, U.S. Department of Commerce, P.O. Box 1450, Alexandria, VA 22313-1450. DO NOT SEND FEES OR COMPLETED FORMS TO THIS ADDRESS. **SEND TO: Commissioner for Patents, P.O. Box 1450, Alexandria, VA 22313-1450.**

If you need assistance in completing the form, call 1-800-PTO-9199 and select option 2.

| Electronic Patent Application Fee Transmittal | | | | | | | | |
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| Application Number: | 17352892 | | | | | | | |
| Filing Date: | 21 | -Jun-2021 | | | | | | |
| Title of Invention: | USE OF A VEGF ANTAGONIST TO TREAT ANGIOGENIC EYE DISORDERS | | | | | | | |
| First Named Inventor/Applicant Name: | George YANCOPOULOS | | | | | | | |
| Filer: | Karl Bozicevic/Kimberly Zuehlke | | | | | | | |
| Attorney Docket Number: | RE | GN-008CIPCON10 | | | | | | |
| Filed as Large Entity | | | | | | | | |
| Filing Fees for Utility under 35 USC 111(a) | | | | | | | | |
| Description | | Fee Code | Quantity | Amount | Sub-Total in
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| Basic Filing: | | | | | | | | |
| Pages: | | | | | | | | |
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| Miscellaneous-Filing: | | | | | | | | |
| Petition: | | | | | | | | |
| Patent-Appeals-and-Interference: | | | | | | | | |
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| Extension-of-Time: | | | | | | | | |

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| STATUTORY OR TERMINAL DISCLAIMER | 1814 | 1 | 170 | 170 |
| | Tot | al in USD | (\$) | 170 |
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| Electronic Acl | knowledgement Receipt |
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| EFS ID: | 44456586 |
| Application Number: | 17352892 |
| International Application Number: | |
| Confirmation Number: | 5070 |
| Title of Invention: | USE OF A VEGF ANTAGONIST TO TREAT ANGIOGENIC EYE DISORDERS |
| First Named Inventor/Applicant Name: | George YANCOPOULOS |
| Customer Number: | 96387 |
| Filer: | Karl Bozicevic/Kimberly Zuehlke |
| Filer Authorized By: | Karl Bozicevic |
| Attorney Docket Number: | REGN-008CIPCON10 |
| Receipt Date: | 07-DEC-2021 |
| Filing Date: | 21-JUN-2021 |
| Time Stamp: | 17:20:26 |
| Application Type: | Utility under 35 USC 111(a) |

Payment information:

| Submitted with Payment | yes | | | | | |
|--|------------------|--|--|--|--|--|
| Payment Type | CARD | | | | | |
| Payment was successfully received in RAM | \$170 | | | | | |
| RAM confirmation Number | E2021B7H21203029 | | | | | |
| Deposit Account | | | | | | |
| Authorized User | | | | | | |
| The Director of the USPTO is hereby authorized to charge indicated fees and credit any overpayment as follows: | | | | | | |
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| File Listing: | | | | | | |
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| Document
Number | Document Description | File Name | File Size(Bytes)/
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| | | | 68698 | | | |
| 1 | | REGN-008CIPCON10_2021-12-0
7_Amendment.pdf | ed2e5354535a93c33f95ba4806079d4e306
2f321 | yes | 10 | |
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| | Document De | scription | Start | E | nd | |
| | Amendment/Req. Reconsiderat | 1 | | 1 | | |
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| | Applicant Arguments/Remarks | Made in an Amendment | 7 | 10 | | |
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| | | REGN-008CIPCON10_2021-12-0 | 25017 | | | |
| 2 | Terminal Disclaimer Filed | 7_Terminal_Disclaimer_Prior_P
at.pdf | 530a143c5bc6a23282322d78b31137c9a5c
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| Information: | | | | | | |
| | | | 38249 | | | |
| 3 | Fee Worksheet (SB06) | fee-info.pdf | 7fd79bad8ce4a26fd26ead8b1975d88a559
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| Information: | | | • | | | |
| | | Total Files Size (in bytes) | : 13 | 31964 | | |

This Acknowledgement Receipt evidences receipt on the noted date by the USPTO of the indicated documents, characterized by the applicant, and including page counts, where applicable. It serves as evidence of receipt similar to a Post Card, as described in MPEP 503.

New Applications Under 35 U.S.C. 111

If a new application is being filed and the application includes the necessary components for a filing date (see 37 CFR 1.53(b)-(d) and MPEP 506), a Filing Receipt (37 CFR 1.54) will be issued in due course and the date shown on this Acknowledgement Receipt will establish the filing date of the application.

National Stage of an International Application under 35 U.S.C. 371

If a timely submission to enter the national stage of an international application is compliant with the conditions of 35 U.S.C. 371 and other applicable requirements a Form PCT/DO/EO/903 indicating acceptance of the application as a national stage submission under 35 U.S.C. 371 will be issued in addition to the Filing Receipt, in due course. New International Application Filed with the USPTO as a Receiving Office

If a new international application is being filed and the international application includes the necessary components for an international filing date (see PCT Article 11 and MPEP 1810), a Notification of the International Application Number and of the International Filing Date (Form PCT/RO/105) will be issued in due course, subject to prescriptions concerning national security, and the date shown on this Acknowledgement Receipt will establish the international filing date of the application.

| <i>Application Number</i>
* 17/352,892 * | Application/Contr | ol No. | Applicant(s)/Patent under
Reexamination | | |
|---|-------------------|------------|--|------------|--|
| 177002,002 | 17/352,892 | | YANCOPOULOS, | George | |
| | Examiner | | Art Unit | | |
| | LOCKARD, JON N | ICCLELLAND | 1647 | | |
| Document Code - DISQ | | Internal | Document - D | O NOT MAIL | |

| TERMINAL
DISCLAIMER | ☑ APPROVED | DISAPPROVED |
|---------------------------------|---|-------------|
| Date Filed:
07 December 2021 | This patent is subject
to a Terminal
Disclaimer | |

| Approved/Disapproved by: | |
|---------------------------------|--|
| /LAWANA R HIXON/ | |
| Technology Center: OPLC | |
| Telephone: <u>(571)272-6074</u> | |
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U.S. Patent and Trademark Office TSS-IFW

Terminal Disclaimer

Part of Paper No. 20211208

PTO/SB/06 (09-11) Approved for use through 1/31/2014. OMB 0651-0032

| P/ | ATENT APPLI | CATIO | N FEE | | RMINATION | | Applicatio | nd to a collection of informati
on or Docket Number
17/352,892 | Filing Date
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| | | | (| Column 1 |) | (Column 2) | | | | |
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(37 CFR 1.16(a), (b), c | or (c)) | | N/A | | N/A | | N/A | | |
|]; | SEARCH FEE
(37 CFR 1.16(k), (i), or | . (m)) | | N/A | | N/A | | N/A | | |
| | EXAMINATION FEE
(37 CFR 1.16(o), (p), c | | | N/A | | N/A | | N/A | | |
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This collection of information is required by 37 CFR 1.16. The information is required to obtain or retain a benefit by the public which is to file (and by the USPTO to process) an application. Confidentiality is governed by 35 U.S.C. 122 and 37 CFR 1.14. This collection is estimated to take 12 minutes to complete, including gathering, preparing, and submitting the completed application form to the USPTO. Time will vary depending upon the individual case. Any comments on the amount of time you require to complete this form and/or suggestions for reducing this burden, should be sent to the Chief Information Officer, U.S. Patent and Trademark Office, U.S. Department of Commerce, P.O. Box 1450, Alexandria, VA 22313-1450. DO NOT SEND FEES OR COMPLETED FORMS TO THIS

ADDRESS. SEND TO: Commissioner for Patents, P.O. Box 1450, Alexandria, VA 22313-1450.

If you need assistance in completing the form, call 1-800-PTO-9199 and select option 2.

| INFORMATION DISCLOSURE
STATEMENT BY APPLICANT | | Application Number
Filing Date
First Named Inventor
Art Unit
Examiner Name | | 17/352,892
June 21, 2021
George D. YANCOPOULOS
1647
Jon McClelland Lockard | | | | | |
|--|------|--|------|--|------------------------|---------------------------|------|---|--|
| Sheet | | 1 | of | 1 | Attorney Docket Number | | REGN | REGN-008CIPCON10 | |
| | | | | U.S. F | ATENT D | OCUMENTS | | | |
| Examiner | Cite | Patent Numb | er | Issue | Date | Name of Patente | e or | Pages, Columns, Lines, Where | |
| Initial* | No. | Number-Kind Code (if kr | own) | YYYY- | MM-DD | Applicant of Cited Docume | | Relevant Passages or Relevant
Figures Appear | |
| | 1 | 6897294 | | 2005-05 | 5-24 | Davis-Smyth e | tal. | | |

| | U.S. PATENT APPLICATION PUBLICATIONS | | | | | | | | |
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| Examiner | Cite | Publication Number | Publication Date | Name of Patentee or | Pages, Columns, Lines, Where | | | | |
| Initial* | No. | | YYYY-MM-DD | Applicant of Cited Document | Relevant Passages or Relevant | | | | |
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| | FOREIGN PATENT DOCUMENTS | | | | | | | | |
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| Examiner
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Country Code-Number-Kind Code (<i>if</i> known) | Publication Date
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Applicant of Cited Document | Pages, Columns, Lines,
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| | | NON PATENT LITERATURE DOCUMENTS | |
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er
Initials* | No. | Include name of the author (in CAPITAL LETTERS), title of the article (when appropriate), title of the item (book, magazine, journal, serial, symposium, catalog, etc.), date, page(s), volume-issue number(s), publisher, city and/or country where published. | Т |
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| Examiner
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*EXAMINER: Initial if reference considered, whether or not citation is in conformance with MPEP 609. Draw line through citation if not in conformance and not considered. Include copy of this form with next communication to applicant.

| Electronic Patent / | Application Fee Transmittal | | | | |
|---|--|----------------------|-----------|--------|-------------------------|
| Application Number: | 17 | 352892 | | | |
| Filing Date: | 21 | -Jun-2021 | | | |
| Title of Invention: | USE OF A VEGF ANTAGONIST TO TREAT ANGIOGENIC EYE DISORDERS | | | | |
| First Named Inventor/Applicant Name: | George YANCOPOULOS | | | | |
| Filer: | Ка | rl Bozicevic/Kimberl | y Zuehlke | | |
| Attorney Docket Number: | RE | GN-008CIPCON10 | | | |
| Filed as Large Entity | | | | | |
| Filing Fees for Utility under 35 USC 111(a) | | | | | |
| Description | | Fee Code | Quantity | Amount | Sub-Total in
USD(\$) |
| Basic Filing: | | | | | |
| Pages: | | | | | |
| Claims: | | | | | |
| Miscellaneous-Filing: | | | | | |
| Petition: | | | | | |
| Patent-Appeals-and-Interference: | | | | | |
| Post-Allowance-and-Post-Issuance: | | | | | |
| Extension-of-Time: | | | | | |

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| Electronic Acl | knowledgement Receipt |
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| EFS ID: | 44540302 |
| Application Number: | 17352892 |
| International Application Number: | |
| Confirmation Number: | 5070 |
| Title of Invention: | USE OF A VEGF ANTAGONIST TO TREAT ANGIOGENIC EYE DISORDERS |
| First Named Inventor/Applicant Name: | George YANCOPOULOS |
| Customer Number: | 96387 |
| Filer: | Karl Bozicevic/Kimberly Zuehlke |
| Filer Authorized By: | Karl Bozicevic |
| Attorney Docket Number: | REGN-008CIPCON10 |
| Receipt Date: | 16-DEC-2021 |
| Filing Date: | 21-JUN-2021 |
| Time Stamp: | 17:36:43 |
| Application Type: | Utility under 35 USC 111(a) |

Payment information:

| Submitted with Payment | yes |
|--|---|
| Payment Type | CARD |
| Payment was successfully received in RAM | \$260 |
| RAM confirmation Number | E2021BFH37171715 |
| Deposit Account | |
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| The Director of the USPTO is hereby authorized to charge | e indicated fees and credit any overpayment as follows: |
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| File Listin | g: | | | | |
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| Document
Number | Document Description | File Name | File Size(Bytes)/
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Part /.zip | Pages
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| | | | 50042 | | |
| 1 | Transmittal Letter | REGN-008CIPCON10_2021-12-1
6_SuppIDS_Trans.pdf | 3932d0d3aedccce743f7dd65adf48c2001d
62041 | no | 2 |
| Warnings: | | | · · · · · · · · · · · · · · · · · · · | | |
| Information: | | | | | |
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| 2 | Information Disclosure Statement (IDS)
Form (SB08) | REGN-008CIPCON10_2021-12-1
6_SuppIDS_SB08A.pdf | d208a68d729acc2bbb65373a6a8ebc1211f
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| This is not an U | SPTO supplied IDS fillable form | | | | |
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| 3 | Fee Worksheet (SB06) | fee-info.pdf | ef41e3f587ac020e1a8dd8cf5d2de0b4a8e2
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| | Attorney Docket No. | REGN-008CIPCON10 |
| | Confirmation No. | 5070 |
| SUPPLEMENTAL INFORMATION | First Named Inventor | George D. Yancopoulos |
| DISCLOSURE STATEMENT | Application Number | 17/352,892 |
| | Filing Date | June 21, 2021 |
| | Group Art Unit | 1647 |
| Address to: | Examiner Name | Jon McClelland Lockard |
| Commissioner for Patents
P.O. Box 1450
Alexandria, VA 22313-1450 | Title: "Use of a VEGF
Eye Disorders" | Antagonist to Treat Angiogenic |

Sir:

Applicant submits herewith documents which may be material to the examination of this application and in respect of which there may be a duty to disclose in accordance with 37 C.F.R. § 1.56. This submission is not intended to constitute an admission that any document referred to therein is "prior art" for this invention unless specifically designated as such. A listing of the documents is shown on enclosed Form PTO/SB/08A.

The publications discussed herein are provided to comply with the duty to disclose in accordance with 37 C.F.R. § 1.56. However, nothing herein is to be construed as an admission that the present invention is not entitled to antedate such publication by virtue of prior invention. Further, the dates of publication provided may be different from the actual publication dates which may need to be independently confirmed

The Examiner is requested to make the documents listed on the enclosed PTO/SB/08A of record in this application. Applicants would appreciate the Examiner initialing and returning the initialed copy of form PTO/SB/08A, indicating the documents cited therein have been considered and made of record herein.

Statements

No statement

PTA Statement under 37 CFR § 1.704(d)(1): Each item of information contained in the information disclosure statement filed herewith:

(i) Was first cited in any communication from a patent office in a counterpart foreign or international application or from the Office, and this communication was not received by any individual designated in § 1.56(c) more than thirty days prior to the filing of the information disclosure statement; or

(ii) Is a communication that was issued by a patent office in a counterpart foreign or international application or by the Office, and this communication was not received by

any individual designated in § 1.56(c) more than thirty days prior to the filing of the information disclosure statement.

IDS Statement under 37 CFR § 1.97(e)(1): Each item of information contained in the information disclosure statement was first cited in any communication from a foreign patent office in a counterpart foreign application not more than three months prior to the filing of the information disclosure statement; or

IDS Statement under 37 CFR § 1.97(e)(2): No item of information contained in the information disclosure statement was cited in a communication from a foreign patent office in a counterpart foreign application, and, to the knowledge of the person signing the certification after making reasonable inquiry, no item of information contained in the information disclosure statement was known to any individual designated in § 1.56(c) more than three months prior to the filing of the information disclosure statement.

Fees

No fee is believed to be due.

The appropriate fee set forth in 37 C.F.R. §1.17(p) accompanies this information disclosure statement.

The Commissioner is hereby authorized to charge any underpayment of fees up to a strict limit of \$3,000.00 beyond that authorized on the credit card, but not more than \$3,000.00 in additional fees due with any communication for the above-referenced patent application, including but not limited to any necessary fees for extensions of time, or credit any overpayment of any amount to Deposit Account No. 50-0815, order number REGN-008CIPCON10.

Respectfully submitted, BOZICEVIC, FIELD & FRANCIS LLP

Date: <u>16 December 2021</u>

By: <u>/Karl Bozicevic, Reg. No. 28,807/</u> Karl Bozicevic Reg. No. 28,807

BOZICEVIC, FIELD & FRANCIS LLP 201 Redwood Shores Parkway, Suite 200 Redwood City, CA 94065 Telephone: (650) 327-3400 Facsimile: (650) 327-3231

> Celltrion Exhibit 1014 Page 1333

UNITED STATES PATENT AND TRADEMARK OFFICE



UNITED STATES DEPARTMENT OF COMMERCE United States Patent and Trademark Office Address: COMMISSIONER FOR PATENTS P.O. Box 1450 Alexandria, Virginia 22313-1450 www.uspto.gov

NOTICE OF ALLOWANCE AND FEE(S) DUE

96387 7590 12/22/2021 Regeneron - Bozicevic, Field & Francis 201 REDWOOD SHORES PARKWAY SUITE 200 REDWOOD CITY, CA 94065

| EXAMINER | | | | |
|--------------|--------------|--|--|--|
| LOCKARD, JON | MCCLELLAND | | | |
| ART UNIT | PAPER NUMBER | | | |
| 1647 | | | | |

DATE MAILED: 12/22/2021

| APPLICATION NO. | FILING DATE | FIRST NAMED INVENTOR | ATTORNEY DOCKET NO. | CONFIRMATION NO. |
|-----------------|-------------|----------------------|---------------------|------------------|
| 17/352,892 | 06/21/2021 | George YANCOPOULOS | REGN-008CIPCON10 | 5070 |

TITLE OF INVENTION: USE OF A VEGF ANTAGONIST TO TREAT ANGIOGENIC EYE DISORDERS

| APPLN. TYPE | ENTITY STATUS | ISSUE FEE DUE | PUBLICATION FEE DUE | PREV. PAID ISSUE FEE | TOTAL FEE(S) DUE | DATE DUE |
|----------------|---------------|---------------|---------------------|----------------------|------------------|------------|
| nonprovisional | UNDISCOUNTED | \$1200 | \$0.00 | \$0.00 | \$1200 | 03/22/2022 |

THE APPLICATION IDENTIFIED ABOVE HAS BEEN EXAMINED AND IS ALLOWED FOR ISSUANCE AS A PATENT. <u>PROSECUTION ON THE MERITS IS CLOSED</u>. THIS NOTICE OF ALLOWANCE IS NOT A GRANT OF PATENT RIGHTS. THIS APPLICATION IS SUBJECT TO WITHDRAWAL FROM ISSUE AT THE INITIATIVE OF THE OFFICE OR UPON PETITION BY THE APPLICANT. SEE 37 CFR 1.313 AND MPEP 1308.

THE ISSUE FEE AND PUBLICATION FEE (IF REQUIRED) MUST BE PAID WITHIN <u>THREE MONTHS</u> FROM THE MAILING DATE OF THIS NOTICE OR THIS APPLICATION SHALL BE REGARDED AS ABANDONED. <u>THIS STATUTORY PERIOD</u> <u>CANNOT BE EXTENDED</u>. SEE 35 U.S.C. 151. THE ISSUE FEE DUE INDICATED ABOVE DOES NOT REFLECT A CREDIT FOR ANY PREVIOUSLY PAID ISSUE FEE IN THIS APPLICATION. IF AN ISSUE FEE HAS PREVIOUSLY BEEN PAID IN THIS APPLICATION (AS SHOWN ABOVE), THE RETURN OF PART B OF THIS FORM WILL BE CONSIDERED A REQUEST TO REAPPLY THE PREVIOUSLY PAID ISSUE FEE TOWARD THE ISSUE FEE NOW DUE.

HOW TO REPLY TO THIS NOTICE:

I. Review the ENTITY STATUS shown above. If the ENTITY STATUS is shown as SMALL or MICRO, verify whether entitlement to that entity status still applies.

If the ENTITY STATUS is the same as shown above, pay the TOTAL FEE(S) DUE shown above.

If the ENTITY STATUS is changed from that shown above, on PART B - FEE(S) TRANSMITTAL, complete section number 5 titled "Change in Entity Status (from status indicated above)".

For purposes of this notice, small entity fees are 1/2 the amount of undiscounted fees, and micro entity fees are 1/2 the amount of small entity fees.

II. PART B - FEE(S) TRANSMITTAL, or its equivalent, must be completed and returned to the United States Patent and Trademark Office (USPTO) with your ISSUE FEE and PUBLICATION FEE (if required). If you are charging the fee(s) to your deposit account, section "4b" of Part B - Fee(s) Transmittal should be completed. If an equivalent of Part B is filed, a request to reapply a previously paid issue fee must be clearly made, and delays in processing may occur due to the difficulty in recognizing the paper as an equivalent of Part B.

III. All communications regarding this application must give the application number. Please direct all communications prior to issuance to Mail Stop ISSUE FEE unless advised to the contrary.

IMPORTANT REMINDER: Maintenance fees are due in utility patents issuing on applications filed on or after Dec. 12, 1980. It is patentee's responsibility to ensure timely payment of maintenance fees when due. More information is available at www.uspto.gov/PatentMaintenanceFees.

PTOL-85 (Rev. 02/11)

PART B - FEE(S) TRANSMITTAL

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Alexandria, Virgin | Patents | | | By t | fax, send to | : (571)-273-2885 |
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| APPLICATION NO. | FILING DATE | | FIRST NAMED INVEN | FOR | ATTORNEY DO | CKET NO. | CONFIRMATION NO. |
| 17/352,892 | 06/21/2021 | • | George YANCOPOUI | .OS | REGN-008CI | PCON10 | 5070 |
| TITLE OF INVENTION | I: USE OF A VEGF ANT | AGONIST TO TREAT | ANGIOGENIC EYE D | ISORDERS | | | |
| APPLN. TYPE | ENTITY STATUS | ISSUE FEE DUE | PUBLICATION FEE D | UE PREV. PAID ISSU | E FEE TOTAL | FEE(S) DUE | DATE DUE |
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| EXAM | AINER | ART UNIT | CLASS-SUBCLASS | | | | |
| LOCKARD, JON | MCCLELLAND | 1647 | 424-134100 | | | | |
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PTOL-85 Part B (08-18) Approved for use through 01/31/2020

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Determination of Patent Term Adjustment under 35 U.S.C. 154 (b) (Applications filed on or after May 29, 2000)

The Office has discontinued providing a Patent Term Adjustment (PTA) calculation with the Notice of Allowance.

Section 1(h)(2) of the AIA Technical Corrections Act amended 35 U.S.C. 154(b)(3)(B)(i) to eliminate the requirement that the Office provide a patent term adjustment determination with the notice of allowance. See Revisions to Patent Term Adjustment, 78 Fed. Reg. 19416, 19417 (Apr. 1, 2013). Therefore, the Office is no longer providing an initial patent term adjustment determination with the notice of allowance. The Office will continue to provide a patent term adjustment determination Letter that is mailed to applicant approximately three weeks prior to the issue date of the patent, and will include the patent term adjustment on the patent. Any request for reconsideration of the patent term adjustment determination (or reinstatement of patent term adjustment) should follow the process outlined in 37 CFR 1.705.

Any questions regarding the Patent Term Extension or Adjustment determination should be directed to the Office of Patent Legal Administration at (571)-272-7702. Questions relating to issue and publication fee payments should be directed to the Customer Service Center of the Office of Patent Publication at 1-(888)-786-0101 or (571)-272-4200.

OMB Clearance and PRA Burden Statement for PTOL-85 Part B

The Paperwork Reduction Act (PRA) of 1995 requires Federal agencies to obtain Office of Management and Budget approval before requesting most types of information from the public. When OMB approves an agency request to collect information from the public, OMB (i) provides a valid OMB Control Number and expiration date for the agency to display on the instrument that will be used to collect the information and (ii) requires the agency to inform the public about the OMB Control Number's legal significance in accordance with 5 CFR 1320.5(b).

The information collected by PTOL-85 Part B is required by 37 CFR 1.311. The information is required to obtain or retain a benefit by the public which is to file (and by the USPTO to process) an application. Confidentiality is governed by 35 U.S.C. 122 and 37 CFR 1.14. This collection is estimated to take 30 minutes to complete, including gathering, preparing, and submitting the completed application form to the USPTO. Time will vary depending upon the individual case. Any comments on the amount of time you require to complete this form and/or suggestions for reducing this burden, should be sent to the Chief Information Officer, U.S. Patent and Trademark Office, U.S. Department of Commerce, P.O. Box 1450, Alexandria, Virginia 22313-1450. DO NOT SEND FEES OR COMPLETED FORMS TO THIS ADDRESS. SEND TO: Commissioner for Patents, P.O. Box 1450, Alexandria, Virginia 22313-1450. Under the Paperwork Reduction Act of 1995, no persons are required to respond to a collection of information unless it displays a valid OMB control number.

Privacy Act Statement

The Privacy Act of 1974 (P.L. 93-579) requires that you be given certain information in connection with your submission of the attached form related to a patent application or patent. Accordingly, pursuant to the requirements of the Act, please be advised that: (1) the general authority for the collection of this information is 35 U.S.C. 2(b) (2); (2) furnishing of the information solicited is voluntary; and (3) the principal purpose for which the information is used by the U.S. Patent and Trademark Office is to process and/or examine your submission related to a patent application or patent. If you do not furnish the requested information, the U.S. Patent and Trademark Office may not be able to process and/or examine your submission, which may result in termination of proceedings or abandonment of the application or expiration of the patent.

The information provided by you in this form will be subject to the following routine uses:

- 1. The information on this form will be treated confidentially to the extent allowed under the Freedom of Information Act (5 U.S.C. 552) and the Privacy Act (5 U.S.C 552a). Records from this system of records may be disclosed to the Department of Justice to determine whether disclosure of these records is required by the Freedom of Information Act.
- 2. A record from this system of records may be disclosed, as a routine use, in the course of presenting evidence to a court, magistrate, or administrative tribunal, including disclosures to opposing counsel in the course of settlement negotiations.
- 3. A record in this system of records may be disclosed, as a routine use, to a Member of Congress submitting a request involving an individual, to whom the record pertains, when the individual has requested assistance from the Member with respect to the subject matter of the record.
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- 5. A record related to an International Application filed under the Patent Cooperation Treaty in this system of records may be disclosed, as a routine use, to the International Bureau of the World Intellectual Property Organization, pursuant to the Patent Cooperation Treaty.
- 6. A record in this system of records may be disclosed, as a routine use, to another federal agency for purposes of National Security review (35 U.S.C. 181) and for review pursuant to the Atomic Energy Act (42 U.S.C. 218(c)).
- 7. A record from this system of records may be disclosed, as a routine use, to the Administrator, General Services, or his/her designee, during an inspection of records conducted by GSA as part of that agency's responsibility to recommend improvements in records management practices and programs, under authority of 44 U.S.C. 2904 and 2906. Such disclosure shall be made in accordance with the GSA regulations governing inspection of records for this purpose, and any other relevant (i.e., GSA or Commerce) directive. Such disclosure shall not be used to make determinations about individuals.
- 8. A record from this system of records may be disclosed, as a routine use, to the public after either publication of the application pursuant to 35 U.S.C. 122(b) or issuance of a patent pursuant to 35 U.S.C. 151. Further, a record may be disclosed, subject to the limitations of 37 CFR 1.14, as a routine use, to the public if the record was filed in an application which became abandoned or in which the proceedings were terminated and which application is referenced by either a published application, an application open to public inspection or an issued patent.
- 9. A record from this system of records may be disclosed, as a routine use, to a Federal, State, or local law enforcement agency, if the USPTO becomes aware of a violation or potential violation of law or regulation.

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JON M LOCK | ARD | Art Unit
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All claims being allowable, PROSECUTION ON THE MERITS IS (OR REMAINS) CLOSED in this application. If not included
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NOTICE OF ALLOWABILITY IS NOT A GRANT OF PATENT RIGHTS. This application is subject to withdrawal from issue at the initiative
of the Office or upon petition by the applicant. See 37 CFR 1.313 and MPEP 1308.
1. This communication is responsive to the Response filed 07 December 2021. | | | | | | | | | | |
| A declaration(s)/affidavit(s) under 37 CFR 1.130(b) was | | | | | | | | | | |
| 2. An election was made by the applicant in response to a res
restriction requirement and election have been incorporated | | | he interview o | n; the | | | | | | |
| 3. ✓ The allowed claim(s) is/are 21-50 (renumbered as claims 1-30, respectively). As a result of the allowed claim(s), you may be eligible to benefit from the Patent Prosecution Highway program at a participating intellectual property office for the corresponding application. For more information, please see http://www.uspto.gov/patents/init_events/pph/index.jsp or send an inquiry to PPHfeedback@uspto.gov. | | | | | | | | | | |
| 4. Acknowledgment is made of a claim for foreign priority under | er 35 U.S.C. § 1 | 19(a)-(d) or (f). | | | | | | | | |
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| 6. DEPOSIT OF and/or INFORMATION about the deposit of E attached Examiner's comment regarding REQUIREMENT F | | | | | | | | | | |
| Attachment(s) 1. Notice of References Cited (PTO-892) 2. ✓ Information Disclosure Statements (PTO/SB/08),
Paper No./Mail Date 3. Examiner's Comment Regarding Requirement for Deposit
of Biological Material 4. Interview Summary (PTO-413), | 6. [| Examiner's Amend Examiner's Statem Other | | | | | | | | |
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Examiner, Art Unit 1647 | | RISTINE J SAOU
nary Examiner, Art | | | | | | | | |
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| U.S. Patent and Trademark Office
PTOL-37 (Rev. 08-13) Notice | of Allowability | Pa | rt of Paper No./I | Mail Date 20211215 | | | | | | |

Notice of Pre-AIA or AIA Status

1. The present application is being examined under the pre-AIA first to invent provisions.

Terminal Disclaimer

 The terminal disclaimer filed on 07 December 2021 disclaiming the terminal portion of any patent granted on this application which would extend beyond the expiration date of U.S.
 Patent No. 9,254,338, U.S. Patent No. 9,669,069, U.S. Patent No. 10,130,681, U.S. Patent No. 10,828,345 and U.S. Patent No. 10,888,601 has been reviewed and is accepted. The terminal disclaimer has been recorded.

Information Disclosure Statement

3. The information disclosure statements (IDS) submitted on 24 November 2021 and 16 December 2021 have been considered by the examiner.

Withdrawn Objections and/or Rejections

4. The rejection of claims 21-50 on the ground of nonstatutory obviousness-type double patenting as set forth at pp. 2-6 of the previous Office action (mailed 28 October 2021) is withdrawn in view of Applicant's submission of a terminal disclaimer (filed 07 December 2021).

Summary

5. Claims 21-50 are allowed.

Advisory Information

Any inquiry concerning this communication or earlier communications from the examiner should be directed to **Jon M. Lockard** whose telephone number is (**571**) **272-2717**. The examiner can normally be reached on Monday through Friday, 8:00 AM to 4:30 PM.

If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, **Joanne Hama**, can be reached on (571) 272-2911. The fax number for the organization where this application or proceeding is assigned is 571-273-8300.

Information regarding the status of an application may be obtained from the Patent Application Information Retrieval (PAIR) system. Status information for published applications may be obtained from either Private PAIR or Public PAIR. Status information for unpublished applications is available through Private PAIR only. For more information about the PAIR system, see http://pair-direct.uspto.gov. Should you have questions on access to the Private PAIR system, contact the Electronic Business Center (EBC) at 866-217-9197 (toll-free). If you would like assistance from a USPTO Customer Service Representative or access to the automated information system, call 800-786-9199 (IN USA OR CANADA) or 571-272-1000.

> /Christine J Saoud/ Primary Examiner, Art Unit 1647

/J.L/ Examiner, Art Unit 1647 December 16, 2021

| | Application/Control No. | Applicant(s)/Patent Under Reexamination | | | | |
|----------------------|-------------------------|---|--|--|--|--|
| Issue Classification | 17/352,892 | YANCOPOULOS, George | | | | |
| | Examiner | Art Unit | | | | |
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| CPC | | | | | | | | | |
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| Symbol | | | Туре | Version | | | | | |
| A61K | / 38 | / 179 | F | 2013-01-01 | | | | | |
| C07K | / 16 | / 22 | I | 2013-01-01 | | | | | |
| C07K | / 14 | / 71 | I | 2013-01-01 | | | | | |
| A61K | / 9 | / 0048 | I | 2013-01-01 | | | | | |
| A61K | / 2039 | / 505 | А | 2013-01-01 | | | | | |
| C07K | / 2319 | / 30 | А | 2013-01-01 | | | | | |
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| /JON M LOCKARD/
Examiner, Art Unit 1647 | 16 December 2021 | Total Claims Allowed: | | | | |
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| (Assistant Examiner) | (Date) | 30 |) | | | |
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Primary Examiner, Art Unit 1647 | 17 December 2021 | O.G. Print Claim(s) | O.G. Print Figure | | | |
| (Primary Examiner) | (Date) | 1 | NONE | | | |
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Page 1 of 3

| | Application/Control No. | Applicant(s)/Patent Under Reexamination |
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| Issue Classification | 17/352,892 | YANCOPOULOS, George |
| | Examiner | Art Unit |
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| INTERNATIONAL CLASSIFICATION | | | | | | | |
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| CLAIMED | | | | | | | |
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| US ORIGINAL CLASSIFICATION | | | | | | | | |
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| CLASS | | | SUBCLASS | | | | | |
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| CROSS REFERENC | ES(S) | | | | | | | |
| CLASS | | SUBCLASS (ONE SUBCLASS PER BLOCK) | | | | | | |
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| /JON M LOCKARD/
Examiner, Art Unit 1647 | 16 December 2021 | Total Claim | s Allowed: |
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| (Assistant Examiner) | (Date) | 30 |) |
| /CHRISTINE J SAOUD/
Primary Examiner, Art Unit 1647 | 17 December 2021 | O.G. Print Claim(s) | O.G. Print Figure |
| (Primary Examiner) | (Date) | 1 | NONE |
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| | Application/Control No. | Applicant(s)/Patent Under Reexamination |
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| Issue Classification | 17/352,892 | YANCOPOULOS, George |
| | Examiner | Art Unit |
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| | Claims r | enumb | ered in t | the san | ne orde | r as pre | sented | by app | licant | 🗆 C | PA (| J T.D | . 🗆 | R.1.47 | 7 |
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Examiner, Art Unit 1647 | 16 December 2021 | Total Claims Allowed: | |
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Page 3 of 3

| | Application/Control No. | Applicant(s)/Patent Under Reexamination |
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| Search Notes | 17/352,892 | YANCOPOULOS, George |
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| | JON M LOCKARD | 1647 |

| CPC - Searched* | | |
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| CPC Combination Sets - Searched* | | | | | |
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| US Classification - Searched* | | | | | | |
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| Class | Subclass Date Examiner | | | | | |
| NONE | | 10/23/2021 | JML | | | |

* See search history printout included with this form or the SEARCH NOTES box below to determine the scope of the search.

| Search Notes | | | | | | | |
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| Search Notes | Date | Examiner | | | | | |
| EAST (USPAT, US-PGPUB, EPO, DERWENT): See attached search history. | 10/23/2021 | JML | | | | | |
| STN (MEDLINE, SCISEARCH, EMBASE, BIOSIS): See attached search history. | 10/23/2021 | JML | | | | | |
| PALM: Inventor search. | 10/23/2021 | JML | | | | | |

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| Search Notes | 17/352,892 | YANCOPOULOS, George | |
| | Examiner | Art Unit | |
| | JON M LOCKARD | 1647 | |

| Interference Search | | | | | | | |
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| US Class/CPC
Symbol | US Subclass/CPC Group | Date | Examiner | | | | |
| | EAST (USPAT, US-PGPUB, EPO, DERWENT): See attached search history. | 12/16/2021 | JML | | | | |
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| INFORMATION DISCLOSURE
STATEMENT BY APPLICANT | | Filing Dat | ed Inventor | 17/352,892 June 21, 2021 George D. YANCOPOULOS 1647 Jon McClelland Lockard | | | | |
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| Sheet | 1 of 1 Atte | | Attorney I | Docket Number | REGN-008CIPCON10 | | | |
| | | | | U.S. F | PATENT C | OCUMENTS | | |
| | | Date | Name of Patentee | | Pages, Columns, Lines, Where | | | |
| Initial* | No. | Number-Kind Code (if kn | own) | ¥ ¥ ¥ ¥ 4 | /M-DD Applicant of Cited Docume | | cument | Relevant Passages or Relevant
Figures Appear |
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Signature | /JON M LOCKARD/ | Date
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*EXAMINER: Initial if reference considered, whether or not citation is in conformance with MPEP 609. Draw line through citation if not in conformance and not considered. Include copy of this form with next communication to applicant.

17/352,892

EAST Search History

EAST Search History (Interference)

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| Ref
| Hits | Search Query | DBs | Default
Operator | Plurals | Time Stamp |
|----------|-------|---|-------|---------------------|---------|---------------------|
| L1 | 2,967 | (flt1 or vegfr1 or (vegf adj r1)) same ((flk1
or kdr or vegfr2 or (vegf adj r2)) or (Flt4
vegfr3 or (vegf adj r3))) | USPAT | OR | ON | 2021/12/16
11:32 |
| L2 | 301 | l1 same ((chimer\$ or fusion) same vegf) | USPAT | OR | ON | 2021/12/16
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| L3 | 2,906 | (flt1 or vegfr1 or (vegf adj r1)) with ((flk1 or kdr or vegfr2 or (vegf adj r2)) or (Flt4 vegfr3 or (vegf adj r3))) | USPAT | OR | ON | 2021/12/16
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| L4 | 159 | I3 with ((chimer\$ or fusion) with vegf) | USPAT | OR | ON | 2021/12/16
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| L5 | 1,906 | aflibercept zaltrap eylea | USPAT | OR | ON | 2021/12/16
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| L6 | 97 | (I2 I4 I5) same ((eye or ocular or retina\$ or
macular) with disorder) | USPAT | OR | ON | 2021/12/16
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| L12 | 1,872 | aflibercept zaltrap eylea.clm. | USPAT | OR | ON | 2021/12/16
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| L13 | 86 | 112 and ((eye or ocular or retina\$ or
macular) with disorder).clm. | USPAT | OR | ON | 2021/12/16
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| | ATTY. DOCKET NO. | APPLICATION NO. | |
| | REGN-008CIPCON10 | 17/352,892 | |
| SUBSTITUTE 1449 | APPLICANT | | |
| INFORMATION DISCLOSURE STATEMENT | REGENERON PHARMACEUTICALS | , INC. | |
| | FILING DATE | GROUP | |
| | June 21, 2021 | 1647 | |

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*Copies of the listed references are either submitted herewith or were previously cited by or submitted to, the Office in a prior application. Pursuant to 37 C.F.R. § 1.97(d) and MPEP §609, the indicated reference may have been previously cited by or submitted to, the Office in a prior application, where the prior application is identified by its U.S. Application Number in this Information Disclosure Statement.

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| INFORMATION DISCLOSURE STATEMENT | REGENERON PHARMACEUTICALS, INC. | | |
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| *Copies of the listed references are either submitted herewith or were previously cited by or submitted to, the Office 1.97(d) and MPEP §609, the indicated reference may have been previously cited by or submitted to, the Office in a identified by its U.S. Application Number in this Information Disclosure Statement. | |

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| | REGN-008CIPCON10 | 17/352,892 |
| | APPLICANT | |
| | REGENERON PHARMACEUTICALS, INC. | |
| | FILING DATE | GROUP |
| | June 21, 2021 | 1647 |

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| EXAMINER | DATE CONSIDERED | |
|---|--|--|
| EXAMINER: Initial if reference considered, whether or not citation is in conformance with MPEP 609. Draw line through citation if not in conformance and not considered. Include copy of this form with next communication to Applicant. | | |
| *Copies of the listed references are either submitted herewith or were previously cited by or submitted to, the Office in | a prior application. Pursuant to 37 C.F.R. § | |

1.97(d) and MPEP \$609, the indicated reference may have been previously cited by or submitted to, the Office in a prior application, where the prior application is identified by its U.S. Application Number in this Information Disclosure Statement.

| | | Page 11 of 12 |
|---|---------------------------------|-----------------|
| SUBSTITUTE 1449
INFORMATION DISCLOSURE STATEMENT | ATTY. DOCKET NO. | APPLICATION NO. |
| | REGN-008CIPCON10 | 17/352,892 |
| | APPLICANT | |
| | REGENERON PHARMACEUTICALS, INC. | |
| | FILING DATE | GROUP |
| | June 21, 2021 | 1647 |

| 1 | NON-PATENT LITERATURE DOCUMENTS - UPDATES TO PREVIOUS IDS CITATI | ONS |
|------|---|--|
| | DOCUMENT (Including Author, Title, Date, Pertinent Pages, etc.) | REFERENCE PROVIDED* |
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17/072,417 |

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| | | Page 12 of 12 |
|---|---------------------------------|-----------------|
| SUBSTITUTE 1449
INFORMATION DISCLOSURE STATEMENT | ATTY. DOCKET NO. | APPLICATION NO. |
| | REGN-008CIPCON10 | 17/352,892 |
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17/072,417 |

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| JON M LOCKARD/ | 12/15/2021 |
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Inventor Information for 17/352892

/J.L./

| Inventor Name | City | State/Country | | | | |
|---|--|-----------------------------|--|--|--|--|
| YANCOPOULOS, GEORGE | YORKTOWN HEIGHTS | NEW YORK | | | | |
| Apple life Contents Petition Info Atty/Agent Info Con | tinuity Data Foreign Data Inventors Applicants Add | vess Fees Post Info Pre Gr. | | | | |
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| From: | PAIR_eOfficeAction@uspto.gov |
| Cc: | PAIR_eOfficeAction@uspto.gov |
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Dec 22, 2021 04:10:17 AM

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| Application | Document | Mailroom Date | Attorney Docket No. |
|-------------|----------|---------------|---------------------|
| 17352892 | NOA | 12/22/2021 | REGN-008CIPCON10 |
| | 1449 | 12/22/2021 | REGN-008CIPCON10 |
| | 1449 | 12/22/2021 | REGN-008CIPCON10 |

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Issue Fee Transmittal Form

| Application Number | Filing Date | First Named Inventor | Atty. Docket No. | Confirmation No. |
|--------------------|-------------|----------------------|------------------|------------------|
| 17352892 | 21-Jun-2021 | George YANCOPOULOS | REGN-008CIPCON10 | 5070 |
| | | TITLE OF INVENTION : | • | • |

USE OF A VEGF ANTAGONIST TO TREAT ANGIOGENIC EYE DISORDERS

| Entity Sta | tus | | Application Type A | | Art Unit | Class - Subclas | s EXAMINER |
|----------------------|----------------|----------------------------|--------------------|-------------|-----------|-----------------|----------------|
| Regular Undiscounted | | Utility under 35 USC 111(a | | 1647 134100 | | 134100 | |
| Issue Fee Due | Publication Du | e | Total Fee(s) Due | 1 | Da | ate Due | Prev. Paid Fee |
| \$1200 | \$0 | | \$1200 | | 22-Mar-20 | 22 | \$0 |

1. Change of Correspondence Address and/or Indication Of Fee Address (37 CFR 1.33 & 1.363)

| Current Correspondence Address: | Current Indicated Fee Address : |
|--|--|
| 96387
Regeneron - Bozicevic, Field & Francis | |
| 201 REDWOOD SHORES PARKWAY
SUITE 200
REDWOOD CITY CA 94065
UNITED STATES
650 327 3400
docket@bozpat.com | |
| Change of correspondence address requested, system
generated AIA/122-EFS form attached | Fee Address indication requested, system generated SB/47-EFS form attached |

2.Entity Status

Change in Entity Status

Applicant certifying micro entity status; system generated Micro Entity certification form attached. See 37 CFR 1.29. Note: Absent a valid certification of micro entity status, issue fee payment in the micro entity amount will not be accepted at the risk of application abandonment.

If this box is checked, you will be prompted to choose a micro entity status on the gross income basis (37 CFR 1.29(a)) or the institution of higher education basis (37 CFR 1.29(d)), and make the applicable certification online.

- Applicant asserting small entity status. See 37 CFR 1.27.
- Note: If the application was previously under micro entity status, checking this box will be taken to be a notification of loss of entitlement to micro entity status.
- Applicant changing to regular undiscounted fee status.
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1. THOMAS TRIOLO

- 2. Karl Bozicevic
- 3.

5.Assignee Name(s) and Residence Data To Be Printed

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|---------------------------------|-----------|----------|---------------|-------------|
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6.Signature

I certify, in accordance with 37 CFR 1.4(d)(4) that I am an attorney or agent registered to practice before the Patent and Trademark Office who has filed and has been granted power of attorney in this application. I also certify that this Fee(s) Transmittal form is being transmitted to the USPTO via EFS-WEB on the date indicated below.

| Signature | /Karl Bozicevic/ | Date | 01-12-2022 |
|-----------|------------------|---------------------|------------|
| Name | Karl Bozicevic | Registration Number | 28807 |

WEB IFEE 1.0

| Electronic Patent Application Fee Transmittal | | | | | |
|---|---------------------------------|-------------------|----------------|-----------------|-------------------------|
| Application Number: | 173 | 352892 | | | |
| Filing Date: | 21- | 21-Jun-2021 | | | |
| Title of Invention: | US | E OF A VEGF ANTAC | 50NIST TO TREA | T ANGIOGENIC EY | E DISORDERS |
| First Named Inventor/Applicant Name: | George YANCOPOULOS | | | | |
| Filer: | Karl Bozicevic/Kimberly Zuehlke | | | | |
| Attorney Docket Number: | REGN-008CIPCON10 | | | | |
| Filed as Large Entity | | | | | |
| Filing Fees for Utility under 35 USC 111(a) | | | | | |
| Description | | Fee Code | Quantity | Amount | Sub-Total in
USD(\$) |
| Basic Filing: | | | | | |
| UTILITY APPL ISSUE FEE | | 1501 | 1 | 1200 | 1200 |
| PUBL. FEE- EARLY, VOLUNTARY, OR NORMAL | | 1504 | 1 | 0 | 0 |
| Pages: | | | | | |
| Claims: | | | | | |
| Miscellaneous-Filing: | | | | | |
| Petition: | | | | | |
| Patent-Appeals-and-Interference: | | | | | |

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| Post-Allowance-and-Post-Issuance: | | | | |
| Extension-of-Time: | | | | |
| Miscellaneous: | | | | |
| | Tot | al in USD | (\$) | 1200 |
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|--------------------------------------|--|--|--|--|--|
| EFS ID: | 44730999 | | | | |
| Application Number: | 17352892 | | | | |
| International Application Number: | | | | | |
| Confirmation Number: | 5070 | | | | |
| Title of Invention: | USE OF A VEGF ANTAGONIST TO TREAT ANGIOGENIC EYE DISORDERS | | | | |
| First Named Inventor/Applicant Name: | George YANCOPOULOS | | | | |
| Customer Number: | 96387 | | | | |
| Filer: | Karl Bozicevic/Kimberly Zuehlke | | | | |
| Filer Authorized By: | Karl Bozicevic | | | | |
| Attorney Docket Number: | REGN-008CIPCON10 | | | | |
| Receipt Date: | 12-JAN-2022 | | | | |
| Filing Date: | 21-JUN-2021 | | | | |
| Time Stamp: | 17:33:41 | | | | |
| Application Type: | Utility under 35 USC 111(a) | | | | |

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|-----------------|------------|------------|---------------------|------------------|
| APPLICATION NO. | ISSUE DATE | PATENT NO. | ATTORNEY DOCKET NO. | CONFIRMATION NO. |
| 17/352,892 | 02/22/2022 | 11253572 | REGN-008CIPCON10 | 5070 |

96387759002/02/2022Regeneron - Bozicevic, Field & Francis201 REDWOOD SHORES PARKWAYSUITE 200REDWOOD CITY, CA 94065

ISSUE NOTIFICATION

The projected patent number and issue date are specified above.

Determination of Patent Term Adjustment under 35 U.S.C. 154 (b)

(application filed on or after May 29, 2000)

The Patent Term Adjustment is 0 day(s). Any patent to issue from the above-identified application will include an indication of the adjustment on the front page.

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INVENTOR(s) (Please see PAIR WEB site http://pair.uspto.gov for additional inventors):

George YANCOPOULOS, Yorktown Heights, NY;

APPLICANT(s) (Please see PAIR WEB site http://pair.uspto.gov for additional applicants):

REGENERON PHARMACEUTICALS, INC., Tarrytown, NY

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| 17352892 | ISSUE.NTF | 02/02/2022 | REGN-008CIPCON10 |
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| Electronically Filed | | | | |
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| PETITION FOR CERTIFICATE | Attorney Docket No. | REGN-008CIPCON10 | | |
| OF CORRECTION | First Named Inventor | George D. Yancopoulos | | |
| Address to: | Patent Number | 11,253,572 | | |
| | Issue Date | February 22, 2022 | | |
| Mail Stop Certificate of Correction Branch | Application Number | 17/352,892 | | |
| Commissioner for Patents | Filing Date | June 21, 2021 | | |
| P.O. Box 1450 | Title: "Use of a VEG | F Antagonist to Treat Angiogenic | | |
| Alexandria, VA 22313-1450 | Eye Disorders" | | | |

Sir:

Transmitted herewith for filing is a Certificate of Correction for the above-identified patent. This request is being submitted to correct typographical errors made during the printing of the patent in a manner that does not correspond to the language (specific symbol) shown in the originally filed specification.

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> Respectfully submitted, BOZICEVIC, FIELD & FRANCIS LLP

Date: 4 March 2022

By: /Karl Bozicevic, Reg. No. 28,807/ Karl Bozicevic Reg. No. 28,807

BOZICEVIC, FIELD & FRANCIS LLP 201 Redwood Shores Parkway, Suite 200 Redwood City, CA 94065 Telephone: (650) 327-3400 Facsimile: (650) 327-3231

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Page <u>1</u> of <u>1</u>

PATENT NO. : 11,253,572

APPLICATION NO. : 17/352,892

ISSUE DATE : February 22, 2022

INVENTOR(S) : George D. Yancopoulos

It is certified that an error appears or errors appear in the above-identified patent and that said Letters Patent is hereby corrected as shown below:

At column 15, lines 36-37, please correct the specification from "gained ETDRS" to read --gained ≥15 ETDRS--.

MAILING ADDRESS OF SENDER (Please do not use customer number below):

BOZICEVIC, FIELD & FRANCIS LLP 201 Redwood Shores Pkwy, Suite 200 Redwood City, California 94065

This collection of information is required by 37 CFR 1.322, 1.323, and 1.324. The information is required to obtain or retain a benefit by the public which is to file (and by the USPTO to process) an application. Confidentiality is governed by 35 U.S.C. 122 and 37 CFR 1.14. This collection is estimated to take 1.0 hour to complete, including gathering, preparing, and submitting the completed application form to the USPTO. Time will vary depending upon the individual case. Any comments on the amount of time you require to complete this form and/or suggestions for reducing this burden, should be sent to the Chief Information Officer, U.S. Patent and Trademark Office, U.S. Department of Commerce, P.O. Box 1450, Alexandria, VA 22313-1450. DO NOT SEND FEES OR COMPLETED FORMS TO THIS ADDRESS. SEND TO: Attention Certificate of Corrections Branch, Commissioner for Patents, P.O. Box 1450, Alexandria, VA 22313-1450.

If you need assistance in completing this form, call 1-800-PTO-9199 and select option 2.

| Electronic Acknowledgement Receipt | | | |
|--------------------------------------|--|--|--|
| EFS ID: | 45146528 | | |
| Application Number: | 17352892 | | |
| International Application Number: | | | |
| Confirmation Number: | 5070 | | |
| Title of Invention: | USE OF A VEGF ANTAGONIST TO TREAT ANGIOGENIC EYE DISORDERS | | |
| First Named Inventor/Applicant Name: | George YANCOPOULOS | | |
| Customer Number: | 96387 | | |
| Filer: | Karl Bozicevic/Kimberly Zuehlke | | |
| Filer Authorized By: | Karl Bozicevic | | |
| Attorney Docket Number: | REGN-008CIPCON10 | | |
| Receipt Date: | 04-MAR-2022 | | |
| Filing Date: | 21-JUN-2021 | | |
| Time Stamp: | 12:40:47 | | |
| Application Type: | Utility under 35 USC 111(a) | | |

Payment information:

| Submitted with Payment | | | no | | | |
|------------------------|---------------------------------------|-----|--|--|---------------------|---------------------|
| File Listing: | | | | | | |
| Document
Number | Document Description | | File Name | File Size(Bytes)/
Message Digest | Multi
Part /.zip | Pages
(if appl.) |
| | | | 21458 | | | |
| 1 | Request for Certificate of Correction | REG | GN-008CIPCON10_2022-03-0
4_Petition_COC.pdf | ac6fb40bebcff9568a3d04842fc5545d9406
c551 | no | 1 |
| Warnings: | | | | | | |

| Information | | | | | |
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| 2 | Request for Certificate of Correction | REGN-008CIPCON10_2022-03-0
4_COC.pdf | 27746
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6a70 | no | 1 |
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| Information | | | | | |
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national seco | This Acknowledgement Receipt evidences receipt on the noted date by the USPTO of the indicated documents, characterized by the applicant, and including page counts, where applicable. It serves as evidence of receipt similar to a Post Card, as described in MPEP 503. New Applications Under 35 U.S.C. 111 If a new application is being filed and the application includes the necessary components for a filing date (see 37 CFR 1.53(b)-(d) and MPEP 506), a Filing Receipt (37 CFR 1.54) will be issued in due course and the date shown on this Acknowledgement Receipt will establish the filing date of the application. National Stage of an International Application under 35 U.S.C. 371 If a timely submission to enter the national stage of an international application is compliant with the conditions of 35 U.S.C. 371 and other applicable requirements a Form PCT/DO/EO/903 indicating acceptance of the application as a national stage submission under 35 U.S.C. 371 will be issued in addition to the Filing Receipt, in due course. New International Application is being filed and the international application includes the necessary components for an international application is being filed and the international application includes the necessary components for an international Application Filed with the USPTO as a Receiving Office If a new international application is being filed and the international application of the International Application Number and of the International Filing Date (Form PCT/RO/105) will be issued in due course, subject to prescriptions concerning national security, and the date shown on this Acknowledgement Receipt will establish the international filing date of the application. | | | | |

UNITED STATES PATENT AND TRADEMARK OFFICE CERTIFICATE OF CORRECTION

 PATENT NO.
 : 11,253,572 B2

 APPLICATION NO.
 : 17/352892

 DATED
 : February 22, 2022

 INVENTOR(S)
 : Yancopoulos

Page 1 of 1

It is certified that error appears in the above-identified patent and that said Letters Patent is hereby corrected as shown below:

In the Specification

At Column 15, Lines 36-37, please correct "gained ETDRS" to read --gained ≥15 ETDRS--.

Signed and Sealed this Twenty-ninth Day of March, 2022

Dunn Han Ŀ

Drew Hirshfeld Performing the Functions and Duties of the Under Secretary of Commerce for Intellectual Property and Director of the United States Patent and Trademark Office

| AO 120 (Rev. 08/10) | | | | |
|---|--------------------------------|---------|---|--|
| Mail Stop 8
TO: Director of the U.S. Patent and Trademark Office
P.O. Box 1450
Alexandria, VA 22313-1450 | | | REPORT ON THE
FILING OR DETERMINATION OF AN
ACTION REGARDING A PATENT OR
TRADEMARK | |
| filed in the U.S. Dist | | orthern | § 1116 you are hereby advised that a court action has been District of West Virginia on the following es 35 U.S.C. § 292.): | |
| DOCKET NO.
1:22-cv-61 | DATE FILED U.S. D
8/2/2022 | | STRICT COURT
Northern District of West Virginia | |
| PLAINTIFF | | | DEFENDANT | |
| REGENERON PHARMA | ACEUTICALS, INC. | | MYLAN PHARMACEUTICALS, INC. | |
| PATENT OR
TRADEMARK NO. | DATE OF PATENT
OR TRADEMARK | | HOLDER OF PATENT OR TRADEMARK | |
| 1 See attached | | | | |
| 2 | | | | |
| 3 | | | | |
| 4 | | | | |
| 5 | | | | |

In the above-entitled case, the following patent(s)/ trademark(s) have been included:

| DATE INCLUDED | INCLUDED BY | |
|----------------------------|--------------------------------|--|
| | | t 🗌 Answer 🗌 Cross Bill 🗌 Other Pleading |
| PATENT OR
TRADEMARK NO. | DATE OF PATENT
OR TRADEMARK | HOLDER OF PATENT OR TRADEMARK |
| 1 | | |
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In the above-entitled case, the following decision has been rendered or judgement issued:

| DECISION/JUDGEMENT | | |
|--------------------|-------------------|------------------|
| CLERK | (BY) DEPUTY CLERK | DATE
8/3/2022 |

Copy 1—Upon initiation of action, mail this copy to Director Copy 3—Upon termination of action, mail this copy to Director Copy 2—Upon filing document adding patent(s), mail this copy to Director Copy 4—Case file copy

| PATENT OR | DATE OF PATENT | HOLDER OF PATENT OR |
|---------------|--------------------|---------------------------------|
| TRADEMARK NO. | OR TRADEMARK | TRADEMARK |
| 7,070,959 | July 4, 2006 | Regeneron Pharmaceuticals, Inc. |
| 9,222,106 | December 29, 2015 | Regeneron Pharmaceuticals, Inc. |
| 9,254,338 | February 9, 2016 | Regeneron Pharmaceuticals, Inc. |
| 9,669,069 | June 6, 2017 | Regeneron Pharmaceuticals, Inc. |
| 9,816,110 | November 14, 2017 | Regeneron Pharmaceuticals, Inc. |
| 10,130,681 | November 20, 2018 | Regeneron Pharmaceuticals, Inc. |
| 10,406,226 | September 10, 2019 | Regeneron Pharmaceuticals, Inc. |
| 10,415,055 | September 17, 2019 | Regeneron Pharmaceuticals, Inc. |
| 10,464,992 | November 5, 2019 | Regeneron Pharmaceuticals, Inc. |
| 10,669,594 | June 2, 2020 | Regeneron Pharmaceuticals, Inc. |
| 10,857,205 | December 8, 2020 | Regeneron Pharmaceuticals, Inc. |
| 10,888,601 | January 12, 2021 | Regeneron Pharmaceuticals, Inc. |
| 10,927,342 | February 23, 2021 | Regeneron Pharmaceuticals, Inc. |
| 10,973,879 | April 13, 2021 | Regeneron Pharmaceuticals, Inc. |
| 11,053,280 | July 6, 2021 | Regeneron Pharmaceuticals, Inc. |
| 11,066,458 | July 20, 2021 | Regeneron Pharmaceuticals, Inc. |
| 11,084,865 | August 10, 2021 | Regeneron Pharmaceuticals, Inc. |
| 11,104,715 | August 31, 2021 | Regeneron Pharmaceuticals, Inc. |
| 11,174,283 | November 16, 2021 | Regeneron Pharmaceuticals, Inc. |
| 11,186,625 | November 30, 2021 | Regeneron Pharmaceuticals, Inc. |
| 11,253,572 | February 22, 2022 | Regeneron Pharmaceuticals, Inc. |
| 11,299,532 | April 12, 2022 | Regeneron Pharmaceuticals, Inc. |
| 11,306,135 | April 19, 2022 | Regeneron Pharmaceuticals, Inc. |
| 11,332,771 | May 17, 2022 | Regeneron Pharmaceuticals, Inc. |

Trials@uspto.gov 571-272-7822 Paper No. 3

UNITED STATES PATENT AND TRADEMARK OFFICE

BEFORE THE PATENT TRIAL AND APPEAL BOARD

APOTEX INC., Petitioner,

v.

REGENERON PHARMACEUTICALS, INC., Patent Owner.

IPR2022-01524 Patent 11,253,572

Mailed: September 28, 2022

Before Cathy Underwood, Trial Paralegal

NOTICE OF FILING DATE ACCORDED TO PETITION AND TIME FOR FILING PATENT OWNER PRELIMINARY RESPONSE

The petition for *inter partes* review, filed in the above proceeding has been accorded the filing date of September 9, 2022.

Patent Owner may file a preliminary response to the petition no later than three months from the date of this notice. The preliminary response is limited to setting forth the reasons why the requested review should not be instituted. Patent Owner may also file an election to waive the preliminary

response to expedite the proceeding. For more information, please consult the Office Patent Trial Practice Guide, 77 Fed. Reg. 48756 (Aug. 14, 2012), which is available on the Board Web site at <u>http://www.uspto.gov/PTAB</u>.

Patent Owner is advised of the requirement to submit mandatory notice information under 37 C.F.R. § 42.8(a)(2) within 21 days of service of the petition.

The parties are encouraged to use the heading on the first page of this Notice for all future filings in the proceeding.

The parties are advised that under 37 C.F.R. § 42.10(c), recognition of counsel pro hac vice requires a showing of good cause. The parties are authorized to file motions for pro hac vice admission under 37 C.F.R. § 42.10(c). Such motions shall be filed in accordance with the "Order -- Authorizing Motion for Pro Hac Vice Admission" in Case IPR2013-00639, Paper 7, a copy of which is available on the Board Web site under "Representative Orders, Decisions, and Notices." **The parties are reminded that, in order for any motion for** *pro hac vice* **admission to be considered by the Board, the requisite fees must first be paid.** The current fee schedule is available at https://www.uspto.gov/learning-and-resources/fees-and-payment/uspto-fee-schedule.

The parties are reminded that unless otherwise permitted by 37 C.F.R. § 42.6(b)(2), all filings in this proceeding must be made electronically in Patent Trial and Appeal Board End to End (PTAB E2E), accessible from the Board Web site at <u>http://www.uspto.gov/PTAB</u>. To file documents, users must register with PTAB E2E. Information regarding how to register with and use PTAB E2E is available at the Board Web site.

If there are any questions pertaining to this notice, please contact Cathy Underwood at 571-272-8358 or the Patent Trial and Appeal Board at 571-272-7822.

PETITIONER:

Teresa Stanek Rea Deborah H. Yellin Shannon M. Lentz CROWELL & MORING LLP <u>TRea@Crowell.com</u> <u>DYellin@Crowell.com</u> <u>SLentz@Crowell.com</u>

PATENT OWNER:

Regeneron – Bozicevic Field & Francis LLP 201 Redwood Shores Parkway Suite 200 Redwood City, CA 94065

NOTICE CONCERNING ALTERNATIVE DISPUTE RESOLUTION (ADR)

The Patent Trial and Appeal Board (PTAB) strongly encourages parties who are considering settlement to consider alternative dispute resolution as a means of settling the issues that may be raised in an AIA trial proceeding. Many AIA trials are settled prior to a Final Written Decision. Those considering settlement may wish to consider alternative dispute resolution techniques early in a proceeding to produce a quicker, mutually agreeable resolution of a dispute or to at least narrow the scope of matters in dispute. Alternative dispute resolution has the potential to save parties time and money.

Many non-profit organizations, both inside and outside the intellectual property field, offer alternative dispute resolution services. Listed below are the names and addresses of several such organizations. The listings are provided for the convenience of parties involved in cases before the PTAB; the PTAB does not sponsor or endorse any particular organization's alternative dispute resolution services. In addition, consideration may be given to utilizing independent alternative dispute resolution firms. Such firms may be located through a standard keyword Internet search.

| CPR
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FOR DISPUTE
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(ABA) |
|---|---|--|---|---|
| Telephone: | Telephone: | Telephone: | Telephone: | Telephone : |
| (212) 949-6490 | (703) 415-0780 | (212) 484-3266 | 41 22 338 9111 | (202) 662-1000 |
| Fax: (212) 949-8859 | Fax: (703) 415-0786 | Fax: (212) 307-4387 | Fax: 41 22 733 5428 | N/A |
| | 241 18th Street, South, | 140 West 51st | 34, chemin des | 1050 Connecticut Ave, |
| 575 Lexington Ave | Suite 700 | Street | Colombettes | NW |
| New York, NY 10022 | Arlington, VA 22202 | New York, NY
10020 | CH-1211 Geneva 20,
Switzerland | Washington D.C. 20036 |
| www.cpradr.org | www.aipla.org | www.adr.org | www.wipo.int | www.americanbar.org |

If parties to an AIA trial proceeding consider using alternative dispute resolution, the PTAB would like to know whether the parties ultimately decided to engage in alternative dispute resolution and the reasons why or why not. If the parties actually engage in alternative dispute resolution, the PTAB would be interested to learn what mechanism (e.g., arbitration,

mediation, etc.) was used and the general result. Such a statement from the parties is not required but would be helpful to the PTAB in assessing the value of alternative dispute resolution to parties involved in AIA trial proceedings. To report an experience with ADR, please forward a summary of the particulars to the following email address: PTAB_ADR_Comments@uspto.gov

| PTO/AIA/80 (07-17) |
|---|
| Approved for use through 03/31/2021. OMB 0651-0035 |
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| POW | ER OF ATTORNEY TO | PROSECU | TE APPLICATIO | ONS BEFOR | E THE USPTC | |
|----------|--|------------------------|--|----------------------|------------------------|--|
| | by revoke all previous powers | of attorney gi | ven in the application | on identified in | the attached | |
| | ent under 37 CFR 3.73(c). | | | | | |
| i neret | by appoint: | | | | | |
| | Practitioners associated with Customs | er Number: 191 | 459 | | | |
| | OR | L | | | | |
| | Practitioner(s) named below (if more | than ten patent pra | ctitioners are to be named, | , then a customer nu | mber must be used): | |
| | Name | Registration
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| | Please change the correspondence address for the application identified in the attached statement
under 37 CFR 3.73(c) to:
X The address associated with Customer Number: 191459
OR | | | | | |
| | Firm or individual name | | | | | |
| | Address | | | | | |
| | City | | State | Zip | | |
| | Country | | | | | |
| | Telephone Email | | | | | |
| Assigne | Assignee name and address: Regeneron Pharmaceuticals, Inc.
777 Old Saw Mill River Road
Tarrytown, NY 10591-5707 | | | | | |
| filed in | of this form, together with a staten
each application in which this form
oners appointed in this form, and n | is used. The state | ment under 37 CFR 3.7 | '3{c} may be comp | leted by one of the | |
| | The individual whose signature a | | Assignee of Record
I below is authorized to | act on behalf of th | ne assignee | |
| Signat | The individual whose signature and title is supplied below is authorized to act on behalf of the assignee. Signature /Frank R. Cottingham/ Date 13 January 2023 | | | | | |
| | Name Frank R. Cottingham, Ph.D., J.D. Telephone | | | | | |
| h | ice President, Associate G | | 1 | roperty | | |
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This collection of information is required by 37 CFR 1.31, 1.32, and 1.33. The information is required to obtain or retain a benefit by the public, which is to update (and by the USPTO to process) the file of a patent or reexamination proceeding. Confidentiality is governed by 35 U.S.C. 122 and 37 CFR 1.11 and 1.14. This collection is estimated to take 18 minutes to complete, including gathering, preparing, and submitting the completed application form to the USPTO. Time will vary depending upon the individual case. Any comments on the amount of time you require to complete this form and/or suggestions for reducing this burden, should be sent to the Chief information Differe, U.S. Patent and Trademark Office, U.S. Department of Commerce, P.O. Box 1450, Alexandria, VA 22313-1450. DO NGT SEND FEES OR COMPLETED FORMS TO THIS ADDRESS, SEND TO: Commissioner for Patents, P.O. Box 1450, Alexandria, VA 22313-1450. If you need assistance in completing the form, call 1-800-PTO-\$129 ond select option 2.

Privacy Act Statement

The **Privacy Act of 1974 (P.L. 93-579)** requires that you be given certain information in connection with your submission of the attached form related to a patent application or patent. Accordingly, pursuant to the requirements of the Act, please be advised that: (1) the general authority for the collection of this information is 35 U.S.C. 2(b)(2); (2) furnishing of the information solicited is voluntary; and (3) the principal purpose for which the information is used by the U.S. Patent and Trademark Office is to process and/or examine your submission related to a patent application or patent. If you do not furnish the requested information, the U.S. Patent and Trademark Office may not be able to process and/or examine your submission, which may result in termination of proceedings or abandonment of the application or expiration of the patent.

The information provided by you in this form will be subject to the following routine uses:

- The information on this form will be treated confidentially to the extent allowed under the Freedom of Information Act (5 U.S.C. 552) and the Privacy Act (5 U.S.C 552a). Records from this system of records may be disclosed to the Department of Justice to determine whether disclosure of these records is required by the Freedom of Information Act.
- A record from this system of records may be disclosed, as a routine use, in the course of presenting evidence to a court, magistrate, or administrative tribunal, including disclosures to opposing counsel in the course of settlement negotiations.
- 3. A record in this system of records may be disclosed, as a routine use, to a Member of Congress submitting a request involving an individual, to whom the record pertains, when the individual has requested assistance from the Member with respect to the subject matter of the record.
- 4. A record in this system of records may be disclosed, as a routine use, to a contractor of the Agency having need for the information in order to perform a contract. Recipients of information shall be required to comply with the requirements of the Privacy Act of 1974, as amended, pursuant to 5 U.S.C. 552a(m).
- 5. A record related to an International Application filed under the Patent Cooperation Treaty in this system of records may be disclosed, as a routine use, to the International Bureau of the World Intellectual Property Organization, pursuant to the Patent Cooperation Treaty.
- A record in this system of records may be disclosed, as a routine use, to another federal agency for purposes of National Security review (35 U.S.C. 181) and for review pursuant to the Atomic Energy Act (42 U.S.C. 218(c)).
- 7. A record from this system of records may be disclosed, as a routine use, to the Administrator, General Services, or his/her designee, during an inspection of records conducted by GSA as part of that agency's responsibility to recommend improvements in records management practices and programs, under authority of 44 U.S.C. 2904 and 2906. Such disclosure shall be made in accordance with the GSA regulations governing inspection of records for this purpose, and any other relevant (*i.e.*, GSA or Commerce) directive. Such disclosure shall not be used to make determinations about individuals.
- 8. A record from this system of records may be disclosed, as a routine use, to the public after either publication of the application pursuant to 35 U.S.C. 122(b) or issuance of a patent pursuant to 35 U.S.C. 151. Further, a record may be disclosed, subject to the limitations of 37 CFR 1.14, as a routine use, to the public if the record was filed in an application which became abandoned or in which the proceedings were terminated and which application is referenced by either a published application, an application open to public inspection or an issued patent.
- A record from this system of records may be disclosed, as a routine use, to a Federal, State, or local law enforcement agency, if the USPTO becomes aware of a violation or potential violation of law or regulation.



ELECTRONIC ACKNOWLEDGEMENT RECEIPT

| APPLICATION #
17/352,892 | RECEIPT DATE / TIME
02/28/2023 02:40:33 PM ET | | EY DOCKET #
I-008CIPCON10 | | | |
|--|--|-------------------------|------------------------------|--|--|--|
| Title of Invention
USE OF A VEGF ANTAGONIST TO TREAT ANGIOGENIC EYE DISORDERS | | | | | | |
| Application Infor | rmation | | | | | |
| APPLICATION TYPE | Utility - Nonprovisional Application under 35 USC 111(a) | PATENT # | 11253572 | | | |
| CONFIRMATION # | 5070 | FILED BY | Kathi Moore | | | |
| PATENT CENTER # | 61663248 | FILING DATE | 06/21/2021 | | | |
| CUSTOMER # | 96387 | FIRST NAMED
INVENTOR | George YANCOPOULOS | | | |
| CORRESPONDENCE
ADDRESS | - | AUTHORIZED BY | Michael Lewis | | | |

Documents

TOTAL DOCUMENTS: 2

| DOCUMENT | PAGES | DESCRIPTION | SIZE (KB) |
|---------------------|-------|---|-----------|
| P35056US13-373c.pdf | 3 | Assignee showing of ownership per 37 CFR 3.73 | 110 KB |
| POA-Regeneron.pdf | 2 | Power of Attorney | 3810 KB |

Digest

| DOCUMENT | MESSAGE DIGEST(SHA-512) |
|---------------------|--|
| P35056US13-373c.pdf | 9B516C78BB3105F2F8119345C7C2FAABBFBE44AC169FF5756
A432FF0EF20C213F61874AA800073A321BB1AE5BCF9B5E3E1
F37509E7540F7F404D71F7C39A7695 |

BDECE05D2F51BBCDB37E674E496644E6B6A6D74B020A14B4 FE816D8015E1EC9283171078BA0A0E1076CA395ADB4C661BA 9E9C16DA415BC049CBFF10B715A0675

This Acknowledgement Receipt evidences receipt on the noted date by the USPTO of the indicated documents, characterized by the applicant, and including page counts, where applicable. It serves as evidence of receipt similar to a Post Card, as described in MPEP 503.

New Applications Under 35 U.S.C. 111

If a new application is being filed and the application includes the necessary components for filing date (see 37 CFR 1.53(b)-(d) and MPEP 506), a Filing Receipt (37 CFR 1.54) will be issued in due course and the date shown on this Acknowledgement Receipt will establish the filing date of the application

National Stage of an International Application under 35 U.S.C. 371

If a timely submission to enter the national stage of an international application is compliant with the conditions of 35 U.S.C. 371 and other applicable requirements a Form PCT/DO/EO/903 indicating acceptance of the application as a national stage submission under 35 U.S.C. 371 will be issued in addition to the Filing Receipt, in due course.

New International Application Filed with the USPTO as a Receiving Office

If a new international application is being filed and the international application includes the necessary components for an international filing date (see PCT Article 11 and MPEP 1810), a Notification of the International Application Number and of the International Filing Date (Form PCT/RO/105) will be issued in due course, subject to prescriptions concerning national security, and the date shown on this Acknowledgement Receipt will establish the International filing date of the application.

PTO/AIA/96 (08-12) Approved for use through 11/30/2020. OMB 0651-0031

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Under the Paperwork Reduction Act of 1995, no persons are required to respond to a collection of information unless it displays a valid OMB control numb |
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| STATEMENT UNDER 37 CFR 3.73(c) |
| Applicant/Patent Owner: Regeneron Pharmaceuticals, Inc. |
| Application No./Patent No.: 11,253,572 Filed/Issue Date: February 22, 2022 |
| Titled: Use of a VEGF Antagonist to Treat Angiogenic Eye Disorders |
| Regeneron Pharmaceuticals, Inc, a corporation |
| (Name of Assignee) (Type of Assignee, e.g., corporation, partnership, university, government agency, etc.) |
| states that, for the patent application/patent identified above, it is (choose one of options 1, 2, 3 or 4 below): |
| 1. 🕑 The assignee of the entire right, title, and interest. |
| 2. An assignee of less than the entire right, title, and interest (check applicable box): |
| The extent (by percentage) of its ownership interest is%. Additional Statement(s) by the owners holding the balance of the interest must be submitted to account for 100% of the ownership interest. |
| There are unspecified percentages of ownership. The other parties, including inventors, who together own the entir right, title and interest are: |
| |
| |
| Additional Statement(s) by the owner(s) holding the balance of the interest <u>must be submitted</u> to account for the enti
right, title, and interest. |
| 3. The assignee of an undivided interest in the entirety (a complete assignment from one of the joint inventors was made) |
| The other parties, including inventors, who together own the entire right, title, and interest are: |
| |
| |
| Additional Statement(s) by the owner(s) holding the balance of the interest <u>must be submitted</u> to account for the entir
right, title, and interest. |
| 4. The recipient, via a court proceeding or the like (<i>e.g.</i> , bankruptcy, probate), of an undivided interest in the entirety (a complete transfer of ownership interest was made). The certified document(s) showing the transfer is attached. |
| The interest identified in option 1, 2 or 3 above (not option 4) is evidenced by either (choose one of options A or B below): |
| |
| A. [-] An assignment from the inventor(s) of the patent application/patent identified above. The assignment was recorded in the United States Patent and Trademark Office at Reel <u>057417</u> , Frame <u>0322</u> , or for which a copy thereof is attached. |
| B. 🗌 A chain of title from the inventor(s), of the patent application/patent identified above, to the current assignee as follows: |
| 1. From: To: |
| The document was recorded in the United States Patent and Trademark Office at |
| Reel, Frame, or for which a copy thereof is attached. |
| 2. From: To: |
| The document was recorded in the United States Patent and Trademark Office at |
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| [Page 1 of 2] |

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| /Michael | W. Lewis/ | | February 28, 2023 | |
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[Page 2 of 2]

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| APPLICATION NUMBER | FILING OR 371(C) DATE | FIRST NAMED APPLICANT | ATTY. DOCKET NO./TITLE |
| 17/352,892 | 06/21/2021 | George YANCOPOULOS | REGN-008CIPCON10 |
| | | | CONFIRMATION NO. 5070 |
| 191459 | | POA ACC | EPTANCE LETTER |
| A&P - Regeneron (Prosect
601 Massachusetts Ave., I
Washington, DC 20001-37 | NW Ó | | OC000000137523336* |
| - | | | Date Mailed: 03/07/2023 |

NOTICE OF ACCEPTANCE OF POWER OF ATTORNEY

This is in response to the Power of Attorney filed 02/28/2023.

The Power of Attorney in this application is accepted. Correspondence in this application will be mailed to the above address as provided by 37 CFR 1.33.

Questions about the contents of this notice and the requirements it sets forth should be directed to the Office of Data Management, Application Assistance Unit, at (571) 272-4000 or (571) 272-4200 or 1-888-786-0101.

/sltorres/

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| 17/352,892 | 06/21/2021 | George YANCOPOULOS | REGN-008CIPCON10 | |
| | | | CONFIRMATION NO. 5070 | |
| 96387 | | POWER | OF ATTORNEY NOTICE | |
| Regeneron - Bozicevic, Fi
201 REDWOOD SHORES
SUITE 200
REDWOOD CITY, CA 940 | S PARKWAY | | CC000000137523335* | |

Date Mailed: 03/07/2023

NOTICE REGARDING CHANGE OF POWER OF ATTORNEY

This is in response to the Power of Attorney filed 02/28/2023.

• The Power of Attorney to you in this application has been revoked by the applicant. Future correspondence will be mailed to the new address of record(37 CFR 1.33).

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/sltorres/

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| 17352892 | N570 | 03/07/2023 | REGN-008CIPCON10 |
| | N570 | 03/07/2023 | REGN-008CIPCON10 |

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UNITED STATES PATENT AND TRADEMARK OFFICE PATENT APPLICATION INFORMATION RETRIEVAL SYSTEM Trials@uspto.gov 571.272.7822 Paper 9 Entered: March 10, 2023

UNITED STATES PATENT AND TRADEMARK OFFICE

BEFORE THE PATENT TRIAL AND APPEAL BOARD

APOTEX INC., Petitioner,

V.

REGENERON PHARMACEUTICALS, INC., Patent Owner.

IPR2022-01524 Patent 11,253,572 B2

Before SUSAN L. C. MITCHELL, ROBERT A. POLLOCK, and RYAN H. FLAX, *Administrative Patent Judges*.

FLAX, Administrative Patent Judge.

DECISION Denying Institution of *Inter Partes* Review 35 U.S.C. § 314

I. INTRODUCTION

Regeneron Pharmaceuticals, Inc. ("Patent Owner") is the owner of U.S. patent 11,253,572B2 ("the '572 patent"). Paper 5, 1. On September 9, 2022, Apotex Inc. ("Petitioner") filed a Petition for *inter partes* review challenging the patentability of claims 1–14 and 26–30 of the '572 patent (claims 15–25 are not challenged). Paper 1, 1 ("Pet."). On December 23, 2022, Patent Owner filed a Preliminary Response to the Petition. Paper 7 ("Prelim. Resp."). No further briefing was requested or authorized.

Under 37 C.F.R. § 42.4(a), we have authority to determine whether to institute trial in an *inter partes* review. We may institute an *inter partes* review if the information presented in the petition filed under 35 U.S.C. § 311, and any preliminary response filed under § 313, shows that there is a reasonable likelihood that Petitioner would prevail with respect to at least one of the claims challenged in the petition. 35 U.S.C. § 314.

After reviewing the parties' submissions, we conclude Petitioner does not demonstrate a reasonable likelihood it would prevail in showing that any challenged claim of the '572 patent is unpatentable under the presented grounds. Therefore, we deny institution of *inter partes* review.¹ Our reasoning is discussed below.

A. REAL PARTIES-IN-INTEREST

Petitioner lists Apotex Inc., Apotex Corp, Apotex Pharmaceutical Holdings Inc, and Aposherm Delaware Holdings Corp. as real parties-in-

¹ We note that there are disputed issues in this proceeding under 35 U.S.C. \$ 325(d)and \$ 314(a). *See* Pet. 6–11; Prelim. Resp. 47–57. However, because we determine institution should be denied on the merits, we do not address these matters.

interest. Pet. 2. PatentOwner identifies itself as the only real party-in-interest. Paper 5, 1.

B. RELATED MATTERS

Petitioner identifies the following as related matters: IPR2021-00881 (concerning U.S. Patent 9,254,338 ("the '338 patent"); IPR2022-00258 (also concerning the '338 patent); IPR2022-00298 (also concerning the '338 patent); IPR2021-00880 (concerning U.S. Patent 9,669,069 ("the '069 patent)); IPR2022-0257 (also concerning the '069 patent); IPR2022-00301 (also concerning the '069 patent); IPR2022-01225 (concerning U.S. Patent 10,130,681 ("the '681 patent"); and IPR2022-01226 (concerning U.S. Patent 10,888,601 ("the '661 patent"). Pet. 3–4. Petitioner also identifies as related *Regeneron Pharms., Inc. v. Mylan Pharms. Inc.*, No. 1:22-cv-00061-TSK (N.D. W. Va), and PGR2021-00035 (concerning U.S. Patent 10,828,345). *Id.* at 5. In addition to the above-listed patents, Petitioner identifies U.S. Patent Application Nos. 17/072,417; 17/112,404; 17/112,063; and 17/350,958 as related. *Id.* Patent Owner identifies the same matters, patents, and applications as related. Paper 5, 2–3.

C. The '572 Patent

The '572 patent issued on February 22, 2022, from U.S. Application 17/352,892, which was filed on June 21, 2021. Ex. 1001, codes (45), (21), (22). The '572 patent ultimately indicates priority to U.S. Provisional Application 61/432,245, filed on January 13, 2011. *Id.* at code (60), 1:7–29. Petitioner declines to challenge whether the '572 patent is entitled such priority. *See, e.g.*, Pet. 1 ("Long before the patent's alleged 2011 priority date").

The '572 patent's abstract states:

The present invention provides methods for treating angiogenic eye disorders by sequentially administering multiple doses of a VEGF antagonist to a patient. The methods of the present invention include the administration of multiple doses of a VEGF antagonist to a patient at a frequency of once every 8 or more weeks. The methods of the present invention are useful for the treatment of angiogenic eye disorders such as age related macular degeneration, diabetic retinopathy, diabetic macular edema, central retinal vein occlusion, branch retinal vein occlusion, and corneal neovascularization.

Id. at Abstract.

As background, the '572 patent states that "[r]elease of vascular endothelial growth factor (VEGF) contributes to increased vascular permeability in the eye and inappropriate new vessel growth," and "inhibiting the angiogenic-promoting properties of VEGF appears to be an effective strategy for treating angiogenic eye disorders." *Id.* at 1:60–65. As further background, the '572 patent identifies that "FDA-approved treatments of angiogenic eye disorders such as AMD and CRVO include the administration of an anti-VEGF antibody called ranibizumab (Lucentis®, Genentech, Inc.) on a monthly basis by intravitreal injection." *Id.* at 1:66– 2:2. The '572 patent indicates that its invention is a response to the need for "new administration regimes" of "less frequent dosing while maintaining a high level of efficacy." *Id.* at 2:6–9.

In summarizing its invention, the '572 patent states:

The present inventors have surprisingly discovered that beneficial therapeutic effects can be achieved in patients suffering from angiogenic eye disorders by administering a VEGF antagonist to a patient at a frequency of once every 8 or more weeks, especially when such doses are preceded by about three doses administered to the patient at a frequency of about 2

to 4 weeks. Thus, according to the methods of the present invention, each secondary dose of VEGF antagonist is administered 2 to 4 weeks after the immediately preceding dose, and each tertiary dose is administered at least 8 weeks after the immediately preceding dose.

Id. at 2:22–33. Relating to this, the '572 patent defines certain terms. For example, "the VEGF antagonist comprises one or more VEGF receptor-based chimeric molecule(s), (also referred to herein as a 'VEGF-Trap' or 'VEGFT')," and an example of this includes a product called "aflibercept," marketed as "EYLEA" by Regeneron Pharmaceuticals, Inc. and approved by the FDA in November 2011 at a dose of 2 mg via intravitreal injection every 4 weeks for three months and then every 8 weeks. *Id.* at 2:47–67.

On the aforementioned FDA-approved dosing regimen, the '572 patent further defines the terms (ultimately used in the claims) "initial dose, "secondary doses," and "tertiary doses" as follows:

the "initial dose" is the dose which is administered at the beginning of the treatment regimen (also referred to as the "baseline dose"); the "secondary doses" are the doses which are administered after the initial dose; and the "tertiary doses" are the doses which are administered after the secondary doses.

Id. at 3:51–58.

The '572 patent describes a series of Examples detailing clinical trials conducted to validate the VEGFT drug and the dosing regimen. *Id.* at 8:12–18:3. Example 4 details two "Phase III Clinical Trials of the Efficacy, Safety, and Tolerability of Repeated Doses of Intravitreal VEGFT in Subjects with Neovascular Age-Related Macular Degeneration" (AMD) (Study 1 and Study 2), which followed dosing regimens using 2 mg doses of aflibercept at the aforementioned initial dose, then two 4-week doses, and then doses every 8-weeks through the end of the 52-week study (the "2Q8"

regimen). Id. at 9:29–14:30. The results of this and other regimens were compared to subjects administered 0.5 mg ranibizumab every 4 weeks (the "RQ4" regimen) by assessing patients' visual acuity based on a Best Corrected Visual Acuity (BCVA) test, which is based on the ability to identify letters. Id. at 9:35-10:7. This disclosure describes the inclusion criteria and exclusion criteria for the participating patients. Id. at 10:50-12:22.

Results of the Example 4 clinical trials are described in TABLE 1, which we reproduce below:

| 1/11/1./1./ 1 | | | | |
|---------------|--|------------------------------------|--------------------------------|---|
| | Ranibizumab
0.5 mg monthly
(RQ4) | VEGFT
0.5 mg monthly
(0.5Q4) | VEGFT
2 mg monthly
(2Q4) | VEGFT
2 mg every 8
weeks ^{iol} (2Q8) |
| Mainte | nance of vision* (% | patients losing <15 | letters) at week 52 | versus baseline |
| Study 1 | 94.4% | 95.9%** | 95.1%** | 95.1%** |
| Study 2 | 94,4% | 96.3%** | 95.6%** | 95.6%** |
| Mesn imp | novement in vision* | (letters) at 52 week | s versus baseline (p | »-valix: vs RQ4)*** |
| Study 1 | 8.1 | 6.9 (NS) | 10.9 (p. <0.61) | 7.9 (NS) |
| Study 2 | 9,4 | 9.7 (NS) | 7.6 (NS) | 8.9 (NS |

TABLE 1

^[s]Following three initial monthly doses

*Visual acuity was measured as the total number of letters read correctly on the Early Treatment Diabetic

Retainputhy Study (ETDRS) eye chart. **Statistically non-inferior based on a non-inferiority margin of 10%, using confidence interval approach (Q5.1% and 95% for Study 1 and Study 2, respectively) ***Test for supersority

NS = non-significant

Id. at 13:9–27. The '572 patent describes that these results showed that the VEGFT therapies usually maintained or improved visual acuity in patients and were not inferior to the ranibizumab treatment based on similar criteria. Id. at 13:28–38.

As Example 5, the '572 patent describes a phase 2 clinical trail using the same drug, also administered at 2 mg doses and, in one arm, at a regimen of three initial doses every four weeks followed by doses every eight weeks, but treating patients with diabetic macular edema (DME). *Id.* at 14:32–15:5. The '572 patent describes that visual acuity in this trial was maintained or improved for all VEGFT study groups. *Id.*

The '572 patent concludes with 30 claims, of which claims 1, 15 (not challenged), 26, and 29 are independent claims. Ex. 1012, 62:12–64:20. Claim 1 is illustrative and reproduced below:

1. A method of treating an angiogenic eye disorder in a patient in need thereof comprising sequentially administering to the patient by intravitreal injection a single initial dose of 2 mg of aflibercept, followed by one or more secondary doses of 2 mg of aflibercept, followed by one or more tertiary doses of 2 mg of aflibercept;

wherein each secondary dose is administered approximately 4 weeks following the immediately preceding dose; and

wherein each tertiary dose is administered approximately 8 weeks following the immediately preceding dose;

wherein the patient achieves a gain in visual acuity within 52 weeks following the initial dose.

Ex. 1001, 23:2–14.

Independent claim 26 is similar to claim 1 in reciting the same drug, at the same dose, and administered the same way, on the same schedule, but adds that the method treats "age related macular degeneration" (AMD) and that "the method is as effective in achieving a gain in visual acuity as monthly administration of 0.5 mg of ranibizumab by intravitreal injection in human subjects with age-related macular degeneration at 52 weeks

following the initial dose." *Id.* at 24:26–44. Independent claim 29 is also similar to claim 1, and essentially the same as claim 26, but differs in requiring effectiveness in "maintaining visual acuity" rather than a *gain* therein. *Id.* at 24:50–67.

D. ASSERTED GROUNDS FOR UNPATENTABILITY

Petitioner asserts the following grounds for the unpatentability of claims 1-14 and 26-30 of the '572 patent:

| Ground | Claims Challenged | 35 U.S.C. § ² | Reference(s)/Basis |
|--------|----------------------|--------------------------|-----------------------------|
| 1 | 1-5, 8-11, 14, 26-30 | 102 | Dixon ³ |
| 2 | 1-5, 8-11, 14, 26-30 | 102 | Regeneron 2008 ⁴ |
| 3 | 1-5, 8-11, 14, 26-30 | 102 | NCT-795 ⁵ |
| 4 | 1-5, 8-11, 14, 26-30 | 102 | NCT-377 ⁶ |

² The '572 patent has an uncontested January 13, 2011, priority date, which is before the AIA revisions to 35 U.S.C. §§ 102 and 103 took effect on March 16, 2013. 35 U.S.C. § 100 (note). Therefore, pre-AIA §§ 102 and 103 apply. Our decision is not impacted by which version of the statute applies.

³ James A. Dixon et al., *VEGF Trap-Eye for the treatment of neovascular age-related macular degeneration*, 18(10) EXPERT OPIN. INVESTIG. DRUGS 1573–80 (2009) (Ex. 1006, "Dixon").

⁴ Regeneron, News Release: *Bayer and Regeneron Dose First Patient in Second Phase 3 Study for VEGF Trap-Eye in Wet Age-Related Macular Degeneration -International study to evaluate efficacy and safety in treating a leading cause of blindness* 1–3 (May 8, 2008) (Ex. 1009, "Regeneron 2008").

⁵ NIH, U.S. National Library of Medicine, ClinicalTrials.gov archive, *History of Changes for Study: NCT00509795, Vascular Endothelial Growth Factor(VEGF)Trap-Eye:Investigation of Efficacy and Safety in Wet Age-Related Macular Degeneration(AMD) (VIEW1)* (Dec. 20, 2012), accessed January 7, 2021, at https://clinicaltrials.gov/ct2/history/NCT00509795?A =8&B=9&C=merged#StudyPageTop (Ex. 1010, "NCT-795").

⁶ NIH, U.S. National Library of Medicine, ClinicalTrials.gov archive, *History of Changes for Study: NCT00637377, VEGF Trap-Eye:*