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1 Serious Adverse Events with Bevacizumab or Ranibizumab for Age-related Macular  
2 Degeneration: Meta-analysis of Individual Patient Data

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5 The Bevacizumab-Ranibizumab International Trials Group ‡

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7 Writing Committee:

8 Maureen G. Maguire, PhD	Department of Ophthalmology, University of Pennsylvania, Philadelphia, Pennsylvania, United States
9 James Shaffer, MS	Department of Ophthalmology, University of Pennsylvania, Philadelphia, Pennsylvania, United States
10 Gui-shuang Ying, PhD	Department of Ophthalmology, University of Pennsylvania, Philadelphia, Pennsylvania, United States
11 Usha Chakravarthy, PhD, FRCS	Institute of Clinical Science, The Queen's University of Belfast, Belfast, Ireland
12 Karina Berg, MD	Department of Ophthalmology, Oslo University Hospital, Oslo, Norway
13 Ragnheiður Bragadóttir MD, PhD	Department of Ophthalmology, Oslo University Hospital, Oslo, Norway
14 Evelyne Decullier, PhD	Hospices Civils de Lyon, Département Recherche Clinique et Innovation, Lyon, France; Université de Lyon, Lyon France
15 Laure Huot, PharmD, PhD	Hospices Civils de Lyon, Département Recherche Clinique et Innovation, Lyon, France; Université de Lyon, Lyon France
16 Laurent Kodjikian, MD, PhD	Hospices Civils de Lyon, Croix-Rousse University Hospital, Department of Ophthalmology; Université de Lyon, Lyon, France
17 Daniel F. Martin, MD	Cole Eye Institute, Cleveland Clinic, Cleveland, Ohio United States
18 Barnaby C. Reeves, DPhil	Clinical Trials and Evaluation Unit, School of Clinical Sciences, University of Bristol, Bristol, UK
19 Chris A. Rogers, PhD	Clinical Trials and Evaluation Unit, School of Clinical Sciences, University of Bristol, Bristol, UK
20 Ann-Sofie M.E. Schauwvlieghe, MD	Department of Ophthalmology, Academic Medical Center, University of Amsterdam, Amsterdam, The Netherlands
21 Reinier O. Schlingemann, MD	Department of Ophthalmology, Academic Medical Center, University of Amsterdam, Amsterdam, The Netherlands

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41 ‡ The members of the Bevacizumab-Ranibizumab International Trials Group are listed in the  
42 Appendix (available at <http://www.aajournal.org>).

43  
44 This article contain additional online-only material. The following should appear online-only:  
45 Appendix and Figures 2 to 5.

46  
47 Corresponding Author:

48 Maureen G. Maguire, PhD 215 615 1501 (V) 215 615 1520 (Fax)

49 [mauirem@mail.med.upenn.edu](mailto:mauirem@mail.med.upenn.edu)

50 Department of Ophthalmology

51 3535 Market Street, Suite 700, Philadelphia PA 19104

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Running head: Meta-analysis of safety of bevacizumab and ranibizumab

Reprints requests to: Maureen Maguire, PhD, CATT Coordinating Center, University of Pennsylvania, 3535 Market Street, Suite 700, Philadelphia, PA 19104-3309

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76 **ABSTRACT**

77

78 **Topic:** A comparison between ranibizumab and bevacizumab of the incidence of systemic  
79 serious adverse events (SAEs) among patients with neovascular age-related macular  
80 degeneration (nAMD) who participated in a large-scale randomized trial. Use of individual  
81 patient data, rather than aggregate data, allowed adjustment for strong predictors of SAEs.

82 **Clinical relevance:** Relative safety of ranibizumab and bevacizumab is important in choosing  
83 an anti-VEGF drug for the hundreds of thousands of patients with nAMD treated each year  
84 worldwide.

85 **Methods:** Results of a Cochrane aggregate meta-analysis of the relative efficacy and safety of  
86 bevacizumab and ranibizumab that used searches of bibliographic databases and clinical trial  
87 registries as of March 14, 2014 and hand searching were reviewed to identify 6 large-scale,  
88 multicenter clinical trials. Individual patient data on SAEs, assigned drug and dosing regimen,  
89 and baseline prognostic factors were requested from the leaders of the 6 trials. A two-stage  
90 approach was used to estimate relative risks and 95% confidence intervals (CIs) from Cox  
91 proportional hazards models adjusting for baseline prognostic factors. The primary outcome  
92 measure was development of  $\geq 1$  SAE; secondary outcome measures were death,  
93 arteriothrombotic events, events associated with systemic anti-VEGF therapy, and events not  
94 associated with systemic anti-VEGF therapy.

95 **Results:** Individual patient data were received from 5 trials to provide information on 3052  
96 patients. There were no large imbalances between drug groups on baseline factors. The  
97 adjusted relative risk (95% CI) for bevacizumab relative to ranibizumab was 1.06 [(0.84, 1.35);  
98  $p=0.61$ ] for  $\geq 1$  SAEs. For secondary outcomes, adjusted relative risks were 0.99 [(0.69, 1.43);  
99  $p=0.97$ ] for death, 0.89 [(0.62, 1.28);  $p=0.53$ ] for arteriothrombotic events, 1.10 [(0.81, 1.50);  
100  $p=0.54$ ] for events related to anti-VEGF treatment, and 1.11 [(0.87, 1.40);  $p=0.40$ ] for events not  
101 related to anti-VEGF treatment.

102 **Conclusion:** Our findings support the absence of large differences in risk of systemic serious  
103 adverse events between these two anti-VEGF drugs; i.e., relative risks of  $\geq 1.5$  are unlikely.  
104 Because additional head-to-head trials are unlikely, any further investigation of differential risk  
105 between anti-VEGF agents will only be achieved through post-marketing surveillance or through  
106 the interrogation of healthcare databases.  
107

108           The management and prognosis of patients with neovascular age-related macular  
109 degeneration (AMD) changed dramatically in 2005 with the release of results from Phase III  
110 clinical trials of intravitreally administered ranibizumab (Lucentis; Genentech, Inc., South San  
111 Francisco, CA), an inhibitor of all active forms of vascular endothelial growth factor (VEGF).<sup>1,2</sup>  
112 On average, eyes treated with ranibizumab gained visual acuity while untreated eyes or eyes  
113 treated with photodynamic laser therapy lost substantial visual acuity. While waiting for approval  
114 from regulatory agencies in the United States and Europe, ophthalmologists began using  
115 intravitreal bevacizumab off label to treat neovascular AMD because it was structurally similar to  
116 ranibizumab (Avastin; Genentech, Inc., South San Francisco, CA), available for use because it  
117 had been approved for treatment of cancer, and was inexpensive. Short-term outcomes related  
118 to vision and retinal morphology after treatment with bevacizumab appeared similar to those of  
119 ranibizumab, leading to rapid adoption of bevacizumab as first-line therapy. The fact that after  
120 ranibizumab was approved by the Food and Drug Administration, ranibizumab was sold for  
121 approximately \$2000 per dose in the United States compared to \$50 for bevacizumab, amplified  
122 the need for comparison of longer term efficacy and safety between the two drugs.<sup>3</sup>

123           Planning for large-scale, multicenter clinical trials of the two drugs was initiated in 6  
124 different countries. These multicenter clinical trials were: the Comparison of Age-related  
125 Macular Degeneration Treatments Trials (CATT) in the United States, the Alternative  
126 Treatments to Inhibit VEGF in Age-related Choroidal Neovascularization (IVAN) in the United  
127 Kingdom, the Groupe d'Etude Français Avastin versus Lucentis dans la DMLA néovasculaire  
128 (GEFAL) in France, the Multicenter Anti-VEGF Trial in Austria (MANTA), Lucentis Compared to  
129 Avastin Study (LUCAS) in Norway, and Bevacizumab and Ranibizumab in Age-related Macular  
130 Degeneration (BRAMD) in the Netherlands.<sup>4-12</sup> In 2011, CATT was the first of the trials to  
131 provide 1-year results.<sup>4</sup> The mean change in visual acuity under treatment with bevacizumab  
132 was non-inferior to the mean change in visual acuity under treatment with ranibizumab. The  
133 results on efficacy from the other multicenter clinical trials have been consistent with no

134 difference or only a small difference in change in visual acuity between drugs after the initiation  
135 of treatment; a recent meta-analysis yielded a mean difference (95% confidence interval [CI]) of  
136 -0.5 letters (-1.6 to +0.6), with a negative difference indicating less improvement in eyes treated  
137 with bevacizumab.<sup>13</sup>

138         However, the results from one of the clinical trials raised concerns on the safety of  
139 bevacizumab relative to ranibizumab. In CATT, the proportion of patients with 1 or more  
140 systemic serious adverse events (SAEs) at 1 year was higher with bevacizumab than  
141 ranibizumab (24.1% vs. 19.0%; adjusted relative risk, 1.29; 95% CI [1.01, 1.66]) and the  
142 elevated risk persisted at 2 years (39.9% vs. 31.7%; adjusted relative risk, 1.30; 95% CI [1.07,  
143 1.57]; P=0.009).<sup>4,5</sup> Rates of death and arteriothrombotic events were similar for the two drugs.  
144 As the results from other clinical trials became available, several groups of investigators  
145 performed meta-analyses of overall SAEs and specific adverse events based on the aggregate  
146 data.<sup>13-19</sup> The most comprehensive analysis of SAEs was a Cochrane review led by Moja  
147 consisting of 3665 patients with 3356 from the 6 multicenter clinical trials noted above and 309  
148 patients from 3 smaller-scale studies.<sup>15</sup> The combined risk ratio for 1 or more systemic adverse  
149 events was 1.08, 95% CI (0.90, 1.31). Similar to the researchers conducting previous meta-  
150 analyses, Moja et al concluded that there was no strong evidence of a difference in risk but that  
151 the data available was not sufficient to rule out clinically important differential risks, particularly  
152 for specific adverse events.

153         The purpose of the present investigation was to use individual patient data, rather than  
154 aggregate data, from the large-scale multicenter clinical trials evaluating bevacizumab and  
155 ranibizumab for treatment of neovascular age-related macular degeneration to estimate the  
156 relative risk of serious systemic adverse events and selected specific SAEs adjusted for  
157 prognostic baseline variables. Although randomization is expected to provide treatment groups  
158 that are balanced on predisposing conditions, small imbalances on strong prognostic factors  
159 such as age, smoking, hypertension, and use of anti-coagulant medications can artificially

160 inflate or deflate the difference in risk between the two drugs. Accounting for covariates also  
161 may increase the precision of the estimates of the relative risk.

## 162 **METHODS**

### 163 **Clinical Trials Included**

164           Investigators for a recent Cochrane aggregate meta-analysis of the relative efficacy and  
165 safety of intravitreal bevacizumab and ranibizumab searched electronic bibliographic databases  
166 and clinical trial registries as of March 14, 2014 and used hand searching to identify 5249  
167 records that might address the topic.<sup>13</sup> Nine trials were identified by the Cochrane investigators.  
168 We targeted for this review the six multicenter, randomized clinical trials that compared  
169 bevacizumab to ranibizumab, reported counts for patients with 1 or more SAEs, had at least 1  
170 patient reported to have an SAE, and had results published or presented at a national meeting  
171 by December 2015. Eligibility criteria for all the trials specified enrollment of eyes with active  
172 neovascularization.

### 173 **Specification of Outcomes and Effect Measures**

174           The primary outcome for the review was the percentage of patients experiencing 1 or  
175 more SAEs as defined by the Food and Drug Administration of the United States and the  
176 European Medicines Agency.<sup>20,21</sup> This definition includes all deaths, life-threatening events,  
177 hospitalizations, events resulting in persistent or significant disability, important medical events,  
178 and congenital anomalies. Secondary outcomes were the specific SAEs of death,  
179 arteriothrombotic events as defined by the Antiplatelet Trialists' Collaboration (APTIC), events  
180 previously associated with systemic anti-VEGF treatment (arteriothrombotic events [including  
181 but not limited to myocardial, cerebellar, and cerebral ischemia and infarction, coronary artery  
182 occlusion, transient ischemic attack, cerebrovascular accidents, and embolism], systemic  
183 hemorrhage [including duodenal, gastric, gastrointestinal, rectal, respiratory tract, urogenital,  
184 cerebral, intracranial hemorrhage and hematoma], cardiac failure [including congestive heart  
185 failure], venous thrombotic events [including pulmonary embolism, deep vein thrombosis, and



186 thrombosis], hypertension [including hypertensive heart disease and accelerated hypertension],  
187 vascular death), and events not previously associated with systemic anti-VEGF treatment.<sup>22-24</sup>  
188 Because of an imbalance reported from CATT, gastrointestinal hemorrhages were also  
189 summarized. The difference in risk was summarized by the relative risk (hazard ratio) and the  
190 associated 95% confidence interval.

## 191 **Data Collection and Statistical Analysis**

192 The Coordinating Center for CATT managed the data and performed the statistical  
193 analyses for the review. The lead author or primary contact person as listed in a registry of  
194 clinical trials was invited to provide individual patient data. Data were to be provided in two  
195 electronic data files containing only de-identified data. The first file contained age at enrollment,  
196 gender, drug (bevacizumab or ranibizumab), dosing regimen (pro re nata, monthly, or treat-and-  
197 extend), study eye (right or left), smoking status at baseline (current, past, or never), diabetes at  
198 baseline (yes or no), use of medications for hypertension at baseline (yes or no), treatment of  
199 the fellow eye with anti-VEGF drugs during the study period (drug and duration of use), use of  
200 aspirin at baseline (yes or no), use of an anti-coagulant at baseline (yes or no), and number of  
201 days between enrollment and the last date of data collection for SAEs. The individual patient  
202 characteristics at baseline were chosen because they are known to be strong prognostic factors  
203 for one or more of the outcomes of interest. The second file contained one record for each SAE  
204 and included the number of days between study enrollment and the SAE, the Medical Dictionary  
205 for Regulatory Activities (MedDRA) code number, and MedDRA preferred term for the SAE.  
206 The period of observation was 2 years after study entry for CATT and IVAN and 1 year for the  
207 other 4 studies.

208 A two-stage approach was used for each meta-analysis.<sup>25,26</sup> In the first stage, a Cox  
209 proportional hazards model of the outcome of interest was used for each individual clinical trial  
210 to provide a relative risk adjusted for baseline prognostic factors and to provide the associated  
211 95% confidence interval for the risk of using bevacizumab compared to using ranibizumab.

212 Only the first observation of the outcome of interest was included in the analysis. The Cox  
213 models included dosing regimen (for CATT and IVAN only because these trials include both  
214 monthly and as-needed regimens), age, gender, smoking status, diabetes status, use of  
215 medications for or a diagnosis of hypertension, use of aspirin, and use of anti-coagulants when  
216 data for these variables were available. For the second stage, OpenMeta[Analyst] statistical  
217 software for meta-analyses was used to produce a weighted average of the trial specific relative  
218 risk from the first stage ([http://www.cebm.brown.edu/open\\_meta/](http://www.cebm.brown.edu/open_meta/) accessed 10/20/2015).  
219 Random effects models using maximum likelihood estimation were chosen to reflect both the  
220 within-study variability (95% CIs estimated in stage 1) and the between-study variability (the  
221 difference between the point estimates from stage 1 and the pooled estimate).<sup>27</sup> Heterogeneity  
222 among trial results was evaluated with the  $I^2$  statistic. For purposes of comparison, an  
223 unadjusted meta-analysis was performed with OpenMeta[Analyst] using aggregate data as for  
224 stage 2 of the adjusted meta-analysis. Individual patient data were not provided from MANTA.<sup>9</sup>  
225 As a secondary analysis, the unadjusted risk estimates for 1 or more SAEs and for death from  
226 based on the publication of 1-year MANTA results were used for the second stage of the  
227 adjusted meta-analysis. Because the conversion from the published data to the other outcomes  
228 of interest could not be made without more details on type of the SAEs, no secondary analyses  
229 were performed for the other outcomes of interest.

230 The data files from the 5 clinical trial groups providing individual patient data were checked  
231 for completeness of the data requested and for consistency with published aggregate results.  
232 Data files for CATT, IVAN, GEFAL, and LUCAS, matched the published aggregate findings for  
233 the safety analysis with respect to number of patients and number of patients with 1 or more  
234 systemic SAE in each treatment group. Serious ocular adverse events were not counted as  
235 systemic adverse events for this analysis.<sup>11</sup> There was 1 less patient assigned to bevacizumab  
236 in the data files from BRAMD than reported in published results.<sup>12</sup> Nine patients in LUCAS, who  
237 had no serious adverse events, were excluded from the efficacy analysis in LUCAS because of

238 serious non-compliance with the treatment protocol and were excluded from the adjusted  
239 analysis in this report. When data on use of medications for hypertension were not available,  
240 data on a diagnosis of hypertension were used instead.

241

242 **RESULTS**

243 The baseline data available from each clinical trial are summarized in Table 1. Among the  
244 five clinical trials providing individual patient data, age, gender, diabetes status, and  
245 hypertension status (as defined in the parent trial) were available in all trials. There were only  
246 small imbalances between the bevacizumab and ranibizumab groups on the baseline  
247 characteristics.

248 There were 403 (26.6%) patients among 1513 treated with bevacizumab and 366 (23.8%)  
249 among 1539 treated with ranibizumab who had 1 or more systemic SAE. The numbers of  
250 patients in each treatment group in each study are provided in Table 2. Adjusted meta-analysis  
251 results are shown in Figure 1 and compared to the unadjusted results in Table 3. The pooled  
252 adjusted relative risk for bevacizumab compared to ranibizumab was 1.06, 95% CI (0.84, 1.35).  
253 The adjusted relative risk differs little from the unadjusted relative risk of 1.08. When the  
254 aggregate data from MANTA was included in the adjusted analysis, the relative risk was 1.09,  
255 95% CI (0.89,1.35). The adjusted relative risk for death was 0.99, 95% CI (0.69, 1.43) (Figure 2  
256 available at <http://www.aaojournal.org>). When the aggregate data from MANTA was included in  
257 the adjusted analysis, the relative risk was 1.01, 95% CI (0.71,1.45). Estimated risk for APTC  
258 arteriothrombotic events was lower for bevacizumab (0.89) but with the 95% confidence interval  
259 spanning (0.62, 1.28) (Figure 3 available at <http://www.aaojournal.org>). The adjusted relative  
260 risks for systemic SAEs related to anti-VEGF treatment and those not related to anti-VEGF  
261 treatment were nearly identical (1.10 and 1.11, respectively) (Figures 4, 5 available at  
262 <http://www.aaojournal.org>). There were too few gastrointestinal hemorrhages reported (1 for  
263 ranibizumab in GEFAL, 1 for ranibizumab in LUCAS) to add any meaningful information to the  
264 imbalance reported in CATT (7 for bevacizumab, 2 for ranibizumab).

265 The percentage of the variability in relative risks due to heterogeneity across studies, rather  
266 than to sampling error, is given by the  $I^2$  statistic in each of the Figures. Heterogeneity was  
267 moderate for the proportion of patients with 1 or more systemic SAE (50%) and systemic SAEs

268 not related to systemic anti-VEGF treatments (59%), substantially less (30%) for  
269 arteriothrombotic events, and 0% for death and events related to systemic anti-VEGF  
270 treatment.

## 271 **DISCUSSION**

272 The individual patient data meta-analyses yielded no significant differences in risk of  
273 systemic SAEs between bevacizumab and ranibizumab. Thus, while the point estimate for  
274 relative risk indicated an approximate 10% increase with bevacizumab relative to ranibizumab  
275 for most categories of SAE, a similar 10% decrease for arteriothrombotic events was found.  
276 However, the confidence intervals for the relative risks spanned values, both for increased risk  
277 and decreased risk with bevacizumab, that would be clinically important for events such as  
278 death, cerebro- and cardio-vascular events, and cancer. The adjusted analyses produced  
279 results indicating less risk with bevacizumab than in the unadjusted analyses; however, the  
280 reduction was minor.

281 Now that 10 years have passed since the introduction of bevacizumab and ranibizumab for  
282 treatment of neovascular age-related macular degeneration, new head-to-head trials are no  
283 longer likely to be performed. Although the recent Cochrane meta-analyses of systemic SAEs  
284 and the unadjusted meta-analysis based on aggregated data reported here did not include the  
285 same set of trials, they yielded similar relative risks of approximately 1.1 for 1 or more SAEs  
286 through 1 or 2 years. A trial in India of 120 patients with no adverse events reported,<sup>28</sup> a trial in  
287 the United States of 28 patients with 2 deaths reported in 20 patients treated with bevacizumab  
288 (1 meckel cell carcinoma and 1 cause unknown),<sup>29</sup> and a trial in Germany registered on  
289 ClinicalTrials.gov but without presentation at a national meeting or in a peer-reviewed journal  
290 were included in the meta-analysis by Moja but not the current one.<sup>30</sup> Moja noted that, in a  
291 personal communication, the German researchers reported SAEs in 21% (22/107) of patients  
292 treated with bevacizumab and in 11% (6/54) of patients treated with ranibizumab.<sup>15</sup> Because

293 small imbalances on strong risk factors such as age, smoking history, hypertension, diabetes,  
294 and aspirin and anti-coagulant use can result in biased estimates of difference in risk, this  
295 review was initiated to find out whether such imbalances might have influenced the result of  
296 meta-analyses that used aggregate data from the clinical trials.

297       There are some weaknesses in this meta-analysis. First, all the trials were of modest size  
298 (<1200 patients each). Second, although there was a common definition of an SAE across  
299 trials, the methods of ascertaining the occurrence of an SAE may have varied among trials.  
300 Third, the dosing intervals varied across the trials. Comparisons between the drugs were made  
301 within each dosing regimen, but the monthly, as needed, and treat and extend approaches were  
302 used among the trials. Fourth, individual patient data could not be obtained for one of the  
303 clinical trials and only a secondary analysis using aggregate data from that trial could be  
304 performed. Fifth, there was moderate heterogeneity across the 5 trials in the proportion of  
305 patients with 1 or more systemic SAE and systemic SAEs not related to systemic anti-VEGF  
306 treatments, due mainly to results from LUCAS. We attribute this to random variation because  
307 eligibility, dose, and visual acuity results in LUCAS were similar to those in the other trials and  
308 the ascertainment of SAEs was made by staff masked to study drug. In addition to the strength  
309 of the study of being able to account for possible imbalances in prognostic factors through use  
310 of patient-level data, the present study employed survival analysis methods that incorporate not  
311 only the occurrence of an SAE but also the time since initiation of treatment, thus providing a  
312 more precise assessment of differential risk than simply comparing the cumulative numbers at  
313 either 1 or 2 years of follow-up.

314       The meta-analyses on individual patient data in this review, as well as previous meta-  
315 analyses on aggregate data, support the conclusion that large differences between  
316 bevacizumab and ranibizumab in risk of systemic serious adverse events; i.e., relative risks of  
317  $\geq 1.5$ , are unlikely. Although the estimated relative risks indicate an approximate 10% increase

318 for most types of SAEs and a 10% decrease in arteriothrombotic events for bevacizumab, these  
319 point estimates have confidence intervals that include up to a 50% increase or decrease in risk.  
320 In the absence of additional large-scale clinical trials, further investigation of the differential risk  
321 of these anti-VEGF agents can be carried out only through epidemiologic surveillance using  
322 administrative or healthcare databases.

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FIGURE LEGEND

Figure 1. Forest Plot for the Adjusted Relative Risk for 1 or More Systemic Serious Adverse Events for Bevacizumab Compared to Ranibizumab.

Studies	Estimate (95% C.I.)
CATT	1.278 (1.055, 1.548)
IVAN	1.043 (0.764, 1.424)
GEFAL	1.154 (0.671, 1.985)
LUCAS	0.569 (0.354, 0.915)
BRAMD	1.337 (0.785, 2.277)
<b>Overall (I<sup>2</sup>=49.63 % , P=0.036)</b>	<b>1.064 (0.841, 1.346)</b>

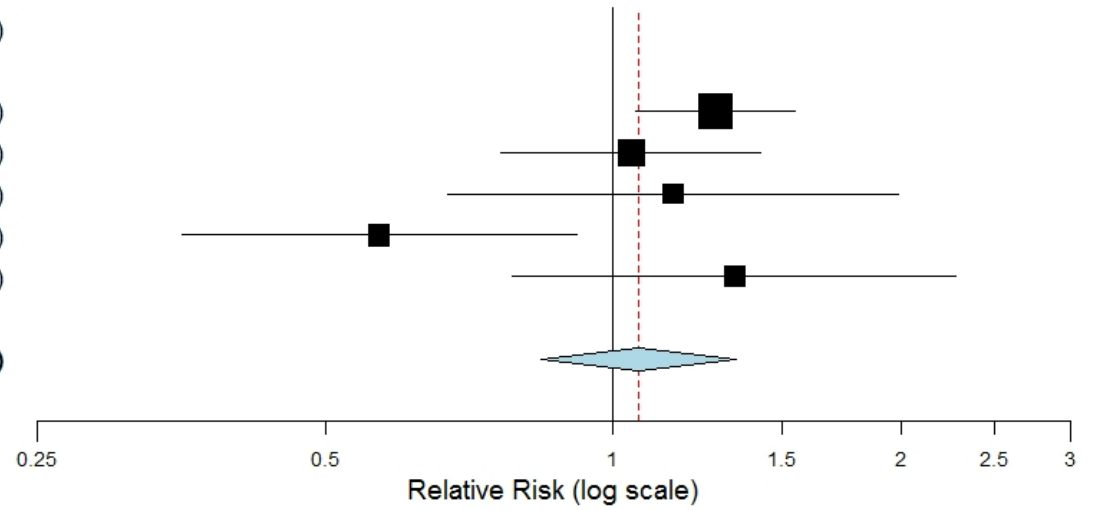


Figure 2. (Supplementary) Forest Plot for the Adjusted Relative Risk for Death for Bevacizumab Compared to Ranibizumab

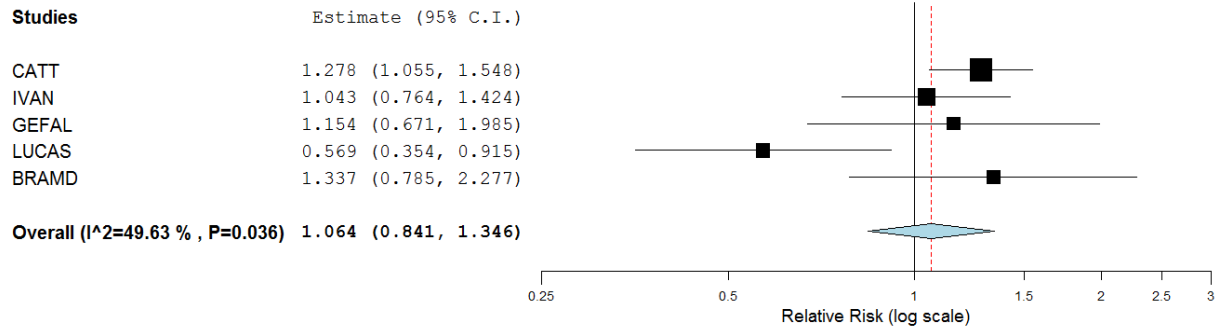


Figure 3. (Supplementary) Forest Plot for the Adjusted Relative Risk for Antiplatelet Trialists' Collaboration (APTC) Arteriothrombotic Event as for Bevacizumab Compared to Ranibizumab

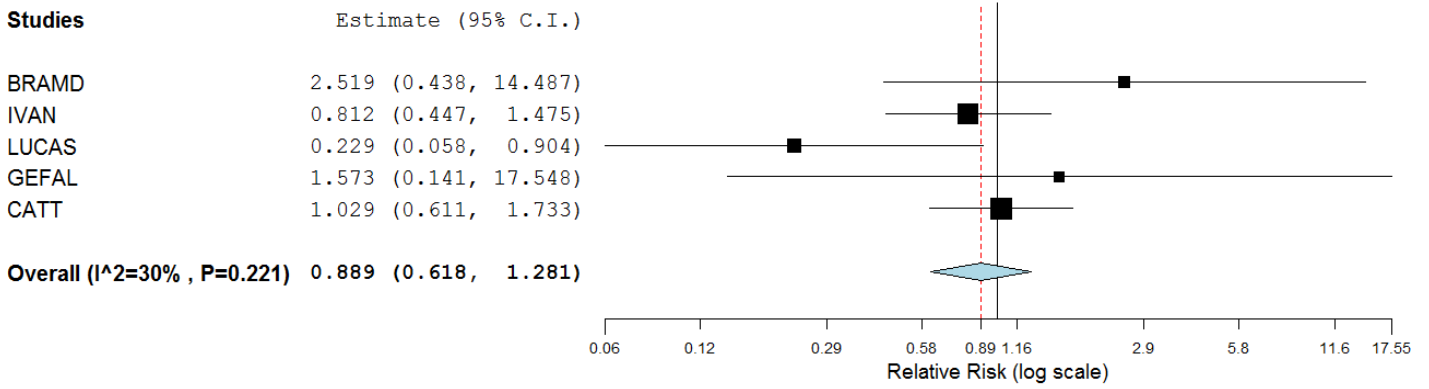


Figure 4. (Supplementary) Forest Plot for the Adjusted Relative Risk for Events Related to Anti-VEGF

Treatment for Bevacizumab Compared to Ranibizumab

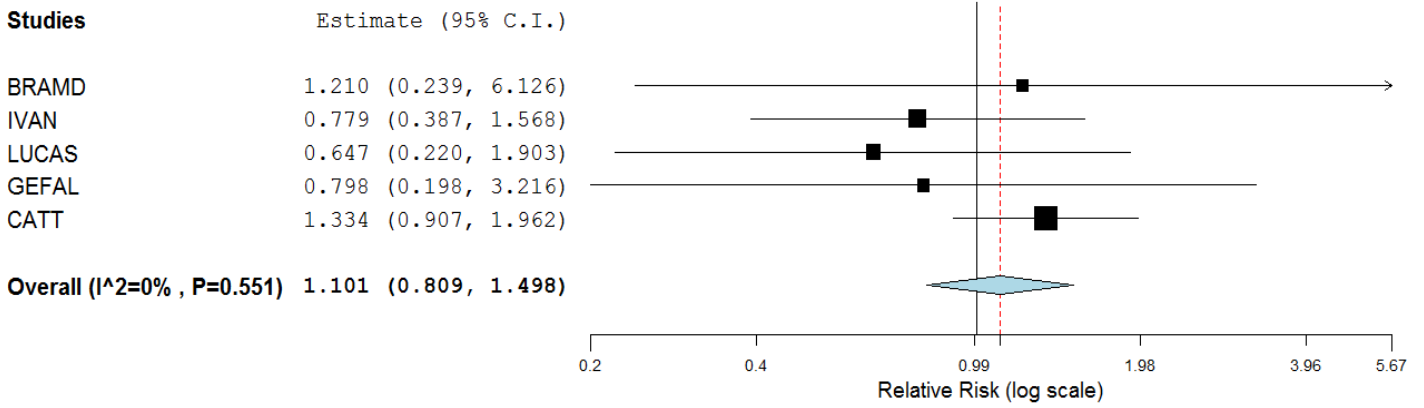




Figure 5. (Supplementary) Forest Plot for the Adjusted Relative Risk for Events Not Related to Anti-VEGF Treatment for Bevacizumab Compared to Ranibizumab

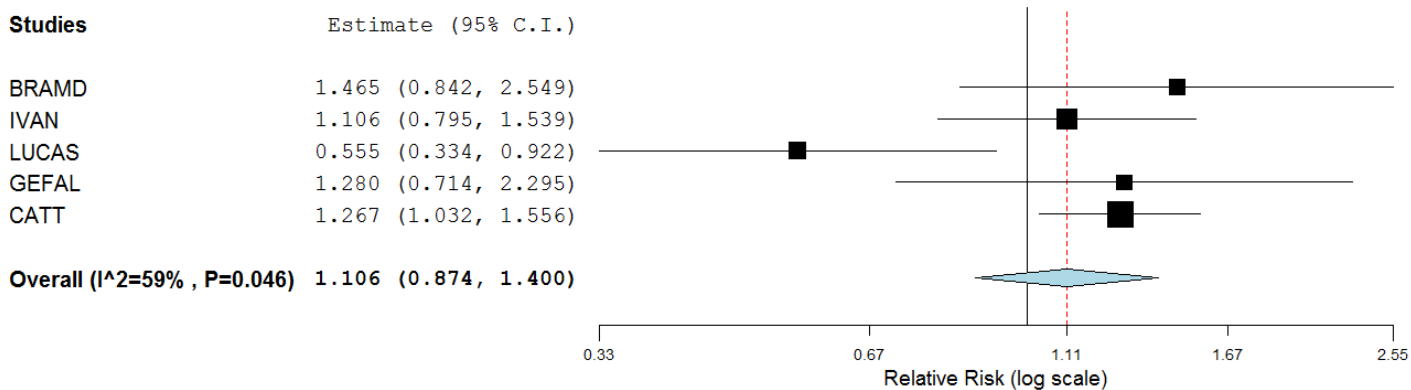


Table 1. Distribution of Baseline Characteristics Available from Each Clinical Trial by Drug

Characteristic	Clinical Trial					Overall
	CATT	IVAN	GEFAL	LUCAS	BRAMD	
Drug, N						
Bevacizumab	586	296	246	220	165	1513
Ranibizumab	599	314	239	221	166	1539
Age (yrs), mean						
Bevacizumab	79.7	77.7	79.5	78.6	77.1	78.8
Ranibizumab	78.8	77.8	79.0	78.0	77.0	78.3
Female (%)						
Bevacizumab	62.1	61.2	62.2	70.6	55.2	62.4
Ranibizumab	61.4	58.9	70.3	64.2	55.4	62.0
Current or past smoker (%)						
Bevacizumab	57.7	62.5	NA	55.5	54.6	58.0
Ranibizumab	56.8	63.7	NA	52.0	51.8	57.0
Diabetic (%)						
Bevacizumab	18.3	9.1	11.8	7.0	10.9	13.0
Ranibizumab	16.7	11.8	10.9	6.4	12.7	12.9
Hypertension (%)						
Bevacizumab	70.3	61.2	61.8	57.9	57.0	63.9
Ranibizumab	68.6	59.9	53.1	53.2	66.9	62.0
Aspirin (%)						
Bevacizumab	50.9	31.4	NA	29.0	NA	41.3
Ranibizumab	45.9	27.1	NA	30.3	NA	37.7
Anticoagulant (%)						
Bevacizumab	16.6	4.4	NA	7.7	NA	11.5
Ranibizumab	17.7	6.1	NA	9.1	NA	12.8

NA = Not available.

Table 2. Systemic Serious Event and its Type from Each Clinical Trial by Drug

Characteristic	CATT	IVAN	GEFAL	LUCAS	BRAMD	Total
<b>N</b>						
Bevacizumab	586	296	246	220	165	1513
Ranibizumab	599	314	239	221	166	1539
<b>≥1 SAE: n (%)</b>						
Bevacizumab	234 (39.9%)	80 (27.0%)	30 (12.2%)	29 (13.2%)	30 (18.2%)	403 (26.6%)
Ranibizumab	190 (31.7%)	81 (25.8%)	24 (10.0%)	45 (20.4%)	26 (15.7%)	366 (23.8%)
<b>Death: n (%)</b>						
Bevacizumab	36 (6.1%)	15 (5.1%)	2 (0.8%)	4 (1.8%)	1 (0.6%)	58 (3.8%)
Ranibizumab	32 (5.3%)	15 (4.8%)	3 (1.3%)	7 (3.2%)	1 (0.6%)	58 (3.8%)
<b>APTC: n (%)</b>						
Bevacizumab	29 (4.9%)	20 (6.8%)	2 (0.8%)	3 (1.4%)	4 (2.4%)	58 (3.8%)
Ranibizumab	28 (4.7%)	25 (8.0%)	1 (0.4%)	9 (4.1%)	2 (0.9%)	65 (4.2%)
<b>VEGF-related: n (%)</b>						
Bevacizumab	62 (10.6%)	14 (4.7%)	4 (1.6%)	6 (2.7%)	3 (1.8%)	89 (5.9%)
Ranibizumab	45 (7.5%)	19 (6.1%)	4 (1.7%)	8 (3.6%)	3 (1.8%)	79 (5.1%)
<b>Not VEGF-related: n (%)</b>						
Bevacizumab	202 (34.4%)	73 (24.7%)	27 (11.0%)	25 (11.4%)	29 (17.6%)	356 (23.5%)
Ranibizumab	170 (28.4%)	70 (22.2%)	20 (8.4%)	40 (18.1%)	23 (13.9%)	323 (21.0%)

NA is not available.

APTC = Antiplatelet Trialists' Collaboration arteriothrombotic events

VEGF is vascular endothelial growth factor

Table 3. Summary of Estimated Relative Risks of Systemic Serious Adverse Events after Treatment with Bevacizumab Compared to Ranibizumab

Systemic Serious Event Type	Bevacizumab	Ranibizumab	Relative Risk (95% CI)		P-value Adjusted Model
	(N=1513) With Event n (%)	(N=1539) With Event n. (%)	Unadjusted	Adjusted	
≥1 event	403 (26.6%)	366 (23.8%)	1.08 (0.90,1.30)	1.06 (0.84,1.35)	0.61
Death	58 ( 3.8%)	58 ( 3.8%)	1.03 (0.72,1.48)	0.99 (0.69,1.43)	0.97
APTC	58 ( 3.8%)	65 ( 4.2%)	0.93 (0.66,1.32)	0.89 (0.62,1.28)	0.53
VEGF-related	89 ( 5.9%)	79 ( 5.1%)	1.16 (0.86,1.56)	1.10 (0.81,1.50)	0.54
Not VEGF-related	356 (23.5%)	323 (21.0%)	1.14 (1.00,1.30)	1.11 (0.87,1.40)	0.40

APTC is Antiplatelet Trialists' Collaboration arteriothrombotic events

VEGF is vascular endothelial growth factor

## Precis

A meta-analysis on individual patient data supports the conclusion that large differences between bevacizumab and ranibizumab in risk of systemic serious adverse events; i.e., relative risks of  $\geq 1.5$ , are unlikely.

## Credit Roster for the Bevacizumab-Ranibizumab International Trials Group

### Credit Roster for the BRAMD

#### Clinical Centers (Ordered by Number of Patients enrolled)

Certified Roles at Clinical Centres: Clinic Coordinator (CC), Data Entry Staff (DE), Participating Ophthalmologist (O), Ophthalmic Photographer (OP); Optical Coherent Tomography Technician (OCT), Principal Investigator (PI), Refractionst (R), Visual Acuity Examiner (VA)

**AMC:** Reinier O. Schlingemann (PI), Frank D. Verbraak (O), Marie van Schooneveld (O), Monique Wezel (CC), Henk Stam (VA/R), Christa Jansen-Kok (VA/R/OCT/DE), Ans Althoff (VA/R/OCT) Douwe Bakker (CC/VA/R/OCT/DE), Anette van der Zee (DE)

**Erasmus MC:** Johannes R. Vingerling (PI), Naus- Postema (O), De Roo Hertoge (O), C. Klaver (O), E. Kilic (O), Yvonne Noordzij (CC/DE/VA/R/OCT), Jeanette Noordzij (CC/DE/VA/R/OCT), Anjo Vermij (VA/R), Ada Hooghart (VA/R)

**LUMC:** Greetje Dijkman (PI), Ingrid Boesten (CC/DE/VA/R), C. Kiewiet de Jonge (VA/R), M. Kromhart-de Haas (VA/R), Cora Mollinger (VA/R), J.W. Zwaan (VA/R), Lou Brink (VA/R), Anneke Boolman (VA/R)

**UMCG:** Johanna Hooymans (PI), Nicole Kamminga (O), Angela Huiskamp (O), Postma (O) Marijke Meinen (CC/DE/R/VA), L. Uwantege (VA/R), H.R. Luurtsema (VA/R), J.F. Eisses (VA/R/OCT), Westra (OP), Ronardel de Lavalette (O), Van De Pol (O), J van Nieuwpoort, j, Tiggelman, T Ezenga, Brouwen, F. Schuit, Ufkes, H. Egberts, G de Fretes

**UMCN:** Carel B. Hoyng (PI), Agnes de Vries (CC), Elke Huntink (CC), Asha Kalisingh (CC), Liesbeth Hoeks (VA/R), Hans Hermans (VA/R), Angeline Rottelveel (VA/R/OCT), J. Weeda (VA/R/OP/OCT), Chantal Van Ast (OCT)

**OMC Haarlem:** Remko van Hierdam (VA/R), Annemieke Coops (VA/R), R.W. Hierden (OCT), Kees Corstanje (VA/R), Madelon Jansen (VA/R/OCT), Nikki van denBerg (VA/R/OCT)

**Zonnestraal Hilversum:** Bianca Ban Leeuwen (VA/R/DE), Gerrie Dantuma (VA/R/OCT), Serge Koning (VA/R/OCT), Maurice van Baekel (OCT/VA/R), Mike Selders (VA/R/OCT), Martine Zandee (VA/R/OCT), Maaïke Goudriaan (VA/R/OCT), Annemarieke Langen (VA/R/OCT), Robert W.H van de Mortel (VA/R/OCT)

#### Resource Centers

**Chairman's Office and Coordinating Center**(Academic Medical Center, Amsterdam, the Netherlands): R.O. Schlingemann, MD Phd(Chair/PI); F.D Verbraak, MD (Vice-Chair; Academic Medical Center, Amsterdam, the Netherlands) M.G.W Dijkgraaf (methodologist), R. De Haan (methodologist), A.H. Zwinderman, SCP (Biostatistician); A.M.E. Schauwvlieghe, MD (Medical

Monitor); Selma Mehmedovic (Protocol Monitor); Jaqueline Van Daele (CRA), Irmgard Corten (Systems Analyst); Eric Veenstra (Financial Administrator); Yolanda Strubel (Database developer)

**OCT Reading Center:** F.D. Verbraak, MD PhD(PI), A.M.E. Schauwvlieghe, MD (reader), Douwe Bakker (reader)

**Fundus Photograph Reading Center** (Reading Centre, Moorfields Eye Hospital): Peto T, MD PhD(PI); Peter Blows (reader)

**Data and Safety Monitoring Committee:** Ringens PJ, MD, Phd (chair); R. Geskus (Biostatistician), MD; R. Van Leeuwen, MD PhD

## **Credit Roster for Comparison of AMD Treatments Trials (CATT)**

### **Clinical Centers (Ordered by Number of Patients Enrolled)**

**Certified Roles at Clinical Centers:** Clinic Coordinator (CC), Data Entry Staff (DE), Participating Ophthalmologist (O), Ophthalmic Photographer (OP); Optical Coherent Tomography Technician (OCT), Principal Investigator (PI), Refractionist (R), Visual Acuity Examiner (VA)

**Vitreoretinal Surgery, PA (Edina, MN):** David F. Williams, MD (PI); Sara Beardsley, COA (VA/R); Steven Bennett, MD (O); Herbert Cantrill, MD (O); Carmen Chan-Tram, COA (VA/R); Holly Cheshier, CRA, COT, OCTC (OP); Kathryn Damato, COT (VA); John Davies, MD (O); Sundeep Dev, MD (O); Julianne Enloe, CCRP, COA (CC); Gennaro Follano (OP/OCT); Peggy Gilbert, COA (VA/R); Jill Johnson, MD (O); Tori Jones, COA (OCT); Lisa Mayleben, COMT (CC/VA/R/OCT); Robert Mittra, MD (O); Martha Moos, COMT, OSA (VA/R); Ryan Neist, COMT (VA/R); Neal Oestreich, COT (CC); Polly Quiram, MD (O); Robert Ramsay, MD (O); Edwin Ryan, MD (O); Stephanie Schindeldecker, OA (VA/R); John Snater, COA (VA); Trenise Steele, COA (VA); Dwight Selders, COA (VA/R); Jessica Tonsfeldt, AO (OP/OCT); Shelly Valardi, COT (VA/R).

**Texas Retina Associates (Dallas, TX):** Gary Edd Fish, MD (PI); Hank A. Aguado, CRA (OP/OCT); Sally Arceneaux (CC/VA/R); Jean Arnwine (CC); Kim Bell, COA (VA/R); Tina Bell (CC/OCT); Bob Boleman (OP); Patricia Bradley, COT (CC); David Callanan, MD (O); Lori Coors, MD (O); Jodi Creighton, COA (VA/R); Timothy Crew, COA (OCT); Kimberly Cummings (OP/OCT); Christopher Dock (OCT); Karen Duignan, COT (VA/R); Dwain Fuller, MD (O); Keith Gray (OP/OCT); Betsy Hendrix, COT, ROUB (OCT); Nicholas Hesse (OCT); Diana Jaramillo, COA (OCT); Bradley Jost, MD (O); Sandy Lash (VA/R); Laura Lonsdale, CCRP (DE); Michael Mackens (OP/OCT); Karin Mutz, COA (CC); Michael Potts (VA/R); Brenda Sanchez (VA/R); William Snyder, MD (O); Wayne Solley, MD (O); Carrie Tarter (VA/R); Robert Wang, MD (O); Patrick Williams, MD (O).

**Southeastern Retina Associates (Knoxville, TN):** Stephen L. Perkins, MD (PI); Nicholas Anderson, MD (O); Ann Arnold, COT (VA/R); Paul Blais (OP/OCT); Joseph Googe, MD (O); Tina T. Higdon, (CC); Cecile Hunt (VA/R); Mary Johnson, COA (VA/R); James Miller, MD (O); Misty Moore (VA/R); Charity K. Morris, RN (CC); Christopher Morris (OP/OCT); Sarah Oelrich, COT (OP/OCT); Kristina Oliver, COA (VA/R); Vicky Seitz, COT (VA/R); Jerry Whetstone (OP/OCT).

**Retina Vitreous Consultants (Pittsburgh, PA):** Bernard H. Doft (PI); Jay Bedel, RN, (CC); Robert Bergren, MD (O); Ann Borthwick (VA/R); Paul Conrad, MD, PHD (O); Amanda Fec (OCT); Christina

Fulwylie (VA/R); Willia Ingram (DE); Shawnique Latham (VA/R); Gina Lester (VA/R); Judy Liu, MD (O); Louis Lobes, MD (O); Nicole M. Lucko, (CC); Holly Mechling (CC); Lori Merlotti, MS, CCRC (CC); Keith McBroom (OCT); Karl Olsen, MD (O); Danielle Puskas, COA (VA/R); Pamela Rath, MD (O); Maria Schmucker (CC); Lynn Schueckler (OCT); Christina Schultz (CC/VA/R); Heather Shultz (OP/OCT); David Steinberg, CRA (OP/OCT); Avni Vyas, MD (O); Kim Whale (VA/R); Kimberly Yeckel, COA, COT (VA/R).

**Ingalls Memorial Hospital/Illinois Retina Associates (Harvey, IL):** David H. Orth, MD (PI); Linda S. Arredondo, RN (CC/VA); Susan Brown (VA/R); Barbara J. Ciscato (CC/VA); Joseph M. Civantos, MD (O); Celeste Figliulo (VA/R); Sohail Hasan, MD (O); Belinda Kosinski, COA (VA/R); Dan Muir (OP/OCT); Kiersten Nelson (OP/OCT); Kirk Packo, MD (O); John S. Pollack, MD (O); Kourous Rezaei, MD (O); Gina Shelton (VA); Shannya Townsend-Patrick (OP/OCT); Marian Walsh, CRA (OP/OCT).

**West Coast Retina Medical Group, Inc. (San Francisco, CA):** H. Richard McDonald, MD (PI); Nina Ansari (VA/R/OCT); Amanda Bye, (OP/OCT); Arthur D. Fu, MD (O); Sean Grout (OP/OCT); Chad Indermill (OCT); Robert N. Johnson, MD (O); J. Michael Jumper, MD (O); Silvia Linares (VA/R); Brandon J. Lujan, MD (O); Ames Munden (OP/OCT); Meredith Persons (CC); Rosa Rodriguez (CC); Jennifer M. Rose (CC); Brandi Teske, COA (VA/R); Yesmin Urias (OCT); Stephen Young (OP/OCT).

**Retina Northwest, P.C. (Portland, OR):** Richard F. Dreyer, MD (PI); Howard Daniel (OP/OCT); Michele Connaughton, CRA (OP/OCT); Irvin Handelman, MD (O); Stephen Hobbs (VA/R/OCT); Christine Hoerner (OP/OCT); Dawn Hudson (VA/R/OCT); Marcia Kopfer, COT (CC/VA/R/OCT); Michael Lee, MD (O); Craig Lemley, MD (O); Joe Logan, COA (OP/OCT); Colin Ma, MD (O); Christophe Mallet (VA/R); Amanda Milliron (VA/R); Mark Peters, MD (O); Harry Wohlsein, COA (OP).

**Retinal Consultants Medical Group, Inc. (Sacramento, CA):** Joel A. Pearlman, MD, PHD (PI); Margo Andrews (OP/OCT); Melissa Bartlett (OCT); Nanette Carlson (CC/OCT); Emily Cox (VA/R); Robert Equi, MD (O); Marta Gonzalez (VA/R/OCT); Sophia Griffin (OP/OCT); Fran Hogue (VA/R); Lance Kennedy (OP/OCT); Lana Kryuchkov (OCT); Carmen Lopez (VA/R); Danny Lopez (OP/OCT); Bertha Luevano (VA/R); Erin McKenna, (CC); Arun Patel, MD (O); Brian Reed, MD (O); Nyla Secor (CC/OCT); Iris R. Sison (CC); Tony Tsai, MD (O); Nina Varghis, (CC); Brooke Waller (OCT); Robert Wendel, MD (O); Reina Yebra (OCT).

**Retina Vitreous Center, PA (New Brunswick, NJ):** Daniel B. Roth, MD (PI); Jane Deinzer, RN (CC/VA/R); Howard Fine, MD MHSC (O); Flory Green (VA/R); Stuart Green, MD (O); Bruce Keyser, MD (O); Steven Leff, MD (O); Amy Leviton (VA/R); Amy Martir (OCT); Kristin Mosenthine (VA/R/OCT); Starr Muscle, RN (CC); Linda Okoren (VA/R); Sandy Parker (VA/R); Jonathan Prenner, MD (O); Nancy Price (CC); Deana Rogers (OP/OCT); Linda Rosas (OP/OCT); Alex Schlosser (OP/OCT); Loretta Studenko (DE); Thea Tantum (CC); Harold Wheatley, MD (O).

**Vision Research Foundation/Associated Retinal Consultants, P.C. (Royal Oak, MI):** Michael T. Trese, MD (PI); Thomas Aaberg, MD (O); Tina Bell (VA/R/OP/OCT); Denis Bezaire, CRA (OP/OCT); Craig Bridges, CRA (OP/OCT); Doug Bryant, CRA (OP/OCT); Antonio Capone, MD (O); Michelle Coleman, RN (CC); Christina Consolo, CRA, COT (OP/OCT); Cindy Cook, RN (CC); Candice DuLong (VA/R); Bruce Garretson, MD (O); Tracy Grooten (VA/R); Julie Hammersley, RN (CC); Tarek Hassan, MD (O); Heather Jessick (OP/OCT); Nanette Jones (VA/R/OP/OCT); Crystal Kinsman (VA/R); Jennifer Krumlauf (VA/R); Sandy Lewis, COT (VA/R/OP/OCT); Heather Locke (VA/R); Alan Margherio, MD (O); Debra Markus, COT (CC/VA/R/OP/OCT); Tanya Marsh, COA (OP/OCT); Serena Neal (CC); Amy Noffke, MD (O); Kean Oh, MD (O); Clarence Pence (OP/OCT); Lisa Preston (VA/R); Paul Raphaelian, MD (O); Virginia R. Regan, RN, CCRP (VA/R); Peter Roberts (OP/OCT); Alan Ruby, MD (O); Ramin Sarrafzadeh, MD, PHD (O); Marissa Scherf (OP/OCT); Sarita Scott (VA/R); Scott Sneed, MD (O);



Lisa Staples (CC); Brad Terry (VA/R/OP/OCT); Matthew T. Trese (OCT); Joan Videtich, RN (VA/R); George Williams, MD (O); Mary Zajechowski, COT, CCRC (CC/VA/R).

**The Retina Institute (St. Louis, MO):** Daniel P. Joseph, MD (PI); Kevin Blinder, MD (O); Lynda Boyd, COT (VA/R); Sarah Buckley (OP/OCT); Meaghan Crow (VA/R); Amanda Dinatale, (OCT); Nicholas Engelbrecht, MD (O); Bridget Forke (OP/OCT); Dana Gabel (OP/OCT); Gilbert Grand, MD (O); Jennifer Grillion-Cerone (VA/R); Nancy Holekamp, MD (O); Charlotte Kelly, COA (VA/R); Ginny Nobel, COT (CC); Kelly Pepple (VA/R); Matt Raeber, (OP/OCT); P. Kumar Rao, MD (O); Tammy Ressel, COT (VA/R); Steven Schremp (OCT); Merrilee Sgorlon (VA/R); Shantia Shears, MA (CC); Matthew Thomas, MD (O); Cathy Timma (VA/R); Annette Vaughn,(OP/OCT); Carolyn Walters, COT (CC/VA/R); Rhonda Weeks, CRC (CC/VA/R); Jarrod Wehmeier (OP/OCT); Tim Wright (OCT).

**The Retina Group of Washington (Chevy Chase, MD):** Daniel M. Berinstein, MD (PI); Aida Ayyad (VA/R); Mohammed K. Barazi, MD (O); Erica Bickhart (CC/VA/R); Tracey Brady (OCT); Lisa Byank, MA (CC); Alysia Cronise, COA (VA/R); Vanessa Denny (VA/R); Courtney Dunn (VA/R); Michael Flory (OP/OCT); Robert Frantz (OP/OCT); Richard A. Garfinkel, MD (O); William Gilbert, MD (O); Michael M. Lai, MD, PHD (O); Alexander Melamud, MD (O); Janine Newgen (VA/R); Shamekia Newton (CC); Debbie Oliver (CC); Michael Osman, MD (O); Reginald Sanders, MD (O); Manfred von Fricken, MD (O).

**Retinal Consultants of Arizona (Phoenix, AZ):** Pravin Dugel, MD (PI); Sandra Arenas (CC); Gabe Balea (OCT); Dayna Bartoli (OP/OCT); John Bucci (OP/OCT); Jennifer A. Cornelius (CC); Scheleen Dickens, (CC); Don Doherty (OP/OCT); Heather Dunlap, COA (VA/R); David Goldenberg, MD (O); Karim Jamal, MD (O); Norma Jimenez (OP/OCT); Nicole Kavanagh (VA/R); Derek Kunimoto, MD (O); John Martin (OP/OCT); Jessica Miner, RN (VA/R); Sarah Mobley, CCRC (CC/VA/R); Donald Park, MD (O); Edward Quinlan, MD (O); Jack Sipperley, MD (O); Carol Slagle (R); Danielle Smith (OP/OCT); Miguelina Yafchak (OCT); Rohana Yager, COA (OP/OCT).

**Casey Eye Institute (Portland, OR):** Christina J. Flaxel, MD (PI); Steven Bailey, MD (O); Peter Francis, MD, PHD (O); Chris Howell, (OCT); Thomas Hwang, MD (O); Shirley Ira, COT (VA/R); Michael Klein, MD (O); Andreas Lauer, MD (O); Teresa Liesegang, COT (CC/VA/R); Ann Lundquist, (CC/VA/R); Sarah Nolte (DE); Susan K. Nolte (VA/R); Scott Pickell (OP/OCT); Susan Pope, COT (VA/R); Joseph Rossi (OP/OCT); Mitchell Schain (VA/R); Peter Steinkamp, MS (OP/OCT); Maureen D. Toomey (CC/VA/R); Debora Vahrenwald, COT (VA/R); Kelly West (OP/OCT).

**Emory Eye Center (Atlanta, GA):** Baker Hubbard, MD (PI); Stacey Andelman, MMSC, COMT (CC/VA/R); Chris Bergstrom, MD (O); Judy Brower, COMT (CC/VA/R); Blaine Cribbs, MD (O); Linda Curtis (VA/R); Jannah Dobbs (OP/OCT); Lindreth DuBois, MED, MMSC, CO, COMT (CC/VA/R); Jessica Gaultney (OCT); Deborah Gibbs, COMT, CCRC (VA/R); Debora Jordan, CRA (OP/OCT); Donna Leef, MMSC, COMT (VA/R); Daniel F. Martin, MD (O); Robert Myles, CRA (OP); Timothy Olsen, MD (O); Bryan Schwent, MD (O); Sunil Srivastava, MD (O); Rhonda Waldron, MMSC, COMT, CRA, RDMS (OCT).

**Charlotte Eye, Ear, Nose & Throat Associates/Southeast Clinical Research (Charlotte, NC):** Andrew N. Antoszyk, MD (PI); Uma Balasubramaniam, COA (OCT); Danielle Brooks, CCRP (VA/R); Justin Brown, MD (O); David Browning, MD, PHD (O); Loraine Clark, COA (OP/OCT); Sarah Ennis, CCRC (VA/R); Susannah Held (OCT); Jennifer V. Helms, CCRC,(CC); Jenna Herby, CCRC (CC); Angie Karow, CCRP (VA/R); Pearl Leotaud, CRA (OP/OCT); Caterina Massimino (OCT); Donna McClain, COA (OP/OCT); Michael McOwen, CRA (OP/OCT); Jennifer Mindel, CRA, COA (OP/OCT); Candace Pereira, CRC (CC); Rachel Pierce, COA (VA/R); Michele Powers (OP/OCT); Angela Price, MPH, CCRC (CC); Jason Rohrer (CC); Jason Sanders, MD (O).

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**Mayo Clinic (Rochester, MN):** Sophie J. Bakri, MD (PI); Nakhleh Abu-Yaghi, MD (O); Andrew Barkmeier, MD (O); Karin Berg, COA (VA/R); Jean Burrington, COA (VA/R); Albert Edwards, MD (O); Shannon Goddard, COA (OP/OCT); Shannon Howard (VA/R); Raymond Iezzi, MD (O); Denise Lewison, COA (OP/OCT); Thomas Link, CRA (OP/OCT); Colin A. McCannel, MD (O); Joan Overend (VA/R); John Pach, MD (O); Margaret Ruszczyk, CCRP (CC); Ryan Shultz, MD (O); Cindy Stephan, COT (VA/R); Diane Vogen (CC).

**Dean A. McGee Eye Institute (Oklahoma City, OK):** Reagan H. Bradford Jr, MD (PI); Vanessa Bergman, COA, CCRC (CC); Russ Burris (OP/OCT); Amanda Butt, CRA (OP/OCT); Beth Daniels, COA (CC); Connie Dwiggin, CCRC (CC); Stephen Fransen, MD (O); Tiffany Guerrero (CC/DE); Darin Haivala, MD (O); Amy Harris (CC); Sonny Icks (CC/DE); Ronald Kingsley, MD (O); Lena Redden (VA/R); Rob Richmond (OP/OCT); Brittany Ross (VA/R); Kammerin White, CCRC (VA/R); Misty Youngberg, COA, CCRC (VA/R).

**Ophthalmic Consultants of Boston (Boston, MA):** Trexler M. Topping, MD (PI); Steve Bennett (OCT); Sandy Chong (VA/R); Mary Ciotti, COA (CC); Tina Cleary, MD (O); Emily Corey (VA/R); Dennis Donovan (OP/OCT); Albert Frederick, MD (O); Lesley Freese (CC/VA/R); Margaret Graham (OP/OCT); Natalya Gud, COA (VA/R); Taneika Howard (VA/R); Mike Jones (OP/OCT); Michael Morley, MD (O); Katie Moses (VA/R); Jen Stone (VA/R); Robin Ty, COA (VA/R); Torsten Wiegand, PHD, MD (O); Lindsey Williams (CC); Beth Winder (CC).

**Tennessee Retina, P.C. (Nashville, TN):** Carl C. Awh, MD (PI); Michelle Amonette (OCT); Everton Arrindell, MD (O); Dena Beck (OCT); Brandon Busbee, MD (O); Amy Dilback (OP/OCT); Sara Downs (VA/R); Allison Guidry, COA (VA/R); Gary Gutow, MD (O); Jackey Hardin (VA/R); Sarah Hines, COA (CC); Emily Hutchins (VA/R); Kim LaCivita, MA (OP/OCT); Ashley Lester (OP/OCT); Larry Malott (OP/OCT); MaryAnn McCain, RN, CNOR (CC); Jayme Miracle (VA/R); Kenneth Moffat, MD (O); Lacy Palazzotta (VA/R); Kelly Robinson, COA (VA/R); Peter Sonkin, MD (O); Alecia Travis (OP/OCT); Roy Trent Wallace, MD (O); Kelly J. Winters, COA (CC); Julia Wray (OP/OCT).

**Retina Associates Southwest, P.C. (Tucson, AZ):** April E. Harris, MD (PI); Mari Bunnell (OCT); Katrina Crooks (VA/R); Rebecca Fitzgerald, CCRC (CC/OCT); Cameron Javid, MD (O); Corin Kew (VA/R); Erica Kill, VAE (VA/R); Patricia Kline (VA/R); Janet Kreienkamp (VA/R); Maricruz Martinez (CC/OCT); Roy Ann Moore, OMA (CC/OCT); Egbert Saavedra, MD (O); LuAnne Taylor, CSC (CC/OCT); Mark Walsh, MD (O); Larry Wilson (OP).

**Midwest Eye Institute (Indianapolis, IN):** Thomas A. Ciulla, MD (PI); Ellen Coyle, COMT (VA/R); Tonya Harrington, COA (VA/R); Charlotte Harris, COA (VA/OCT); Cindi Hood (OCT); Ingrid Kerr, COA (VA/R); Raj Maturi, MD (O); Dawn Moore (OCT); Stephanie Morrow, COA (OP); Jennifer Savage, COA (VA); Bethany Sink, COA (CC/VA/R); Tom Steele, CRA (OP); Neelam Thukral, CCRC (CC/OCT); Janet Wilburn, COA (CC).

**National Ophthalmic Research Institute (Fort Myers, FL):** Joseph P. Walker, MD (PI); Jennifer Banks (VA/R); Debbie Ciampaglia (OP/OCT); Danielle Dyshanowitz (VA/R); Jennifer Frederick, CRC (CC); A. Tom Ghuman, MD (O); Richard Grodin, MD (O); Cheryl Kiesel, CCRC (CC); Eileen Knips,

RN, CCRC, CRA (OP/OCT); Jonathan McCue (VA/R); Maria Ortiz (VA/R); Crystal Peters, CCRC (CC); Paul Raskauskas, MD (O); Etienne Schoeman (OP/OCT); Ashish Sharma, MD (O); Glenn Wing, MD (O), Rebecca Youngblood (CC).

**University of Wisconsin Madison (Madison, WI):** Suresh R. Chandra, MD (PI); Michael Altaweel, MD (O); Barbara Blodi, MD (O); Kathryn Burke, BA (VA/R); Kristine A. Dietzman, (CC); Justin Gottlieb, MD (O); Gene Knutson (OP/OCT); Denise Krolnik (OP/OCT); T. Michael Nork, MD (O); Shelly Olson (VA/R); John Peterson, CRA (OP/OCT); Sandra Reed (OP/OCT); Barbara Soderling (VA/R); Guy Somers (VA/R); Thomas Stevens, MD (O); Angela Wealti, (CC).

**Duke University Eye Center (Durham, NC):** Srilaxmi Bearely, MD (PI); Brenda Branchaud (VA/R); Joyce W. Bryant, COT, CPT (CC/VA/R); Sara Crowell (CC/VA); Sharon Fekrat, MD (O); Merritt Gammage (OP/OCT); Cheala Harrison, COA (VA/R); Sarah Jones (VA); Noreen McClain, COT, CPT, CCRC (VA/R); Brooks McCuen, MD (O); Prithvi Mruthyunjaya, MD (O); Jeanne Queen, CPT (OP/OCT); Neeru Sarin, MBBS (VA/R); Cindy Skalak, RN, COT (VA/R); Marriner Skelly, CRA (OP/OCT); Ivan Suner, MD (O); Ronnie Tomany (OP/OCT); Lauren Welch (OP/OCT).

**University of California-Davis Medical Center (Sacramento, CA):** Susanna S. Park, MD, PHD (PI); Allison Cassidy (VA/R); Karishma Chandra (OP/OCT); Idalew Good (VA/R); Katrina Imson (CC); Sashi Kaur (OP/OCT); Helen Metzler, COA, CCRP (CC/VA/R); Lawrence Morse, MD, PHD (O); Ellen Redenbo, ROUB (OP/OCT); Marisa Salvador (VA/R); David Telander, MD (O); Mark Thomas, CRA (OCT); Cindy Wallace, COA (CC).

**University of Louisville School of Medicine, KY (Louisville, KY):** Charles C. Barr, MD (PI); Amanda Battcher (VA/R); Michelle Bottorff, COA (CC/OCT); Mary Chasteen (VA/R); Kelly Clark (VA/R); Diane Denning, COT (OCT); Debra Schoen (OP); Amy Schultz (OP); Evie Tempel, CRA, COA (OP); Lisa Wheeler, COT (VA/R); Greg K. Whittington, MPS, PSY (CC).

**Retina Associates of Kentucky (Lexington, KY):** Thomas W. Stone, MD (PI); Todd Blevins (OP/OCT); Michelle Buck, COT, (VA/R/OCT); Lynn Cruz, COT (CC); Wanda Heath (VA/R); Diana Holcomb (VA/R); Rick Isernhagen, MD (O); Terri Kidd, COA (OCT); John Kitchens, MD (O); Cathy Sears, CST, COA (VA/R); Ed Slade, CRA, COA (OP/OCT); Jeanne Van Arsdall, COA (VA/R); Brenda VanHoose, COA (VA/R); Jenny Wolfe, RN (CC); William Wood, MD (O).

**Colorado Retina Associates (Denver, CO):** John Zilis, MD (PI); Carol Crooks, COA (VA/R); Larry Disney (VA/R); Mimi Liu, MD (O); Stephen Petty, MD (O); Sandra Sall, ROUB, COA (CC/VA/R/OP/OCT).

**University of Iowa Hospitals & Clinics (Iowa City, IA):** James C. Folk, MD (PI); Tracy Aly, CRA (OP/OCT); Abby Brotherton (VA); Douglas Critser, CRA (OP/OCT); Connie J. Hinz, COT, CCRC (CC/VA/R); Stefani Karakas, CRA (OP/OCT); Valerie Kirschner (VA); Cheyanne Lester (VA/R); Cindy Montague, CRA (OP/OCT); Stephen Russell, MD (O); Heather Stockman (VA/R); Barbara Taylor, CCRC (VA/R); Randy Verdick, FOPS (OP/OCT), Jean Walshire (CC).

**Retina Specialists (Towson, MD):** John T. Thompson, MD (PI) ; Barbara Connell (VA/R); Maryanth Constantine (CC); John L. Davis Jr (VA/R); Gwen Holsapple (VA/R); Lisa Hunter (OP/OCT); C. Nicki Lenane (CC/VA/R/OP/OCT); Robin Mitchell (CC); Leslie Russel, CRA (OP/OCT); Raymond Sjaarda, MD (O).

**Retina Consultants of Houston (Houston, TX):** David M. Brown, MD (PI); Matthew Benz, MD (O); Llewellyn Burns (OCT); JoLene G. Carranza, COA, CCRC (CC); Richard Fish, MD (O); Debra Goates

(VA/R); Shayla Hay (VA/R); Theresa Jeffers, COT (VA/R); Eric Kegley, CRA, COA (OP/OCT); Dallas Kubecka (VA/R); Stacy McGilvra (VA/R); Beau Richter (OCT); Veronica Sneed, COA (VA/R); Cary Stoeber (OCT); Isabell Tellez (VA/R); Tien Wong, MD (O).

**Massachusetts Eye and Ear Infirmary/Harvard Vanguard Medical Associates (Boston, MA):**

Ivana Kim, MD (PI); Christopher Andreoli, MD (O); Leslie Barresi, CRA, COA, OCT-C (VA/OP/OCT); Sarah Brett (OP); Charlene Callahan (OP); Karen Capaccioli (OCT); William Carli, COA (VA/R/OCT); Matthew Coppola, COA (VA); Nicholas Emmanuel (CC); Claudia Evans, OD (VA/R); Anna Fagan, COA (VA/R); Marcia Grillo (OCT); John Head, CRA, OCT-C (OP/OCT); Troy Kieser, COA, OCT-C (CC/VA/R); Elaine Lee, COA (VA); Ursula Lord, OD (VA/R); Edward Miretsky (CC); Kate Palitsch (OP/OCT); Todd Petrin, RN (OCT); Liz Reader (CC); Svetlana Reznichenko, COA (VA); Mary Robertson, COA (VA); Justin Smith, OD (VA/R); Demetrios Vavvas, MD, PHD (O).

**Palmetto Retina Center (West Columbia, SC):** John Wells, MD (PI); Cassie Cahill (VA/R); W. Lloyd Clark, MD (O); Kayla Henry (VA/R); David Johnson, MD (O); Peggy Miller (CC/VA/R); LaDetrick Oliver, COT (OP/OCT); Robbin Spivey (OP/OCT); Tiffany Swinford (VA/R); Mallie Taylor (CC).

**Retina and Vitreous of Texas (Houston, TX):** Michael Lambert, MD (PI); Kris Chase (OP/OCT); Debbie Fredrickson, COA (VA/R); Joseph Khawly, MD, FACS (O); Valerie Lazarte (VA/R); Donald Lowd (OP/OCT); Pam Miller (CC); Arthur Willis, MD (O).

**Long Island Vitreoretinal Consultants (Great Neck, NY):** Philip J. Ferrone, MD (PI); Miguel Almonte (OCT); Rachel Arnott, (CC); Ingrid Aviles (VA/R/OCT); Sheri Carbon (VA/R); Michael Chitjian (OP/OCT); Kristen D'Amore (CC); Christin Elliott (VA/R); David Fastenberg, MD (O); Barry Golub, MD (O); Kenneth Graham, MD (O); AnnMarie Laverna (CC); Laura Murphy (VA/R); Amanda Palomo (VA/R); Christina Puglisi (VA/R); David Rhee, MD (O); Juan Romero, MD (O); Brett Rosenblatt, MD (O); Glenda Salcedo (OP/OCT); Marianne Schlameuss, RN (CC); Eric Shakin, MD (O); Vasanti Sookhai (VA/R).

**Wills Eye Institute/ Mid Atlantic Retina (Philadelphia, PA):** Richard Kaiser, MD (PI); Elizabeth Affel, MS, OCT-C (OCT); Gary Brown, MD (O); Christina Centinaro (CC); Deborah Fine, COA (OCT); Mitchell Fineman, MD (O); Michele Formoso (CC); Sunir Garg, MD (O); Lisa Grande (VA/R); Carolyn Herbert (VA/R); Allen Ho, MD (O); Jason Hsu, MD (O); Maryann Jay (OCT); Lisa Lavetsky (OCT); Elaine Liebenbaum (OP); Joseph Maguire, MD (O); Julia Monsonago (OP/OCT); Lucia O'Connor (OCT); Lisa Pierce (CC); Carl Regillo, MD (O); Maria Rosario (DE); Marc Spirn, MD (O); James Vander, MD (O); Jennifer Walsh (VA/R).

**Ohio State University Eye Physicians & Surgeons-Retina Division (Dublin, OH):** Frederick H. Davidorf, MD (PI); Amanda Barnett (OP/OCT); Susie Chang, MD (O); John Christoforidis, MD (O); Joy Elliott (CC); Heather Justice (VA/R); Alan Letson, MD (O); Kathrynne McKinney, COMT (CC); Jeri Perry, COT (VA/R); Jill A. Salerno, COA (CC); Scott Savage (OP); Stephen Shelley (OCT).

**Retina Associates of Cleveland (Beachwood, OH):** Lawrence J. Singerman, MD (PI); Joseph Coney, MD (O); John DuBois (OP/OCT); Kimberly DuBois, LPN, CCRP, COA (VA/R); Gregg Greanoff, CRA (OP/OCT); Dianne Himmelman, RN, CCRC (CC); Mary Ilc, COT (VA/R); Elizabeth Mcnamara (VA/R/OP); Michael Novak, MD (O); Scott Pendergast, MD (O); Susan Rath, PA-C (CC); Sheila Smith-Brewer, CRA (OP/OCT); Vivian Tanner, COT, CCRP (VA/R); Diane E. Weiss, RN, (CC); Hernando Zegarra, MD (O).

**Retina Group of Florida (Fort Lauderdale, FL):** Lawrence Halperin, MD (PI); Patricia Aramayo (OCT); Mandeep Dhalla, MD (O); Brian Fernandez, MD (OP/OCT); Cindy Fernandez, MD (CC); Jaclyn

Lopez (CC); Monica Lopez (OCT); Jamie Mariano, COA (VA/R); Kellie Murphy, COA (OCT); Clifford Sherley, COA (VA/R); Rita Veksler, COA (OP/OCT).

**Retina-Vitreous Associates Medical Group (Beverly Hills, CA):** Firas Rahhal, MD (PI); Razmig Babikian (DE); David Boyer, MD (O); Sepideh Hami (DE); Jeff Kessinger (OP/OCT); Janet Kurokouchi (CC); Saba Mukarram (VA/R); Sarah Pachman (VA/R); Eric Protacio (OCT); Julio Sierra (VA/R); Homayoun Tabandeh, MD, MS, FRCP (O); Adam Zamboni (VA/R).

**Elman Retina Group, P.A. (Baltimore, MD):** Michael Elman, MD (PI); Jennifer Belz (CC); Tammy Butcher (CC); Theresa Cain (OP/OCT); Teresa Coffey, COA (VA/R); Dena Firestone (VA/R); Nancy Gore (VA/R); Pamela Singletary (VA/R); Peter Sotirakos (OP/OCT); JoAnn Starr (CC).

**University of North Carolina at Chapel Hill (Chapel Hill, NC):** Travis A. Meredith, MD (PI); Cassandra J. Barnhart, MPH (CC/VA/R); Debra Cantrell, COA (VA/R/OP/OCT); RonaLyn Esquejo-Leon (OP/OCT); Odette Houghton, MD (O); Harpreet Kaur (VA/R); Fatoumatta NDure, COA (CC).

**Ophthalmologists Enrolling Patients but No Longer Affiliated with a CATT Center:** Ronald Glatzer, MD (O); Leonard Joffe, MD (O); Reid Schindler, MD (O).

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**Coordinating Center (University of Pennsylvania, Philadelphia, PA):** Maureen G. Maguire, PhD (PI); Mary Brightwell-Arnold, SCP (Systems Analyst); Ruchira Glaser, MD (Medical Monitor); Judith Hall (Protocol Monitor); Sandra Harkins (Staff Assistant); Jiayan Huang, MS (Biostatistician); Alexander Khvatov, MS (Systems Analyst); Kathy McWilliams, CCRP (Protocol Monitor); Susan K. Nolte (Protocol Monitor); Ellen Peskin, MA, CCRP (Project Director); Maxwell Pistilli, MS, MEd (Biostatistician); Susan Ryan (Financial Administrator); Allison Schnader (Administrative Coordinator); Gui-Shuang Ying, PhD (Senior Biostatistician).

**OCT Reading Center (Duke University, Durham, NC):** Glenn Jaffe, MD (PI); Jennifer Afrani-Sakyi (CATT PowerPoint Presentations); Brannon Balsley (OCT Technician Certifications); Linda S. Bennett (Project Manager); Adam Brooks (Reader/SD-Reader); Adrienne Brower-Lingsch (Reader); Lori Bruce (Data Verification); Russell Burns (Senior Technical Analyst/Senior Reader/SD Reader/OCT Technician Certifications); Dee Busian (Reader); John Choong (Reader); Lindsey Cloaninger (Reader Reliability Studies/ Document Creation/CATT PPT Files); Francis Char DeCroos (Research Associate); Emily DuBois (Data Entry); Mays El-Dairi (Reader/SD-Reader); Sarah Gach (Reader); Katelyn Hall (Project Manager/Reader Reliability Studies/ Data Verification/Document Creation); Terry Hawks (Reader); ChengChenh Huang (Reader); Cindy Heydary (Senior Reader/Quality Assurance Coordinator/SD Reader/Data Verification); Alexander Ho (Reader, Transcription); Shashi Kini (Data Entry/Transcription); Michelle McCall (Data Verification); Daaimah Muhammad (Reader Feedback); Jayne Nicholson (Data Verification); Jeanne Queen (Reader/SD-Reader); Pamela Rieves (Transcription); Kelly Shields (Senior Reader); Cindy Skalac (Reader); Adam Specker (Reader); Sandra Stinnett (Biostatistician); Sujatha Subramaniam (Reader); Patrick Tenbrink (Reader); Cynthia Toth, MD (Director of Grading); Aaron Towe (Reader); Kimberly Welch (Data Verification); Natasha Williams (Data Verification); Katrina Winter (Senior Reader); Ellen Young (Senior Project Manager).

**Fundus Photograph Reading Center (University of Pennsylvania, Philadelphia, PA):** Juan E. Grunwald, MD (PI); Judith Alexander (Director); Ebenezer Daniel, MBBS, MS, MPH, PhD (Director); Elisabeth Flannagan (Administrative Coordinator); E. Revell Martin (Reader); Candace Parker (Reader); Krista Sepielli (Reader); Tom Shannon (Systems Analyst); Claressa Whearry (Data Coordinator).

**National Eye Institute, National Institutes of Health:** Maryann Redford, DDS, MPH (Program Officer).

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**Clinic Monitoring Committee:** Ellen Peskin, MA, CCRP (chair); Mary Brightwell-Arnold, SCP; Joan DuPont; Maureen G. Maguire, PhD; Kathy McWilliams, CCRP; Susan K. Nolte.

**Data and Safety Monitoring Committee:** Lawrence M. Friedman, MD (chair); Susan B. Bressler, MD; David L. DeMets, PhD; Martin Friedlander, MD, PhD; Mark W. Johnson, MD; Anne Lindblad, PhD; Douglas W. Losordo, MD, FACC; Franklin G. Miller, PhD.

### Credit Roster for IVAN

Site	Investigators	Research team
<b>Aintree University Hospitals:</b>	Mr Ahmed Kamal Thomas Papathomas Ahmed Khalil	Pauline Guinness Richard Hancock Maria Dangler-Harles Jane Blocksage Samantha Lorensen
<b>Aston University Birmingham<sup>1</sup>, Birmingham Optegra Eye Hospital<sup>2</sup>:</b>	Mr J.M.Gibson <sup>1</sup> Katie Pedwell <sup>1</sup> Jane Pitt <sup>2</sup>	Ajith Kumar S. Al-Husainy S.Sreekantam M.Hanratty Tara Clark
<b>Queen's University Belfast<sup>3</sup>, Belfast Health and Social Care Trust<sup>4</sup>:</b>	Prof Usha Chakravarthy <sup>3</sup> Dr Lisa Kelly <sup>4</sup> Dr Karen Gilvray <sup>4</sup>	Ms Pamela Jamison Mrs Georgina Sterret Mr Vittorio Silvestri

Dr Deirdre Burns  
Ms Rebecca Denham  
Mrs Joanne Grattan  
Mrs Teresa Rice  
Mrs Lenore Ponisi

**Aston University Birmingham<sup>5</sup>, Birmingham & Midland Eye Centre<sup>6</sup>:**

Mr J.M.Gibson<sup>5</sup>  
Katherine Brown<sup>6</sup>  
Bethan Swain<sup>6</sup>

A.Kumar  
G.Bliss  
P.Cikatricis  
K.Damer

**Royal Blackburn Hospital:**

Mrs Salwa Abugreen  
Mohamed Alarbie  
Debra Myerscough

H Patel  
N Nixon  
M Anderson  
T Thompson  
P Richardson

**Bradford Teaching Hospitals:**

Mr Faruque Ghanchi  
Mrs Helen Devonport  
Miss Nicci Atkinson

Julie Dixon  
Tony Dook  
Cara Phillips  
Tomas Cudrnak  
Charlotte Hazel  
Mary Elliott

**Sussex Eye Hospital:**

Mr Anthony Casswell  
Dr Gek Ong  
Mr Edward Hughes

Mrs Susan Bennett  
Mr Nick White  
Mrs Catriona Gardiner  
Mr Stephen Turner  
Mrs Tenesa Sargent

**Bristol Eye Hospital:**

Miss Claire Bailey

**Cambridge University Hospitals NHS Foundation Trust:**

Mr Douglas Newman  
Mr Kevin McNally

Haris Papanikolaou  
Dawn Russell-Hermanns  
Katherine Martin  
Jo Fielden

**Frimley Park Hospital NHS Foundation Trust:**

Mrs Geeta Menon  
Mrs Manju Chandran  
Mr Gulrez Ansari

Mrs Bhavani Mathapati  
Mr Nitin Jain  
Mrs Lorraine North  
Mrs Jincy Jose  
Nadeem Rob

**The Queen Elizabeth Hospital, King's Lynn NHS Foundation Trust:**

Mr R J Pushapanathan  
Mr I Ali

**Department of Eye and Vision Science, Institute of Ageing and Chronic Disease, University of Liverpool<sup>7</sup> and St. Paul's Eye Unit, Royal Liverpool University Hospitals NHS Trust<sup>8</sup>:**

Prof Simon P Harding<sup>7</sup>  
Sandra Taylor<sup>8</sup>  
Valerie Tompkin<sup>8</sup>

Karen Hawkins  
Jerry Sharp  
Stephen Pearson

Martin Hodgson  
William Hooley  
Gillian Lewis

**Maidstone Hospital:**

Mr Frank Ahfat  
Mr. Luke Membrey

Mrs. Margaret Gurney  
Mr. Clive Wood  
Dr. Shabeeba Hannan  
Mr. Syed Idris Haider  
Mr. Paul Adley

**Central Manchester University Hospitals NHS Foundation Trust, Manchester Academic Health Sciences Centre & School of Biomedicine, University of Manchester:**

Mr Paul N. Bishop

Ekaterina Varimezova-Georgieva

Tariq M Aslam

**Newcastle upon Tyne Hospital Foundation Trust:**

Mr S James Talks  
Rajeev Kak

Alan Branon  
Kapka Nenova  
Kevin Gales

**Nottingham University Hospitals Trust:**

Mr Alexander Foss

Karen Armstrong-Owen

**Oxford Eye Hospital, Oxford University Hospitals:**

Miss Susan Downes  
Miss Shahrnaz Izadi  
Mr Robert Purbrick

Mrs Alexina Fantato  
Mrs Ivy Samuel  
Miss Vicky Hart  
Mrs Anna Rudenko  
Mr Lewis Smith  
Mr Charles Cottrill  
Miss Paula Hedges

**Sheffield Teaching Hospitals NHS Foundation Trust<sup>9</sup>, South Warwickshire NHS Foundation Trust<sup>10</sup>:**

Mr Christopher Brand<sup>9</sup>  
Dr Hibba Abdulkarim<sup>9</sup>  
Mrs Uma Thakur<sup>10</sup>

Mrs Helen Pokora  
Mr Andy Jubb  
Mrs Katy Kelly  
Mr Fahd Quhill  
Mrs Mary Freeman

**Southend Hospital:**

Mr Niral Karia  
Mr A Krishnan

Ms Maria Shipman  
Mr John Williams  
Mr Chris Johnson

**Faculty of Medicine, University of Southampton<sup>11</sup>, Wellcome Trust Clinical Research Facility, University Hospital Southampton NHS Foundation Trust<sup>12</sup>, Southampton Eye Unit, University Hospital Southampton NHS Foundation Trust<sup>13</sup>:**

Prof Andrew John Lotery<sup>11</sup>  
Marie Anne Nelson<sup>12</sup>  
Suresh Thulasidharan<sup>13</sup>

Claire O'Brien  
Kevin Oxlade  
Caitrin Watkins  
Maria Gemenetzi  
Gabiella De Salvo

**Sunderland Eye infirmary, Sunderland, Institute of Genetic Medicine, Newcastle University<sup>14</sup>, Sunderland Eye infirmary, Sunderland<sup>15</sup>:**

Mr David Steel<sup>14</sup>

Steve Dodds



Eoghan Millar<sup>14</sup>  
Vinna Manjunath<sup>15</sup>

Shelagh Thomson  
Martyn Hallowell  
Hugh Harris  
Paula Foley

**Torbay Hospital:**

Mr Mick Cole  
Yinka Osoba  
Sanjay Dhir

Annette Field  
Sharon Criddle  
Karin Tilley  
Eddy Doyle  
Debbie Knowles

**Royal Wolverhampton Hospitals NHS Trust:**

Mr Yit C Yang  
Nirodhini Narendran  
Swathi Paneerselvam

Jas Purewal  
Mary Stott  
Bhagal Bhagal  
Sharon Hughes  
Gurminder Sahota  
Jenny Nosek

**Credit Roster for LUCAS**

Karina Berg MD, Department of Ophthalmology, Oslo University Hospital, Oslo, Norway

Ragnheiður Bragadóttir MD PhD, Department of Ophthalmology, Oslo University Hospital, Oslo, Norway

Terje R. Pedersen MD PhD, Department of Endocrinology, Morbid Obesity and Preventive Medicine, Oslo University Hospital, Oslo, Norway

Leiv Sandvik PhD, Department of Biostatistics and Epidemiology, Oslo University Hospital, Oslo, Norway

Emina Hadzalic MD, Department of Ophthalmology, Betanien Hospital, Skien, Norway

Inger Gjertsen MD, Department of Ophthalmology, Vestre Viken Hospital Trust, Drammen, Norway

Vegard Forsaa MD, Department of Ophthalmology, Stavanger University Hospital, Stavanger, Norway

Lars Haakon Berger MD, Department of Ophthalmology, Østfold Hospital, Moss, Norway

Bettina Kinge MD PhD, The Retina Clinic, Aleris, Oslo, Norway

Hans Henschien MD, Department of Ophthalmology, Vestfold Hospital, Tønsberg, Norway

Kristian Fossen MD, Department of Ophthalmology, University hospital of Northern Norway, Tromsø, Norway

Slavica Markovic MD, Department of Ophthalmology, Innlandet Hospital, Elverum, Norway