

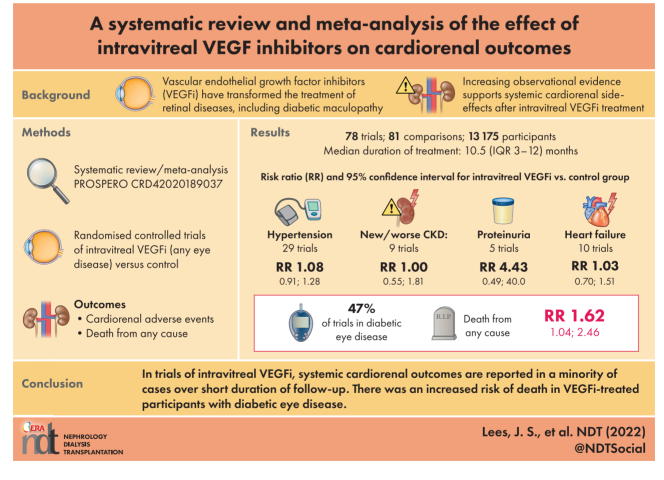
A systematic review and meta-analysis of the effect of intravitreal VEGF inhibitors on cardiorenal outcomes

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GRAPHICAL ABSTRACT



ABSTRACT

Background. Vascular endothelial growth factor inhibitors (VEGFis) have transformed the treatment of many retinal diseases, including diabetic maculopathy. Increasing evidence supports systemic absorption of intravitreal VEGFi and development of significant cardiorenal side effects.

Methods. We conducted a systematic review and metaanalysis (PROSPERO: CRD42020189037) of randomised controlled trials of intravitreal VEGFi treatments (bevacizumab, ranibizumab and aflibercept) for any eye disease. Outcomes of interest were cardiorenal side effects (hypertension, protein-

opment of significant cardiorenal side effects. "The Author(s) 2022. Published by Oxford University Press on behalf of the ERA. This is an Open Access article distributed under the terms of the Creative Commons Attribution-NonCommercial License (https://creativecommons.org/licenses/by-nc/4.0/), which permits non-commercial re-use, distribution, and reproduction in any medium, provided the original work is properly cited. For commercial re-use, please contact journals.permissions@oup.com

What is already known about this subject?

- Intravitreal vascular endothelial growth factor inhibitors (VEGFis) are commonly used in the treatment of diabetic eye disease.
- There is increasing evidence of systemic absorption of intravitreal VEGFi associated with exacerbation of cardiorenal side effects such as hypertension, proteinuria, decline in kidney function and heart failure.

What this study adds?

- Trials of intravitreal VEGFi do not routinely report cardiorenal side effects, although mechanistically these side effects are plausible, especially in patients with diabetes and pre-existing kidney disease.
- In patients with diabetic eye disease (who commonly also have kidney disease), treatment with intravitreal VEGFi is associated with an increased risk of death, with potential implications for obtaining informed consent.

What impact this may have on practice or policy?

- Additional scrutiny of post-licensing observational data may improve recognition of safety concerns in VEGFi-treated patients.
- Monitoring for cardiorenal side effects should be considered, especially in high-risk patients with diabetes and kidney disease who are treated with intravitreal VEGFi.

uria, kidney function decline and heart failure). Fixed effects meta-analyses were conducted where possible.

Results. There were 78 trials (81 comparisons; 13175 participants) that met the criteria for inclusion: 47% were trials in diabetic eye disease. Hypertension (29 trials; 8570 participants) was equally common in VEGFi and control groups {7.3 versus 5.4%; relative risk [RR] 1.08 [95% confidence interval (CI) 0.91–1.28]}. New or worsening heart failure (10 trials; 3384 participants) had a similar incidence in VEGFi and control groups [RR 1.03 (95% CI 0.70–1.51)]. Proteinuria (5 trials; 1902 participants) was detectable in some VEGFi-treated participants (0.2%) but not controls [0.0%; RR 4.43 (95% CI 0.49–40.0)]. Kidney function decline (9 trials; 3471 participants) was similar in VEGFi and control groups. In participants with diabetic eye disease, the risk of all-cause mortality was higher in VEGFi-treated participants [RR 1.62 (95% CI 1.04–2.46)].

Conclusion. In trials of intravitreal VEGFi, we did not identify an increased risk of cardiorenal outcomes, although these outcomes were reported in only a minority of cases. There was an increased risk of death in VEGFi-treated participants with diabetic eye disease. Additional scrutiny of post-licensing observational data may improve the recognition of safety concerns in VEGFi-treated patients.

Keywords: CKD, diabetes mellitus, hypertension, proteinuria, systematic review

INTRODUCTION

Vascular endothelial growth factor inhibitors (VEGFis) have transformed the treatment of many retinal diseases [1] but are most commonly used in the management of diabetic macular oedema (DME) [2], neovascular age-related macular degeneration (nAMD) [3] and retinal vein occlusion [4]. The VEGFi ranibizumab (Lucentis, Novartis UK, London, UK) and aflibercept (Eylea, Regeneron Pharmaceuticals, Tarrytown, NY, USA) were specifically designed for intravitreal treatment. Bevacizumab (Avastin, Genentech, South San Francisco, CA, USA) was originally developed for systemic administration in the treatment of various cancers but is used extensively offlabel across the globe (less so in the UK) for retinal disease because of the significant cost savings associated with using a more established treatment. Brolucizumab (Beovu, Novartis UK) is also licenced by the US Food and Drug Administration (2019) and the European Medicines Agency (2020) after landmark studies showed efficacy for the treatment of nAMD and DME. Due to its more recent marketing authorisation, brolucizumab is not considered further in this review.

A treatment course of intravitreal VEGFi typically consists of a loading phase of monthly injections over 2–4 months followed by an extension phase based on the treatment response. These treatments are commonly used in patients with diabetes: \approx 50% of patients with type 1 diabetes and >25% of patients with type 2 diabetes have evidence of diabetic retinopathy [5]. Although only 2–4% of patients with diabetes require ophthalmic treatment, the absolute number of people with any diabetic eye disease is forecast to increase in Europe from 6.4 million to 8.6 million in 2050, which will substantially increase the number of patients eligible for intravitreal VEGFi treatment [6].

In the case of bevacizumab, intravitreal VEGFi is administered at <15% of the intravenous dose and was previously thought to exert predominantly local effects within the eye [7]; however, increasing evidence supports pronounced systemic absorption [8]. In 56 patients with age-related macular degeneration, intravitreal administrations of ranibizumab, aflibercept and bevacizumab were all rapidly detectable in the circulation [9]. Ranibizumab (48-kDa monoclonal antibody fragment) was cleared relatively quickly (within days), but aflibercept (115-kDa fusion protein) and bevacizumab (149-kDa full-length monoclonal antibody) accumulated over repeated doses and suppressed free plasma VEGF [9] for at least 7 days [9] and up to 30 days after intravitreal injection [10]. In 82 patients with nAMD, intravitreal administration of bevacizumab was associated with both de novo blood pressure (BP) dysregulation and exacerbation of pre-existing hypertension [11], although this has not been a consistent finding in all studies nor for all intravitreal VEGFi treatments [12]. Systemic VEGFi, when used primarily as an anticancer therapy, is almost universally associated with the development or exacerbation of hypertension [13]. Hypertension and endothelial damage associated with VEGFi are associated with end-organ damage, including heart failure [14], nephropathy and kidney failure [13]. Caution is advised in administering systemic VEGFi in patients with pre-existing hypertension, proteinuria, cardiovascular disease and severe kidney impairment, and monitoring for hypertension, proteinuria and heart failure is recommended [15]. No such monitoring recommendations exist for intravitreal VEGFi administration.

Particularly in patients with diabetes—already at a higher risk of end-organ damage—the potential for systemic absorption of intraocular therapies and accelerated albuminuria, heart failure and progression to end-stage kidney disease is a major concern. The cardiorenal side effects of intravitreal VEGFis may be identifiable from initial trials: any cardiorenal safety signal seen in these groups should highlight the need for greater vigilance in patients receiving VEGFi, either systemically for cancer or locally for ophthalmological indications.

The aims of this review were to identify the prevalence of cardiorenal side effects after intravitreal administration of VEGFi and to identify factors associated with cardiorenal side effects after intravitreal VEGFi administration.

MATERIALS AND METHODS

This review was prospectively registered on PROSPERO (CRD42020189037) and reported according to the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) methodology. PICOS (Population, Intervention, Comparison, Outcome, Setting) criteria for inclusion are detailed in Table 1.

Types of studies

Randomised controlled trials that report outcomes in those receiving intravitreal VEGFi treatments versus control groups [no treatment/sham or non-VEGFi treatment (e.g. laser photocoagulation)] were eligible for inclusion. We included trials that administered a minimum of two intravitreal injections of VEGFi with follow-up for at least 4 weeks with a minimum of 20 participants.

Types of participants

Participants receiving VEGFi treatment for any eye disease, with any baseline level of eye disease, cardiovascular disease and kidney function, were eligible for inclusion. The following was recorded for all eligible studies: VEGFi type, number of injections (intended or mean administered dose), duration of VEGFi treatment and duration of follow-up. We intended to record baseline demographics for the included populations: sex, age, ethnicity, kidney function [estimated glomerular filtration rate (eGFR)], BP, urinary protein content (spot urinary protein:creatinine ratio or albumin:creatinine

Table 1: PICOS criteria for study inclusion.

Population	Any maculopathy
	Any baseline level of eye disease, cardiovascular disease
	and kidney function
Intervention	Intravitreal VEGF inhibitor, with a minimum of two
	injections:
	Bevacizumab
	Ranibizumab
	Aflibercept
Comparison	Control group
	No treatment
	Sham treatment
	Non-VEGFi treatment, e.g. laser photocoagulation
Outcomes	Cardiorenal outcomes:
	Hypertension
	Proteinuria
	New or worsening heart failure
	Heart failure hospitalisation
	New CKD (de novo reduction in eGFR to
	<60 ml/min/1.73 m ²)
	Decrease in eGFR by \geq 30%
	Doubling serum creatinine
	Need for dialysis or transplant
	eGFR slope
	Arterial thrombotic cardiovascular events: MI, stroke
	peripheral arterial disease
	Venous thromboembolism: pulmonary embolus,
	deep vein thrombosis
	Death from cardiovascular cause (MI, stroke, HF)
	Death from kidney failure
	All-cause mortality
Setting	Randomised controlled trials

ratio or 24-h urine protein:albumin ratio), hypertension (or prescription of antihypertensive medications), diabetes (%) and cardiovascular disease, including coronary artery disease, myocardial infarction (MI), stroke, peripheral arterial disease (%) and heart failure (%) and severity [ejection fraction (EF)] if available.

Types of interventions

We examined the following VEGFis delivered as intravitreal injections with a minimum of two injections, assuming there were follow-up data available at least 4 weeks after the interventions started: bevacizumab, ranibizumab and aflibercept.

Outcome measures

We extracted data on the following outcome measures, where available: hypertension, proteinuria, new or worsening heart failure, heart failure hospitalisation, new CKD (defined as *de novo* reduction in eGFR to <60 ml/min/1.73 m²), a decrease in eGFR by \geq 30%, doubling of serum creatinine, need for dialysis or transplant, eGFR slope, arterial thrombotic cardiovascular events (MI, stroke, peripheral arterial disease), venous thromboembolism (pulmonary embolus, deep vein thrombosis), all-cause mortality, death from a cardiovascular cause (MI, stroke, heart failure) and death from kidney failure.

Search strategy

The search period spanned 1966 to the end of May 2020. We searched PubMed, Cochrane Library (CENTRAL), Google (for grey literature) and the ISRCTN registry (for ongoing studies) for relevant studies. Hand searching was performed, including references of included articles and references from previous reviews of intravitreal VEGFi therapy. No language restriction was applied to eligible reports (although there were no identified reports that were not published in English). Abstracts were eligible for inclusion if relevant data were available (although no studies were included in abstract-only form).

Search terms

We used the following search terms to identify eligible reports: (vascular endothelial growth factor OR VEGF OR bevacizumab OR ranibizumab OR aflibercept) AND (intravitreal OR intraocular) AND (clinical trial OR randomized controlled trial).

Review methods

All possible randomised controlled trials were identified independently by two researchers (J.S.L. and S.J.H.D.) and entered into Mendeley Reference Manager software. Two researchers independently assessed titles and abstracts of all possible relevant studies. When eligibility was not clear from the title and/or abstract, the full article was reviewed. Differences were resolved by discussion between the two researchers. Data were abstracted by two researchers (J.S.L. and S.J.H.D.) using a pre-specified form. Where available, the trial registration identifier (from clinical trial registries) was extracted to identify repeat publications for each trial. The first relevant trial publication was included, as this tended to contain the most complete baseline demographic information. We performed an exploratory search using the trial identifier from each included trial to perform a targeted search for later trial publications with the maximum published duration of follow-up that also reported cardiorenal outcomes.

Statistical analysis

We conducted frequentist meta-analysis to report risk ratios (RRs) and 95% confidence intervals (CIs) using fixed effects models, stratified for VEGFi type, where adequate data were available for outcomes of interest. Weights were assigned by the inverse variance method. Statistical heterogeneity was assessed using I^2 (\geq 50% was considered to represent significant heterogeneity) and τ^2 (as an estimate of between-study heterogeneity). Meta-regression models were used to assess potential sources of heterogeneity, including VEGFi type, number of injections, duration of treatment and treated eye disease (diabetic versus non-diabetic indication). Variables accounting for heterogeneity among studies were identified if their inclusion in the model resulted in a significant reduction in τ^2 . Meta-regression identified treated eye disease (diabetic

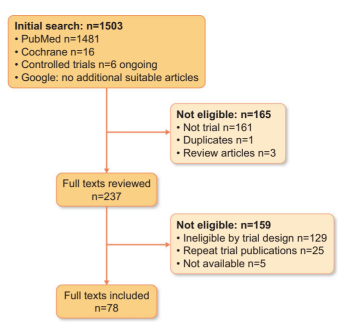


Figure 1: Flow chart of included studies and reasons for exclusion.

versus non-diabetic) as a significant source of heterogeneity for death outcome: forest plots were generated for all cardiorenal outcomes and were stratified by diabetic versus non-diabetic eye disease. Evidence of publication bias was sought by visual inspection of funnel plots and trim-and-fill analysis.

RESULTS

Trial characteristics

Of 1503 articles identified on systematic searching, there were 78 eligible full texts containing 81 comparisons in 13 175 participants (Fig. 1). There were 31 comparisons of intravitreal bevacizumab {mean 4.0 injections [standard deviation (SD) 2.4]}, 44 comparisons of intravitreal ranibizumab [mean 8.3 injections (SD 6.3)] and 6 comparisons of intravitreal aflibercept [mean 7.2 injections (SD 1.6)]. The median duration of treatment was 10.5 months [interquartile range (IQR) 3–12] and the median duration of follow-up was 12 months (IQR 6–12) (Table 2).

There were 37 trials performed for diabetic indications (27 diabetic macular oedema and 10 proliferative diabetic nephropathy), 11 trials for nAMD, 23 for retinal vein occlusion and 10 for other indications (most commonly for neovascular glaucoma or polypoidal choroidal vasculopathy; Table 2). In the 44 trials conducted for non-diabetic indications, the presence or absence of diabetes at baseline was recorded in 7 trials (15.9%).

Across all trials, there was a male preponderance of 55.3% and the mean age was 63 years (SD 8). Ethnicity was recorded in 58.0% of studies; of these, 80.9% of trials had a Caucasian majority.

Baseline comorbidities

Baseline BP or a history of hypertension was recorded in nine trials (11.0%; 1690 participants), unspecified

			- - -	Men	Age (years),	ıent,	Controls,		Injections	Duration of treatment	Duration of follow-up
n		Country	Lye disease	(%)	mean	и	и	VЕGHI type	(n), mean	(months)	(months)
	SN	A	nAMD	40.2	78.4	121	63	Ranibizumab	10	24	24
ACTRN12607000577415 Brazil	Braz		Neovascular glaucoma	60.0	60.8	20	20	Bevacizumab	ŝ	2	24
NCT01325181 Korea	Korea		Chronic central serous chorioretinonathy	82.4	50.8	16	18	Ranibizumab	б	7	12
NCT01909791 USA	USA		Diabetic macular oedema	62.3	59	226	476	Aflibercept	8.3	24	24
	Italy		Branch retinal vein occlusion	58.3	67	153	154	Ranibizumab	8	12	12
NR Lebanon	Lebano	u	nAMD	50.0	75	32	30	Bevacizumab	2.4	9	9
NCT01489189 USA	USA		Proliferative diabetic retinopathy	56.0	53.5	102	114	Ranibizumab	10	24	24
NCT01135914 Canada	Canada	~	Diabetic macular oedema	60.0	61.7	148	72	Ranibizumab	6	12	12
NCT00996437 USA	USA		Proliferative diabetic retinopathy	48.0	58	125	136	Ranibizumab	б	2	4
NR India	India		Diabetic macular oedema	40.0	57.7	12	18	Bevacizumab	2.6	6	9
NCT00943072 International	Internat	tional	Central retinal vein occlusion	57.0	66.3	114	74	Aflibercept	9	9	12
NCT00061594 USA	NSA		nAMD	48.7	77	280	143	Ranibizumab	11	12	12
NCT00485836 USA	USA		Central retinal vein occlusion	57.0	68	132	130	Ranibizumab	5.7	9	9
NCT00485836 USA	USA		Central retinal vein occlusion	57.0	68	130	130	Ranibizumab	5.7	9	9
NCT00486018 USA	NSA		Branch retinal vein occlusion			132	131	Ranibizumab	5.7	9	12
NR International	Internatio	nal	nAMD	44.8	77.7	140	143	Ranibizumab	22	24	24
	USA		Diabetic macular oedema	62.8	63.6	182	181	Ranibizumab	8.7	12	12
NCT02050828 USA	USA		Diabetic macular oedema	59.0	61.3	48	46	Ranibizumab	Э	3	3
ND I opened	I abanon		Ductificanting dishatic rationarthy	52.4	696	12	<i>cy</i>	Donihizmuch	7	~	~
F01223612	теранон		r ionrerative urabeue reuniopaury Diabetic macular oedema	53.6 63.6	00.0 57 7	1/	70	Ranihizumah	/.c	+ <u>-</u> 1	4 12
	China		Central retinal vein occlusion	54.8	54.6	16	16	Bevacizumab	2.4	6	6
NCT00445003 USA	USA		Diabetic macular oedema	56.0	63	375	479	Ranibizumab	6	12	24
NCT00906685 Sweden	Sweden		Central retinal vein occlusion	60.0	70.5	30	30	Bevacizumab	4	6	9
NCT01746563 Brazil	Brazil		Proliferative diabetic retinopathy	46.4	52.6	14	15	Ranibizumab	2	1	9
NCT01280929 Portugal	Portugal		Proliferative diabetic retinopathy	74.3	57	22	13	Ranibizumab	5	12	12
NCT01941329 International	Internati	onal	Proliferative diabetic retinopathy	63.0	55.2	41	46	Bevacizumab	ю	ю	12
OzuBevaCRVOME-1 Egypt	Egypt		Central retinal vein occlusion	66.6	68.8	30	30	Bevacizumab	4.3	9	9
NCT01994291 International	Internatio	onal	Diabetic macular oedema	62.1	62.3	66	66	Ranibizumab	ю	б	4
NCT01489189 USA	USA		Proliferative diabetic retinopathy	56.0	52	191	203	Ranibizumab	19.2	60	60
-	German	y	Branch retinal vein occlusion			126	118	Ranibizumab	4.71	9	9
	Switzer	land	nAMD	32.5	78.5	19	21	Ranibizumab	3	1	12
NCT00056823 USA	USA		nAMD	46.9	74.1	106	56	Ranibizumab	24	24	24
UMIN000001546 Japan	Japan		Branch retinal vein occlusion	39.5	68.4	22	21	Bevacizumab	2.2	12	12

1able 2: Continued.												
Author	Vear	C leith	Country	Etre di cease	Men (%)	Age (years), '	Treatment,	Controls,	VF GFi trune	Injections	Duration of treatment	Duration of follow-up (monthe)
10 mm t	TCUT		f mmnoo	the machine	(0/)	IIICAIL		=	ATATINA		(empront)	
Hoerauf et al. [71]	2016	NCT01396083	Germany	Central retinal vein occlusion	59.7	66.1	124	119	Ranibizumab	4.52	9	9
Ishibashi <i>et al.</i> [72]	2015	NCT00989989	International	Diabetic macular oedema	54.0	61.4	132	131	Ranibizumab	7	12	12
Karadzic et al. [73]	2015	NR	Serbia	Branch retinal vein occlusion	60.0	60.5	11	6	Bevacizumab	2	3	б
Kinge et al. [74]	2010	NCT00567697	Norway	Central retinal vein occlusion	55.1	72	15	14	Ranibizumab	4.3	9	9
Koh <i>et al.</i> [75]	2012	NCT00674323	International	Polypoidal choroidal	65.0	63	19	21	Ranibizumab	3.9	9	9
Korobelnik <i>et al.</i> [76]	2014	NCT01331681	International	vascunopaury Diabetic macular oedema	62.2	64.1	135	132	Aflibercept	8	12	12
Korobelnik et al. [76]	2014	NCT01363440	USA	Diabetic macular oedema	53.4	62.4	151	154	Aflibercept	8	12	12
Kriechbaum [77]	2014	NR	Austria	Diabetic macular oedema	40.0	59	15	15	Bevacizumab	6	12	12
Kumar et al. [78]	2019	NR	India	Branch retinal vein occlusion	51.7	56.2	15	45	Ranibizumab	б	3	9
Lai <i>et al.</i> [79]	2018	NCT03459144	China	Polypoidal choroidal	63.1	62	34	23	Ranibizumab	б	12	12
				vasculopathy								
Lang <i>et al.</i> [80]	2018	NCT01131585	Germany	Diabetic macular oedema	62.5	63.5	85	43	Ranibizumab	5	12	12
Li <i>et al.</i> [81]	2019	NCT02259088	China	Diabetic macular oedema	46.4	58.7	307	77	Ranibizumab	7.9	12	12
Lim <i>et al.</i> [82]	2016	ACTRN12611000888965	Australia	Diabetic macular oedema	73.2	66.4	17	24	Bevacizumab	2.71	9	9
Lucatto et al. [83]	2017	NR	Brazil	Central retinal vein occlusion	60.0	59.5	14	21	Bevacizumab	NR	12	12
Massin et al. [84]	2010	NCT00284050	France	Diabetic macular oedema	53.6	64	102	49	Ranibizumab	10.2	12	12
Mitchell et al. [85]	2011	NCT00687804	Australia	Diabetic macular oedema	58.2	63.5	234	111	Ranibizumab	7	12	12
Moradian et al. [86]	2011	NCT00370851	Iran	Branch retinal vein occlusion	41.9	57.6	42	39	Bevacizumab	2	2	ю
Motta <i>et al.</i> [87]	2019	NCT02308644	Italy	Diabetic macular oedema	48.8	61.8	20	21	Bevacizumab	2	2	ŝ
Nguyen et al. [88]	2012	NCT00473330	USA	Diabetic macular oedema	57.3	62.7	252	130	Ranibizumab	21.2	24	24
Nguyen et al. [88]	2012	NCT00473382	USA	Diabetic macular oedema	56.2	62.1	250	127	Ranibizumab	21.2	24	24
Nguyen et al. [89]	2019	NCT02302079	USA	Diabetic macular oedema	50.0	61.5	64	32	Ranibizumab	4	3	9
Ogura <i>et al.</i> [90]	2014	NCT01012973	Japan	Central retinal vein occlusion	55.6	61.5	106	71	Aflibercept	8.5	12	18
Parodi et al. [91]	2012	NCT01327222	Italy	Juxtafoveal choroidal	38.1	71.5	11	10	Bevacizumab	NR	9	9
				neovascularisation								
Parodi et al. [92]	2010	NR	Italy	nAMD	31.5	47.9	19	35	Bevacizumab	3.8	24	24
Parodi et al. [93]	2015	UMIN000005014	Italy	Branch retinal vein occlusion		66.8	17	18	Bevacizumab	9	12	12
Patwardhan et al. [94]	2011	NR	India	Vitreous haemorrhage	100.0	26	10	10	Bevacizumab	NR	3	ŝ
Pielen et al. [95]	2015	NCT00562406	Germany	Branch retinal vein occlusion	50.0	66.3	20	10	Ranibizumab	б	33	9
Preti <i>et al.</i> [96]	2013	NCT01389505	Brazil	Proliferative diabetic retinopathy	66.6	56	35	35	Bevacizumab	2	1	9
Raizada <i>et al.</i> [97]	2015	NR	Kuwait	Diabetic macular oedema		53.9	22	22	Bevacizumab	б	3	б
Rajendram et al. [98]	2012	2007-000847-89	UK	Diabetic macular oedema	68.8	64.2	42	38	Bevacizumab	ŝ	б	24

YarTrial IDCountryByc diseaseMen(yeas), neanTreatment, nOmtols, nInjections12012NCT01178697IranCountryEyc disease(%)mennnVEGFitype(n).men12010NCT001585619BrazilDiabetic macular occlusion55.0605032.2Bevacizumab11.32000NCT00056836USAnAMDS5.2772611Rambizumab5.92011NRGreecePolypoidal choroidal43.3671414Bevacizumab5.92010NCT0036836USAAMD37.2771911Rambizumab6.82011NRGreecePolypoidal choroidal43.3671414Bevacizumab5.92017NCT0036832USADiabetic macular ocdema37.17825728242011OH463-69GermanyPolifierative diabetic retriopathy73.066687.9264.42011NRNarcPakistanProlifierative diabetic retriopathy73.0667.027282.62011NRNRPakistanProlifierative diabetic retriopathy73.0663.72.68.44.42011NRProlifierative diabetic retriopathy73.0672.32.78.44.42012NRProlifierative diabetic retriopathy73							Age					of	of
2012 NCT01178697 Iran Central retinal vein occlusion 55.0 60 50 32 Bevacizumab 11.3 2020 NCT02985619 Brazil Diabetic macular oedema 41.5 62.4 478 238 Ranibizumab 59 2006 NCT00256356 USA nAMD 35.2 77 26 11 Ranibizumab 59 2009 NR Greece Retinal angiomatous proliferation 37.0 77 19 11 Ranibizumab 68 2011 NR Greece Rolpoidal choroidal 43.3 67 14 14 $Bevacizumab$ 68 2017 NR Pakistan Proliferative diabetic retinopathy 73.0 56.6 68 19 Bevacizumab 7 2017 NR Pakistan Proliferative diabetic retinopathy 73.0 56.6 68 19 Bevacizumab 7 2017 NR Diabetic macular oedema 51.0 $65.$	Author	Year	Trial ID	Country	Eye disease	Men (%)	(years), mean	Treatment, n	Controls n		Injections (n) , mean	treatment (months)	follow-up (months)
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	Ramezani <i>et al.</i> [99]	2012	NCT01178697	Iran	Central retinal vein occlusion	55.0	60	50	32	Bevacizumab	11.3	12	12
$ \begin{array}{llllllllllllllllllllllllllllllllllll$	Rodrigues et al. [100]	2020	NCT02985619	Brazil	Diabetic macular oedema	41.5	62.4	478	238	Ranibizumab	24	24	24
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	Rosenfeld et al. [101]	2006	NCT00056836	USA	nAMD	35.2	77	26	11	Ranibizumab	5.9	9	9
	Rouvas et al. [102]	2009	NR	Greece	Retinal angiomatous proliferation	37.0	77	19	11	Ranibizumab	9	12	12
2009 2005-003288-21 Austria nAMD 33.1 78 25 Bevacizumab 3 2017 NR Pakistan Proliferative diabetic retinopathy 33.1 78 25 Bevacizumab 3 2007 NCT00336323 USA Diabetic macular oedema 61.0 65 73 62 Ranibizumab 7 2011-004463-69 Germany Radiation retinopathy 81.0 67 23 27 Bevacizumab 7 2016 NR USA Diabetic macular oedema 61.0 65 73 25 Ranibizumab 7 2016 NR USA Diabetic macular oedema 50.0 63.2 116 116 414 109 2017 ISRCTN32207582 UK Proliferative diabetic retinopathy 66.8 51.2 51 84 414 101 2017 ISRCTN32207582 UK Proliferative diabetic retinopathy 66.8 51.2 51 84 414 111	Rouvas et al. [103]	2011	NR	Greece	Polypoidal choroidal	43.3	67	14	14	Bevacizumab	6.8	12	12
2009 2005-003288-21 Austria nAMD 32.1 78 25 Bevacizumab 3 55 2017 NR Pakistan Proliferative diabetic retinopathy 73.0 56.6 68 19 Bevacizumab 3 1 2007 NCT00336323 USA Diabetic macular oedema 61.0 65 73 62 Ranibizumab 5 1 2020 2011-004463-69 Germany Radiation retinopathy 81.0 67 23 27 Bevacizumab 7 2016 NR USA Diabetic macular oedema 50.0 63.2 116 116 Afibercept 4.4 2010 2017 ISRCTN32207582 UK Proliferative diabetic retinopathy 66.8 51.2 51 50 Bevacizumab 2.6 101 2017 ISRCTN32207582 UK Proliferative diabetic retinopathy 66.8 51.2 51 80 8.4 1010 2017 ISRCTN3325075 France Branch					vasculopathy								
5] 2017 NR Pakistan Proliferative diabetic retinopathy 73.0 56.6 68 19 Bevacizumab 2 2007 NCT00336323 USA Diabetic macular oedema 61.0 65 73 62 Ranibizumab 5 1 2020 2011-004463-69 Germany Radiation retinopathy 81.0 67 23 27 Bevacizumab 7 2016 NR USA Diabetic macular oedema 50.0 63.2 116 116 Afibercept 4.4 2010 2017 ISRCTN32207582 UK Proliferative diabetic retinopathy 66.8 51.2 51 50 Bevacizumab 2.6 110] 2017 NCT0159950 Iran Diabetic macular oedema 50.7 61 36.3 13 Ranibizumab 2.6 111 2017 NCT0159950 France Branch retinal vein occlusion 50.4 66.1 429 13 Ranibizumab 7.1 2014 ACTRN15607000262404	Sacu et al. [104]	2009	2005-003288-21	Austria	nAMD	32.1	78	25	25	Bevacizumab	б	ŝ	4
2007 NCT00336323 USA Diabetic macular oedema 61.0 65 73 62 Ranibizumab 5 1 2020 2011-004463-69 Germany Radiation retinopathy 81.0 67 23 27 Bevacizumab 7 2016 NR USA Diabetic macular oedema 50.0 63.2 116 116 Afibercept 4.4 2010 2017 ISRCTN33207582 UK Proliferative diabetic retinopathy 66.8 51.2 51 50 Bevacizumab 2.6 110] 2017 NCT01599650 France Branch retinal vein occlusion 50.4 66.1 429 13 Ranibizumab 11 2017 NCT01599650 France Branch retinal vein occlusion 50.4 66.1 429 13 Ranibizumab 11 2017 NCT01599650 France Branch retinal vein occlusion 50.7 61 36.3 13 Ranibizumab 11 2017 NCT01599650 France Branch retinal vein occlusion 50.7 61 429 13 Ranibizumab	Sameen et al. [105]	2017	NR	Pakistan	Proliferative diabetic retinopathy	73.0	56.6	68	19	Bevacizumab	2	2	ю
2020 2011-004463-69 Germany Radiation retinopathy 81.0 67 23 27 Bevacizumab 7 2016 NR USA Diabetic macular oedema 50.0 63.2 116 116 Afibercept 4.4 2017 ISRCTN32207582 UK Proliferative diabetic retinopathy 66.8 51.2 51 50 Bevacizumab 2.6 2012 NCT00370669 Iran Diabetic macular oedema 50.7 61 36.3 13 Ranibizumab 11 2017 NCT01599650 France Branch retinal vein occlusion 50.4 66.1 429 13 Ranibizumab 11 2017 NCT01599650 France Branch retinal vein occlusion 50.4 66.1 429 13 Ranibizumab 11 2017 NCT01599650 France Branch retinal vein occlusion 50.4 66.3 15 21 Ranibizumab 7.1 2014 ACTRN12607000262404 Australia Branch retinal vein occlusion 59.5 61 14 14 18 20.1 2010	Scott et al. [106]	2007	NCT00336323	USA	Diabetic macular oedema	61.0	65	73	62	Ranibizumab	Ŋ	12	12
2016 NR USA Diabetic macular oedema 50.0 63.2 11.6 11.6 Aflibercept 4.4 2017 ISRCTN32207582 UK Proliferative diabetic retinopathy 66.8 51.2 51 50 Bevacizumab 2.6 2012 NCT00370669 Iran Diabetic macular oedema 50.7 61 36.3 13 Ranibizumab 2.1 2017 NCT01599650 France Branch retinal vein occlusion 50.4 66.1 429 13 Ranibizumab 11 2017 NCT01599650 France Branch retinal vein occlusion 59.7 66.3 15 2.1 Ranibizumab 11 2014 ACTRN12607000262404 Australia Branch retinal vein occlusion 59.7 66.3 15 2.1 Ranibizumab 7.1 2010 ISRCTN83325075 UK nAMD 59.5 81 14 12 Bevacizumab 7.1 2010 ISRCTN83325075 UK nAMD 59.5 81	Seibel et al. [107]	2020	2011-004463-69	Germany	Radiation retinopathy	81.0	67	23	27	Bevacizumab	7	9	7
2017 ISRCTN32207582 UK Proliferative diabetic retinopathy 66.8 51.2 51 50 Bevacizumab 2.6 2012 NCT00370669 Iran Diabetic macular oedema 50.7 61 363 13 Ranibizumab 11 2017 NCT01599650 France Branch retinal vein occlusion 50.4 66.1 429 13 Ranibizumab 11 2017 NCT01599650 France Branch retinal vein occlusion 59.7 66.3 15 21 Ranibizumab 11 2014 ACTRN12607000262404 Australia Branch retinal vein occlusion 59.7 66.3 15 21 Ranibizumab 7.1 2014 ACTRN12607000262404 Australia Branch retinal vein occlusion 59.7 66.3 15 21 Ranibizumab 7.1 2010 ISRCTN83325075 UK nAMD 59.5 81 14 12 Bevacizumab 4.5 2008 NR Iran Nervascular plaucoma 32.1	Shah <i>et al.</i> [108]	2016	NR	NSA	Diabetic macular oedema	50.0	63.2	116	116	Aflibercept	4.4	12	12
2012 NCT00370669 Iran Diabetic macular oedema 50.7 61 363 13 Ranibizumab 11 2017 NCT01599650 France Branch retinal vein occlusion 50.4 66.1 429 13 Ranibizumab 11 2017 NCT01599650 France Branch retinal vein occlusion 59.7 66.3 15 21 Ranibizumab NR 2014 ACTRN1260700262404 Australia Branch retinal vein occlusion 47.2 67.9 65 66 Bevacizumab 7.1 2010 ISRCTN83325075 UK nAMD 59.5 81 14 14 Bevacizumab 4.5 2008 NR Switzerland nAMD 32.1 78 10 17 60 121 63 Ranibizumab 10 2009 NR Iran Neovascular glancoma 80.7 60 121 63 Ranibizumab 10	Sivaprasad et al. [109]	2017	ISRCTN32207582	UK	Proliferative diabetic retinopathy	66.8	51.2	51	50	Bevacizumab	2.6	24	24
2017 NCT01599650 France Branch retinal vein occlusion 50.4 66.1 429 13 Ranibizumab 11 2017 NCT01599650 France Branch retinal vein occlusion 59.7 66.3 15 21 Ranibizumab NR 2014 ACTRN1260700262404 Australia Branch retinal vein occlusion 47.2 67.9 65 66 Bevacizumab 7.1 2010 ISRCTN83325075 UK nAMD 59.5 81 14 14 Bevacizumab 4.5 2008 NR Switzerland nAMD 32.1 78 14 12 Bevacizumab 3 2009 NR Iran Nervascular glancoma 80.7 60 121 63 Ranibizumab 10	Soheilian et al. [110]	2012	NCT00370669	Iran	Diabetic macular oedema	50.7	61	363	13	Ranibizumab	11	24	24
2017 NCT01599650 France Branch retinal vein occlusion 59.7 66.3 15 21 Ranibizumab NR 2014 ACTRN1260700262404 Australia Branch retinal vein occlusion 47.2 67.9 65 66 Bevacizumab 7.1 2010 ISRCTN83325075 UK nAMD 59.5 81 14 14 Bevacizumab 4.5 2008 NR Switzerland nAMD 32.1 78 14 12 Bevacizumab 3 2009 NR Iran Nervascular glancoma 80.7 60 121 63 Ranibizumab 10	Tadayoni et al. [111]	2017	NCT01599650	France	Branch retinal vein occlusion	50.4	66.1	429	13	Ranibizumab	11	24	24
2014 ACTRN12607000262404 Australia Branch retinal vein occlusion 47.2 67.9 65 66 Bevacizumab 7.1 2010 ISRCTN83325075 UK nAMD 59.5 81 14 Bevacizumab 4.5 2010 ISRCTN83325075 UK nAMD 59.5 81 14 14 Bevacizumab 4.5 2008 NR Switzerland nAMD 32.1 78 14 12 Bevacizumab 3 2009 NR Iran Nervascular elancoma 80.7 60 121 63 Ranihizumab 10	Tadayoni et al. [111]	2017	NCT01599650	France	Branch retinal vein occlusion	59.7	66.3	15	21	Ranibizumab	NR	12	12
2010 ISRCTN83325075 UK nAMD 59.5 81 14 14 Bevacizumab 4.5 2008 NR Switzerland nAMD 32.1 78 14 12 Bevacizumab 3 2009 NR Iran Nervascular elaucoma 80.7 60 121 63 Ranihizumab 10 2	Tan <i>et al.</i> [112]	2014	ACTRN12607000262404	Australia	Branch retinal vein occlusion	47.2	67.9	65	99	Bevacizumab	7.1	12	12
2008 NR Switzerland nAMD 32.1 78 14 12 Bevacizumab 3 2009 NR Iran Neovascular glaucoma 80.7 60 121 63 Ranihizumah 10 2	Tufail <i>et al.</i> [113]	2010	ISRCTN83325075	UK	nAMD	59.5	81	14	14	Bevacizumab	4.5	9	9
2009 NR Iran Neovascular glaucoma 80.7 60 121 63 Ranihizumah 10	Weigert et al. [114]	2008	NR	Switzerland	nAMD	32.1	78	14	12	Bevacizumab	ю	33	9
	Yazdani et al. [115]	2009	NR	Iran	Neovascular glaucoma	80.7	60	121	63	Ranibizumab	10	24	24

Table 2: Continued.

	Experimenta				
Study	Events Tota	I Events Total	Risk Ratio	RR	95%-CI Weight
Non diabetic					
Brown 2010	0 132	2 0.5 130		0.49	[0.02; 14.55] 0.5%
Brown 2010	0 130	0.5 130		0.50	[0.02; 14.77] 0.5%
Brown 2006	12 280) 12.0 143		0.51	[0.24; 1.11] 7.2%
Tadayoni 2017 (2)	48 429			0.73	[0.20; 2.68] 1.8%
Tadayoni 2017 (1)	41 363			0.73	[0.20; 2.71] 1.8%
Brown 2009	17 140			0.75	[0.42; 1.35] 10.4%
Ogura 2014	4 106			0.89	[0.21; 3.87] 1.6%
Rosenfeld 2006	80 478			1.05	[0.74; 1.49] 23.1%
Hoerauf 2016	5 124			1.20	[0.33; 4.36] 1.9%
Abraham 2010	19 12			1.41	[0.63; 3.18] 4.2%
Bandello 2018	10 153			2.01	[0.70; 5.75] 2.3%
Brown 2011	1 132			2.98	[0.12; 72.42] 0.2%
Tan 2014	2 15			6.94	[0.36; 134.55] 0.2%
Bashshur 2007	0 32				0.0%
Common effect mode		5 1399	Ŷ	0.99	[0.78; 1.25] 55.6%
Heterogeneity: $I^2 = 0\%$, τ	$^{-} = 0, p = 0.69$				
Diabetic					
Bhavsar 2013	0 125	5 3.0 136		0.16	[0.01; 2.98] 1.5%
Li 2019	19 307			0.48	[0.23; 0.98] 7.3%
Baker 2019	4 226			0.53	[0.18; 1.56] 4.7%
Scott 2007	2 68			0.56	[0.05; 5.84] 0.7%
Lang 2018	14 85			0.79	[0.37; 1.67] 5.4%
Massin 2010	9 102	2 5.0 49		0.86	[0.31; 2.44] 3.1%
Ishibashi 2015	8 132	2 7.0 131	<u> </u>	1.13	[0.42; 3.04] 3.2%
Gross 2018	38 19	28.0 203	h -	1.44	[0.92; 2.25] 12.4%
Nguyen 2012	5 250) 1.0 127		2.54	[0.30; 21.51] 0.6%
Rajendram 2012	1 42			2.72	[0.11; 64.75] 0.2%
Elman 2010	25 375			2.90	[1.45; 5.82] 4.4%
Mitchell 2011	3 234			3.33	[0.17; 63.88] 0.3%
Figueira 2018	1 4				[0.14; 80.28] 0.2%
Nguyen 2012	4 252			4.65	[0.25; 85.74] 0.3%
Motta 2019	0 20				0.0%
Common effect mode			¢	1.20	[0.93; 1.54] 44.4%
Heterogeneity: $I^2 = 40\%$,	τ ⁻ = 0.2270, <i>p</i> =	0.06			
Common effect mode	508	5 3485	-	1.08	[0.91; 1.28] 100.0%
Heterogeneity: $I^2 = 18\%$,					
Test for subgroup differen			.01 0.1 1 10 10	0	

Figure 2: Forest plot of RRs for hypertension: frequentist meta-analysis using fixed and random effects models, stratified by diabetic versus non-diabetic eye disease.

cardiovascular disease in two trials (2.4%; 1115 participants) trials, previous MI or stroke in one trial (1.2%; 702 participants) and eGFR in one trial (1.2%; 41 participants: Table 2). Baseline proteinuria or a history of heart failure was not reported in any trial.

Cardiorenal outcomes

Cardiorenal outcomes were reported in only a minority of trials. Of these, hypertension was recorded most often in 29 trials (35.4%; 8570 participants). Hypertension was not more common in those treated with VEGFi versus controls [7.3 versus 5.4%; RR 1.08 (95% CI 0.91–1.28), P = .369; Fig. 2].

New or worsening heart failure was recorded in 10 trials (12.2%; 3384 participants), with an incidence of 2.8% versus 3.2% in VEGFi-treated patients and controls, respectively [RR 1.03 (95% CI 0.70–1.51), P = .894; Fig. 3A]; proteinuria was recorded in 5 trials (6.1%; 1902 participants) and was detectable in some VEGFi-treated participants (0.2%) but not controls [0.0%; RR 4.43 (95% CI 0.49–40.0), P = .185; Fig. 3B]. *De novo* CKD or nephropathy was recorded in nine trials (11.0%; 3471 participants), with a similar proportion in the VEGFi (1.8%) versus control groups [1.4%; RR 1.00 (95% CI 0.55–1.81), P = 1.00; Fig. 3C]; however, absolute values of eGFR were not recorded in any trial. Meta-regression analyses did not identify any variation in heterogeneity according to

(A)	Experim	ental	Co	ontrol			
Study			Events		Risk Ratio	RR	95%-Cl Weight
Diabetic Mitchell 2011 Gale 2018 Bhavsar 2013	1 0 0	234 99 125	3 1 1	111 99 136		0.33	[0.02; 1.50] 8.3% [0.01; 8.08] 3.1% [0.01; 8.82] 2.9%
Shah 2016 Nguyen 2012 Baker 2019	0 5 8	23 252 226	1 6 22	27 130 476		0.43 0.77	[0.02; 9.13] 2.8% [0.13; 1.38] 16.2% [0.35; 1.69] 29.0%
Scott 2007 Elman 2010 Nguyen 2012 Lang 2018	1 24 9 1	68 375 250 85	0 17 2	19 479 127 43		1.80	[0.04; 20.15] 1.6% [0.98; 3.31] 30.6% [0.50; 10.42] 5.4% 0.0%
Common effect mode Heterogeneity: $I^2 = 27\%$		1737	0.20	1647	\$	1.03	[0.70; 1.51] 100.0%
Common effect mode Heterogeneity: $I^2 = 27\%$ Test for subgroup differe	$\tau^2 = 0.2829$			1647 NA)	0.1 0.51 2 10	1.03	[0.70; 1.51] 100.0%

(B)

Study	Experim Events			ntrol Total	F	lisk Ratio		RR	95%-CI Weight
Non diabetic Brown 2010	0	132	0	130					0.0%
Brown 2010	0	130	0	130					0.0%
Brown 2011	0	132	0	131					0.0%
Diabetic									
Elman 2010	1	375	0	479				- 3.83	[0.16; 93.77] 46.7%
Ishibashi 2015	2	132	0	131	-			- 4.96	[0.24; 102.37] 53.3%
Common effect mode		507		610				4.43	[0.49; 40.02] 100.0%
Heterogeneity: $I^2 = 0\%$, τ	$p^{2} = 0, p = 0$.91							
Common effect mode Heterogeneity: $I^2 = 0\%$, τ	$p^2 = 0, p = 0$			1001	I				[0.49; 40.02] 100.0%
Test for subgroup differer	ices: $\chi_0^2 = 0$.00, df	= 0 (<i>p</i> = 1	VA) 0.01	0.1	1	10	100	

(C)

Study	Experim Events		Co Events	ntrol Total	Risk Ratio	RR	95%–Cl Weight
Non diabetic Heier 2006	1	106	0	56		1.59	[0.07; 38.44] 2.8%
Diabetic Bhavsar 2013 Massin 2010 Baker 2019 Elman 2010 Nguyen 2012 Li 2019 Nguyen 2012 Gale 2018 Common effect mode Heterogeneity: $I^2 = 0\%$, 1		125 102 226 375 252 307 250 99 1736 0.83	2 1 11 3 2 1 0	136 49 476 479 130 77 127 99 1573		- 3.00	[0.03; 7.52]5.8%[0.16; 2.04]30.7%[0.14; 5.07]11.4%[0.32; 4.58]17.1%
Common effect mode Heterogeneity: $I^2 = 0\%$, Test for subgroup different	$p^2 = 0, p = 0$			1629).77)	0.1 0.51 2 10	1.00	[0.55; 1.81] 100.0%

Figure 3: Forest plot of RRs for (A) heart failure, (B) proteinuria and (C) CKD: frequentist meta-analysis using fixed and random effects models, only recorded in trials of diabetic eye disease.

VEGFi subtype, number of injections, duration of treatment or eye disease treated (Supplementary Table S1). Funnel plots and trim-and-fill analyses did not detect any substantial evidence of publication bias for any of the outcomes (although the proteinuria outcome was not tested due to insufficient data; Supplementary Figs. S2–S4).

Arterial thrombotic cardiovascular events and death

In 39 comparisons (48.1%) including 10 133 participants, the absolute incidence of arterial thrombotic cardiovascular events (MI, stroke and peripheral arterial disease) was similar in the VEGFi-treated groups compared with controls [3.2 versus 3.0%, respectively; RR 1.19 (95% CI 0.95-1.48), P = .122; Supplementary Fig. S1). In 41 comparisons (50.6%; 9877 participants), the rate of all-cause mortality was similar in the VEGFi and control groups [1.6 versus 1.3%; RR 1.24 (95% CI 0.89–1.73), P = .198]. Meta-regression analysis identified an increased risk of death in patients treated for diabetic eye disease (Supplementary Table S1 and Supplementary Fig. S5). In the subgroup of participants treated for diabetic eye disease, the rate of all-cause mortality was higher in the VEGFitreated group [RR 1.62 (95% CI 1.04–2.46), P = .020; Fig. 4]. Funnel plots and trim-and-fill analyses did not detect any substantial evidence of publication bias for cardiovascular event (Supplementary Fig. S6) or all-cause mortality models (Supplementary Fig. S7).

Longer follow-up studies

Three studies were identified as constituting extended follow-up from initial trials and reporting cardiorenal adverse events or death [16–18]. Of these, none were suitable for additional analysis: two were excluded, as all treatment groups were eligible to receive intravitreal VEGFi [17, 19] and one was excluded because it reported total systemic adverse events per group without quantifying cardiorenal events and/or death [18].

DISCUSSION

We have not identified an increased risk of cardiorenal outcomes—including hypertension, proteinuria, heart failure and *de novo* CKD—in randomised controlled trials of intravitreal VEGFis nor have we identified an increased risk of arterial thrombotic cardiovascular events, even though populations in which these agents are used are at high risk for these events. However, there are insufficient reported data to definitively confirm or refute any link between these agents and adverse cardiorenal outcomes. In the subgroup of patients treated for diabetic eye disease, we identified an increased risk of death associated with VEGFi treatment.

This is the first systematic review to explore the incidence and reporting of renal adverse events and heart failure after intravitreal VEGFi. Prior reviews have made a disproportionate effort to capture arterial thrombotic cardiovascular events and death [20–23], although mechanistically hypertension, heart failure and CKD are more likely sequelae [24]. In a systematic review of systematic reviews, intravitreal VEGFi treatments were not found to be associated with an increased risk of systemic adverse events, predominantly focusing on arterial thrombotic cardiovascular events and death [20]. However, in restricted analyses in participants with diabetic eye disease, associations have been demonstrated between intravitreal VEGFi and risk of stroke and vascular death [22] and with allcause mortality [25]. We have similarly found an association between intravitreal VEGFi and death in the subset of patients treated for diabetic eye disease.

If intravitreal VEGFis are associated with a higher risk of premature death, this may have important implications for informed consent for patients with diabetes and kidney disease, who are already at higher risk from their underlying disease. However, the absolute risk of death associated with VEGFis may be outweighed by benefits to quality of life, such as preservation of visual acuity in VEGFi-treated patients. We do not believe the current trial data are adequate to quantify differences in the absolute risks-of death with treatment and visual loss without-to inform the consent process. First, only a minority of trials report cardiorenal, arterial thrombotic and death events: the risk of death or other significant events may be underestimated due to underreporting. Second, the duration of follow-up in trials is relatively short: except one trial with a 5-year follow-up, the maximum observation period in the included trials was 2 years. This is unlikely to be long enough to capture the cumulative risks associated with prolonged treatment with intravitreal VEGFis, particularly as kidney disease and heart failure may not manifest clinically until much later in the disease course. Third, patients in higher-risk populations, such as those with diabetes, preexisting CKD, heart failure or a combination of these may be at increased risk, but this is not adequately recorded to assess in current trials. Fourth, it has been observed that patients with multiple medical conditions are underrepresented in clinical trials [26], with higher recorded adverse events in the general population compared with the trial populations [27]: this is also likely to be true for trials of intravitreal treatments. We may be able to quantify absolute risks of visual loss, cardiorenal side effects and death through the analysis of large, longterm, real-world databases ('big data'); however, with many confounding factors in population studies, particularly in cohorts of patients requiring VEGFis for diabetic eye disease, we acknowledge that it will be challenging to identify causal relationships.

We have not identified an increased risk of cardiorenal outcomes in intravitreal VEGFi-treated participants in the published trials. Since hypertension is an almost ubiquitous sequela in patients treated with intravenous VEGFi [24], it may be that the definitions or thresholds used within the trials were not sensitive enough to detect treatment-related hypertension. None of the 29 trials reporting hypertension as an outcome specified a definition of hypertension and there were inadequate raw data reported to assess absolute changes in BP over the treatment period. Similarly, reporting of renal outcomes were inconsistent, insensitive and non-specific across studies. One trial [28] reported >10 renal complications (including 'acute kidney injury', 'acute renal failure', 'chronic

Non diabetic $1 adayoni 2017 (2)$ 5 429 2 13 0.08 $0.02; 0.35$ 6.59 Tadayoni 2017 (1) 3 363 0 13 0.26 $0.01; 4.79$ 1.69 Tan 2014 0 15 1 21 0.66 $0.22; 0.61$ 2.12 Abraham 2010 2 121 2 63 0.52 $0.08; 3.61$ 4.49 Brown 2006 11 478 6 238 0.91 $0.34; 2.44$ 13.49 Brown 2006 5 280 2 143 1.28 $0.25; 6.50$ 4.49 Tufail 2010 1 65 0 66 3.05 $[0.13; 73.42]$ 0.89 Bashshur 2007 0 32 0 30 0.09 Brown 2010 132 131 0.09 0.09 Brown 2010 0 126 0 0.09 Haterbach 2018 0 24 119 0.09	Study
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Figure 4: Forest plot of RRs for all-cause mortality: frequentist meta-analysis using fixed and random effects models, stratified by diabetic versus non-diabetic eye disease.

kidney disease', 'kidney failure', 'end-stage kidney disease'), many of which are likely to be overlapping. It is impossible to know whether raw data values are available in the trial site files but have not reached publication. This review illustrates the potential utility of presenting absolute values of pre- and post-treatment BP, proteinuria, kidney function (e.g. eGFR) and objective measures of heart failure (e.g. N-terminal brain natriuretic peptide and left ventricular EF) in patients treated with VEGFis by any route. These would be substantially more informative and sensitive to detecting treatment-related side effects.

Recent evidence highlights the importance of assessing and reporting important systemic outcomes in trials, particularly in high-risk groups. Rosiglitazone was previously used widely for glycaemic control in type 2 diabetes; however, scrutiny of the published trial data showed that rosiglitazone was unexpectedly associated with increased odds of MI and death [29]. Bardoxolone methyl was tested to slow progression of diabetic kidney disease, but the trial was stopped early due to an increased rate of cardiovascular events in the treatment group compared with placebo [30]. Atrasentan, tested in patients with diabetic kidney disease, was shown to reduce the risk of renal decline or end-stage kidney disease, but there was a signal towards an increased rate of heart failure hospitalisations [31]. In clinical practice, hypertension, proteinuria, renal decline and heart failure are all common in patients with diabetes. In the absence of guidelines to test and monitor for cardiorenal side effects after introducing intravitreal VEGFi, it may be difficult to distinguish whether new or worsened cardiorenal effects are related to the progression of diabetic complications or whether they could have been exacerbated by intravitreal treatments. There are numerous published case reports of de novo renal sequelae after intravitreal VEGFi-particularly bevacizumab and aflibercept-including proteinuria, hypertension, heart failure and progressive renal injury [7, 32-35]. Ranibizumab is similarly absorbed, but is cleared far more quickly due to its structure and size. Although ranibizumab is associated with fewer reports of cardiorenal side effects compared with bevacizumab and aflibercept, it has also been associated with hypertension, thrombotic microangiopathy and renal injury [36-38]. It is desirable to scrutinise population and prescription data to assess the prevalence of these cardiorenal sequelae after intravitreal VEGFi treatments. If concerns about cardiorenal safety are confirmed, this may encourage trialists, regulators and guideline developers to monitor for these potential sequelae, adjust ophthalmic treatment regimens and provide better information for informed consent for patients.

We acknowledge some limitations of this work. First and most important, the cardiorenal adverse events were secondary outcomes in these trials: only a limited number of trials reported the incidence of our cardiorenal outcomes of interest, limiting the power to detect a signal for cardiorenal side effects. We did not find evidence of additional reporting of systemic adverse events in secondary trial publications. Second, the baseline cardiometabolic phenotype of the participants in these trials was not well-described: both comorbidities and absolute values of markers of cardiorenal disease (eGFR, BP, proteinuria, EF) were rarely reported, although it is likely that vital signs/BP were measured in most. It was not possible to identify subgroups who may be at higher risk of cardiorenal sequelae in the trial populations. Third, in the limited trials that reported rates of cardiorenal adverse events, there was a limited duration of follow-up to detect these risks over the longer term—a common issue in prospective trials. Repeated exposure to systemic absorption of VEGFis over a longer time period may be associated with a high risk of developing cardiorenal side effects that are not detectable within the relatively short follow-up (range 3–60 months, but the majority <24 months).

CONCLUSION

In published trials of intravitreal VEGFis, we did not identify an increased risk of cardiorenal outcomes—including hypertension, proteinuria, heart failure and *de novo* CKD however, there are insufficient data definitively to confirm or refute any link between these agents and adverse cardiorenal outcomes. In keeping with previous analyses, we did not identify an increased risk of arterial thrombotic cardiovascular events, but there was an increased risk of death in the subgroup of patients treated with intravitreal VEGFis for diabetic indications. However, there is increasing evidence for systemic cardiorenal sequelae of intravitreal VEGFis. Additional scrutiny of post-licensing population data may help identify if there are implications for cardiorenal safety and monitoring when prescribing these medications, particularly in high-risk patients with diabetes.

SUPPLEMENTARY DATA

Supplementary data are available at *ndt* online.

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AUTHORS' CONTRIBUTIONS

All authors designed the research and critically reviewed and approved the final manuscript. J.S.L. and S.J.H.D. performed the search and extracted data. J.S.L. collated the data, performed statistical analysis and wrote the first draft of the manuscript.

DATA AVAILABILITY STATEMENT

Extracted data and analysis code will be available from the project GitHub repository on publication (https://github.com/jennifer-s-lees/vegf_sr_meta_analysis_public).

CONFLICT OF INTEREST STATEMENT

J.S.L. has received personal honoraria from Bristol-Myers Squibb, Pfizer and AstraZeneca, outside the submitted work. P.B.M. has received personal honoraria from Vifor, Pharmacosmos, Napp, AstraZeneca, GlaxoSmithKline and Astellas and grants from Boehringer Ingelheim. The University of Glasgow, which employs N.N.L., has received research grant funding from Roche Diagnostics, AstraZeneca and Boehringer Ingelheim (outside the submitted work) and has received speaker fees/advisory board fees from Roche, Pharmacosmos, AstraZeneca and Novartis. The results presented in this article have not been published previously in whole or in part except in abstract form.

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