

Finerenone in Heart Failure and Chronic Kidney Disease with Type 2 Diabetes: the FINE-HEART Pooled Analysis of Cardiovascular, Kidney, and Mortality Outcomes

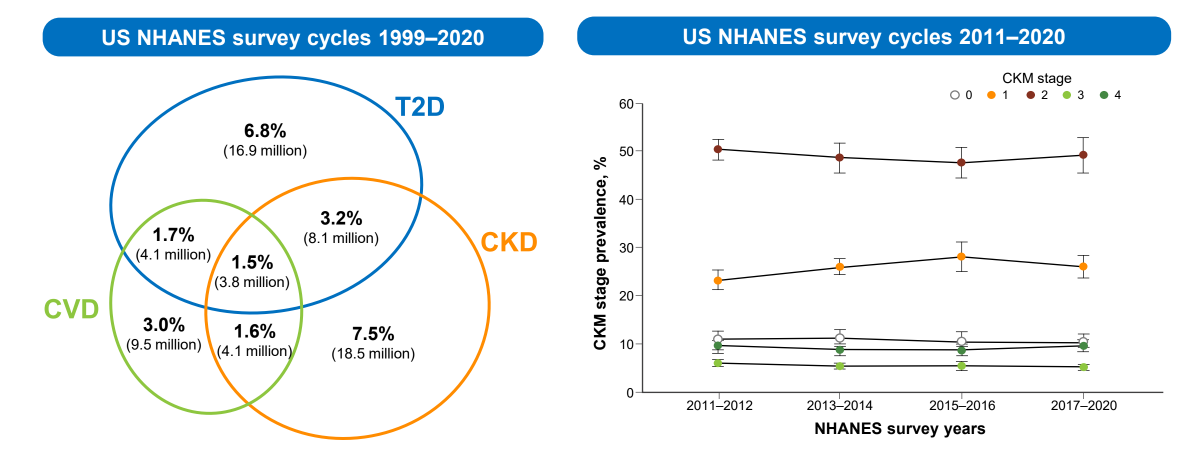
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PROSPERO CRD42024570467



# Strong Epidemiological Overlap of Cardiovascular, Metabolic, and Kidney Disorders



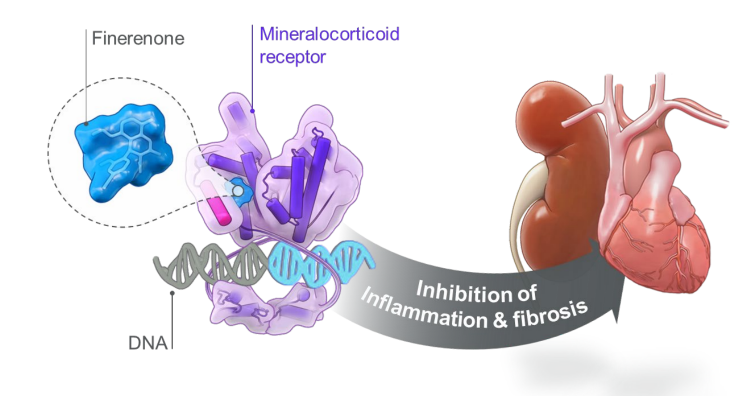






# Could the Non-Steroidal MRA, Finerenone, Modify Risk across the Cardio-Kidney-Metabolic Spectrum?

- Finerenone is a non-steroidal MRA that has been studied in RCTs of patients with T2D and CKD and separately in patients with HF (with and without T2D).
- However, none of these trials were individually powered to evaluate treatment effects on mortality outcomes or effects in key subgroups.







#### Design of FINE-HEART Umbrella Program



**Prospectively Registered:** PROSPERO CRD42024570467

(n=18,991 Participants)







Pooling data in the FINE-HEART program increased precision to robustly assess the efficacy and safety of the non-steroidal MRA finerenone on important cardio-kidney outcomes and is enriched for participants with a high burden of CKM multimorbidity.





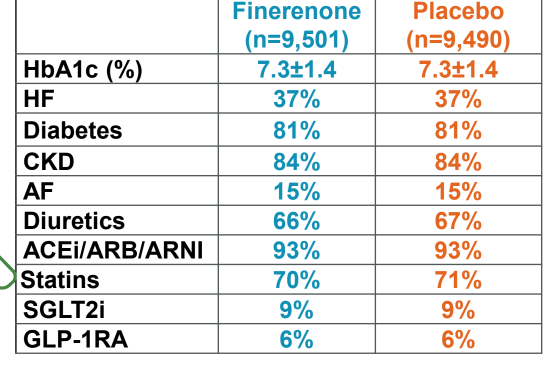
### **Study Designs of the Individual Trials**

	FINEARTS-HF	FIDELIO-DKD and FIGARO-DKD
Validly Randomized	6,001	12,990
Countries	37	48
Patient population	HFmrEF or HFpEF	CKD and T2D
Inclusion criteria	<ul> <li>Adults (≥40 years)</li> <li>Symptomatic HF</li> <li>LVEF ≥40%</li> <li>Elevation natriuretic peptides</li> <li>Structural heart disease</li> <li>Recent diuretic use</li> </ul>	<ul> <li>Adults (≥18 years old)</li> <li>T2D</li> <li>UACR ≥ 30 mg/g</li> <li>Maximally tolerated RASi</li> </ul>
<b>Exclusion criteria</b>	Potassium ≤5.0 mmol/L	Potassium ≤4.8 mmol/L
Dosage and titration	eGFR ≤60: 10 up to 20 mg eGFR >60: 20 up to 40 mg (potentially down to 10 mg)	eGFR <60: 10 up to 20 mg eGFR ≥60: 20 mg (potentially down to 10 mg)
Study duration	2.6 years	2.6 years (FIDELIO-DKD) 3.4 years (FIGARO-DKD)

#### **Baseline Characteristics of FINE-HEART Integrated Population**

Finerenone	Placebo	
(n=9,501)	(n=9,490)	
67±10	67±10	
36%	35%	
72%	72%	
31±6	31±6	
135±15	134±15	
4.4±0.5	4.4±0.5	
59±21	59±21	
1%	1%	
29%	29%	
27%	26%	
44%	44%	
283	293	
[46-836]	[47-855]	
20%	20%	
31%	31%	
49%	50%	
	(n=9,501) 67±10 36% 72% 31±6 135±15 4.4±0.5 59±21 1% 29% 27% 44% 283 [46-836] 20% 31%	

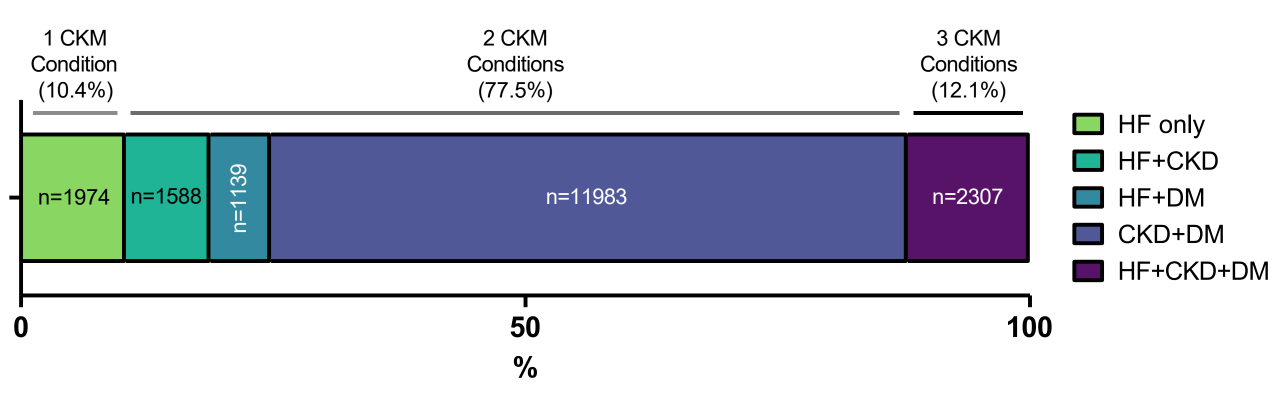




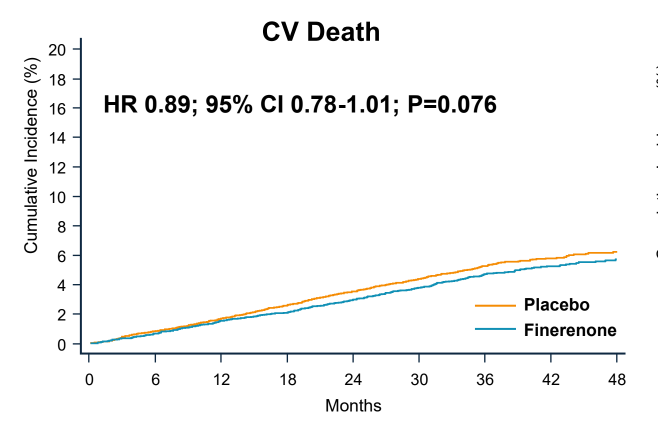


### High Burden of Cardio-Kidney-Metabolic Disease Overlap

#### **Baseline CKM Status in FINEHEART**



#### **Primary Endpoint: CV Death**



**CV Death (including Unknown Death)** Cumulative Incidence (%) 18 HR 0.88; 95% CI 0.79-0.98; P=0.025 16 14 12 -10 -4 **Placebo** 2 **Finerenone** 0 12 18 24 30 36 42 Months

**Primary Analysis: CV Death Excluding Unknown Deaths** 

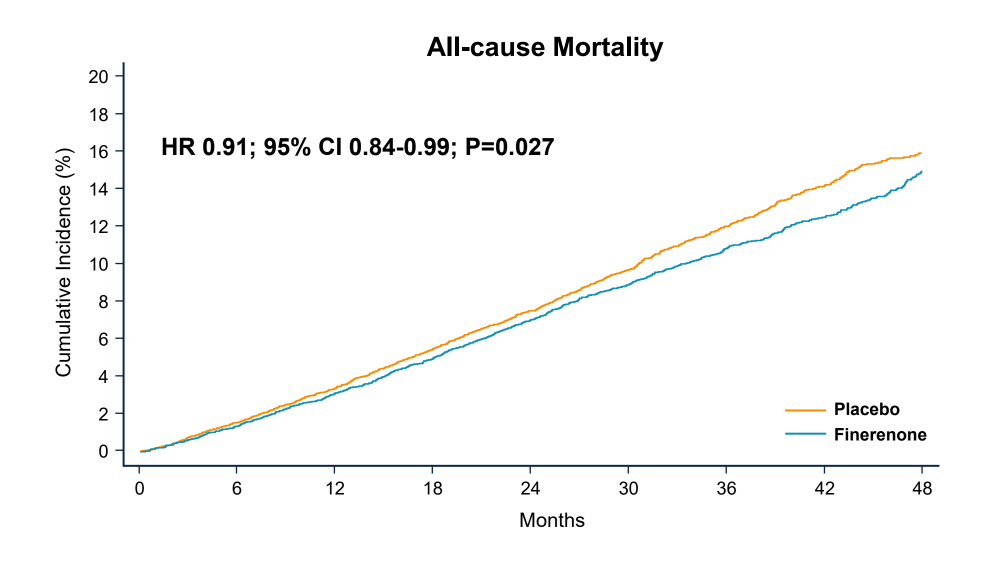
Finerenone 421 (4.4%) vs. Placebo 471 (5.0%)

Prespecified Sensitivity Analysis:

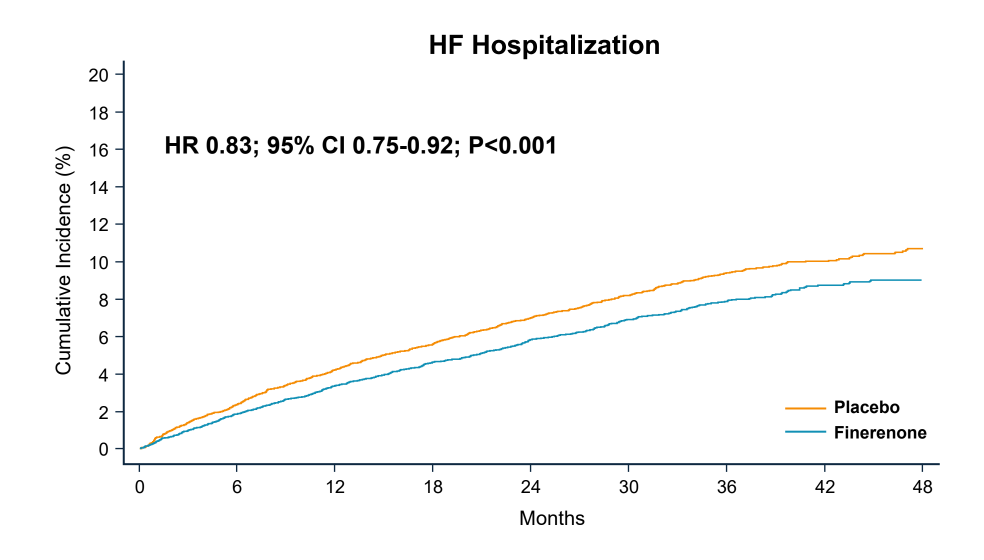
CV Deaths Including Unknown Deaths

Finerenone 627 (6.6%) vs. Placebo 703 (7.4%)

#### **Secondary Endpoint: All-Cause Mortality**

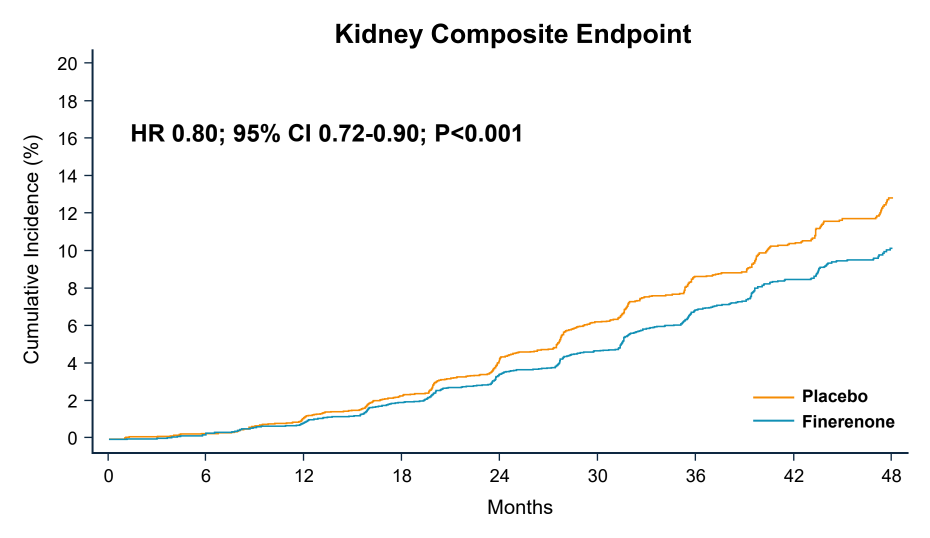


#### **Secondary Endpoint: HF Hospitalization**



#### **Secondary Endpoint: Kidney Composite Endpoint**

sustained eGFR decline of ≥50%, kidney failure\*, or death due to kidney failure



\*sustained eGFR < 15 ml/min/1.73m<sup>2</sup>, chronic dialysis, or kidney transplantion

## **Summary of Prespecified Efficacy Endpoints**

Outcome		HR (95% CI)	P-value
Primary Endpoint			
CV death (excluding undetermined death)		0.89 (0.78–1.01)	0.076
Prespecified sensitivity analysis:  CV death (including undetermined death)		0.88 (0.79–0.98)	0.025
Secondary Endpoints			
Kidney Composite Endpoint	<b>⊢</b>	0.80 (0.72–0.90)	<0.001
HF Hospitalization	<b>⊢</b>	0.83 (0.75–0.92)	<0.001
CV Death or HF Hospitalization	<b>⊢</b>	0.85 (0.78–0.93)	<0.001
New-onset Atrial Fibrillation	<u> </u>	0.83 (0.71–0.97)	0.018
Major Adverse Cardiovascular Events	re-l	0.91 (0.85–0.98)	0.010
All-cause Death	<b>-</b>	0.91 (0.84–0.99)	0.027
All-cause Hospitalization	<b>\rightarrow</b>	0.95 (0.91–0.99)	0.025
All-cause Death or All-cause Hospitalization		0.94 (0.91–0.98)	0.007
	0.5 1	2	

Favors placebo

Favors finerenone

# **Broad Consistency Across 17 Prespecified Subgroups for the Primary Endpoint (CV Death)**

Category	Finerenone (n=9501)	Placebo (n=9490)		HR (95% CI)
eatings. J	n/N	n/N		
Age			l I	
≤ Median	149/5071	179/5053	<del> </del>	0.84 (0.68–1.05
>Median	272/4430	292/4437	<b>₩</b>	0.91 (0.77–1.07)
Sex			i	
Male	265/6111	298/6216	<b>₩</b>	0.87 (0.74–1.03)
Female	156/3390	173/3274	<b>⊢</b>	0.89 (0.72–1.11)
Race			i	
Asian	56/1910	57/1946	<b>⊢</b>	0.98 (0.68–1.42)
Black	7/300	11/308	<del></del>	0.58 (0.22–1.53)
Other	15/476	19/447	<b>—</b>	0.72 (0.37–1.44)
White	343/6815	384/6789	<b>₩</b>	0.89 (0.77–1.03)
Region			į	
Asia	56/1808	55/1815	<b>—</b>	0.99 (0.68–1.44
Eastern Europe	176/3001	187/2941	<b>⊢</b>	0.93 (0.76–1.14)
Latin America	40/1041	69/1034	<b>-</b> → ;	0.58 (0.39–0.85)
North America	43/1259	50/1261	<b>→</b>	0.85 (0.57–1.28)
Western Europe, Oceania, Others	107/2392	110/2439	$\mapsto$	0.98 (0.75–1.28)
Baseline BMI (kg/m²)			į	
<30 mg/m <sup>2</sup>	210/4591	237/4616	₩	0.87 (0.73–1.05)
≥30kg/m²	210/4880	234/4856	н	0.89 (0.74–1.07)
Baseline Systolic Blood Pressure (mmHg)			i	
≤ Median	254/4790	257/4786	₩	1.00 (0.84–1.19)
>Median	1664/4707	214/4701	<b>⊢</b>	0.76 (0.62-0.93)
Baseline Serum Potassium				
≤4.5 mmol/L	284/6746	308/6419	<b>₩</b>	0.91 (0.77–1.06)
>4.5 mmol/L	137/3024	163/3068	<b>⊢</b>	0.86 (0.69-1.08)

**Favours finerenone** 

Favours placebo

Category	Finerenone (n=9501)	Placebo (n=9490)		HR (95% CI)
Category	n/N	n/N		
KDIGO Risk Categories			!	
Low risk	48/1052	50/1034	<u> </u>	0.94 (0.63–1.39)
Moderately increased risk	84/1545	88/1455	<u> </u>	0.89 (0.66–1.20)
High risk	1203/3184	161/3318	<b>⊢</b>	0.78 (0.61–0.98)
Very high risk	157/3616	161/3577	<b>—</b>	0.96 (0.77–1.20)
History of HF				
Present	273/3488	299/3520	<b>₩</b>	0.92 (0.78–1.08)
Absent	148/6013	172/5970	ьф	0.85 (0.68-1.06)
History of Diabetes Mellitus				
Present	294/7715	343/7714	<b>₩</b>	0.85 (0.73–1.00)
Absent	127/1786	128/1776	<b>I</b>	0.98 (0.77–1.25)
History of CKD				
Present	330/7949	363/7929	<b>₩</b>	0.90 (0.77–1.04)
Absent	91/1552	108/1561	<b>⊢</b>	0.84 (0.64–1.11)
Cardio-Kidney-Metabolic Conditions				
1 Condition	58/996	61/978	<b>-</b>	0.93 (0.65–1.33)
2 Conditions	250/7359	286/7351	Ю	0.87 (0.74–1.03)
3 Conditions	113/1146	124/1161	$\vdash \!$	0.91 (0.71–1.18)
GLP-1RA at Baseline			!	
No	403/8925	453/8956	$\mapsto$	0.88 (0.77–1.01)
Yes	18/576	18/534	<b>—</b>	1.05 (0.54–2.07)
SGLT2i at Baseline				
No	375/8672	422/8629	$\mapsto$	0.88 (0.76–1.01)
Yes	46/829	49/861	<u> </u>	0.96 (0.64–1.44)

**Favours finerenone** 

Favours placebo

### **Safety Outcomes**

	Finerenone	Placebo
	n=9,482	n=9,467
Any serious adverse event	35%	37%
Any ae leading to treatment discontinuation	5%	5%
Any potassium >5.5 mmol/L	17%	8%
Any potassium >6.0 mmol/L	3%	1%
Any potassium <3.5 mmol/L	5%	10%
Hyperkalemia	13%	6%
Hyperkalemia leading to discontinuation	1.3%	0.5%
Hyperkalemia leading to hospitalization	0.8%	0.2%
Hyperkalemia leading to death	0%	0%
Acute kidney injury	4%	3%
Acute kidney injury leading to discontinuation	0.2%	0.1%
Acute kidney injury leading to hospitalization	2%	1%
Systolic blood pressure<100mmHg	11%	7%
Gynecomastia or breast hyperplasia	0.2%	0.2%

Treatment-emergent adverse events are defined as any adverse event occurring in any patient who has received at least one dose of study drug and within 3 days of permanent discontinuation. This safety table includes 1 patient who was randomized to placebo but who actually received finerenone.

#### Conclusions

- The FINE-HEART participant-level pooled analysis represents the largest analysis of the effects of the non-steroidal MRA finerenone across the CKM spectrum.
- While this pooled analysis failed to demonstrate significant reductions in cardiovascular death, finerenone was associated with significantly lower deaths of any cause, cardiovascular events, and kidney outcomes.
- Treatment effects were consistent across all tested clinical subgroups including those with multiple intersecting CKM conditions and on background SGLT2i or GLP-1RA.
- No new or unexpected safety signals were uncovered in this pooled analysis.

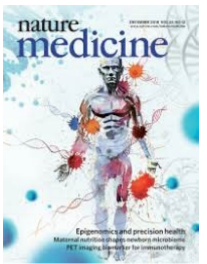
The totality of the evidence supports the disease-modifying potential of finerenone in broad, high-risk patient populations encompassing cardiovascular, kidney, and metabolic diseases.

#### Full Details Available Online in Nature Medicine





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### In Memory of the Late Dr. George Bakris (1952-2024)



A pioneer in cardio-kidney-metabolic research, physician, leader, colleague, and dear friend