

European Heart Journal (2023) **00**, 1–14 European Society https://doi.org/10.1093/eurheartj/ehad283

Dapagliflozin and diuretic utilization in heart failure with mildly reduced or preserved ejection fraction: the DELIVER trial

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Received 4 February 2023; revised 20 April 2023; accepted 1 May 2023

Abstract

Aims

Dapagliflozin reduced the combined risk of worsening heart failure or cardiovascular death among patients with heart failure with mildly reduced or preserved ejection fraction. In this study, the safety and efficacy of dapagliflozin according to background diuretic therapy and the influence of dapagliflozin on longitudinal diuretic use were evaluated.

Methods and results

In this pre-specified analysis of the Dapagliflozin Evaluation to Improve the LIVEs of Patients With Preserved Ejection Fraction Heart Failure (DELIVER) trial, the effects of dapagliflozin vs. placebo were assessed in the following subgroups: no diuretic, non-loop diuretic, and loop diuretic furosemide equivalent doses of <40, 40, and >40 mg, respectively. Of the 6263 randomized patients, 683 (10.9%) were on no diuretic, 769 (12.3%) were on a non-loop diuretic, and 4811 (76.8%) were on a loop diuretic at baseline. Treatment benefits of dapagliflozin on the primary composite outcome were consistent by diuretic use categories ($P_{\rm interaction} = 0.64$) or loop diuretic dose ($P_{\rm interaction} = 0.57$). Serious adverse events were similar between dapagliflozin and placebo arms, irrespective of diuretic use or dosing. Dapagliflozin reduced new initiation of loop diuretics by 32% [hazard ratio (HR) 0.68; 95% confidence interval (CI): 0.55–0.84, P < 0.001] but did not influence discontinuations/disruptions (HR 0.98; 95% CI: 0.86–1.13, P = 0.83) in follow-up. First sustained loop diuretic dose increases were less frequent, and sustained dose decreases were more frequent in patients treated with dapagliflozin: net difference of -6.5% (95% CI: -9.4 to -3.6; P < 0.001). The mean dose of loop diuretic increased over time in the placebo arm, a longitudinal increase that was significantly attenuated with treatment with dapagliflozin (placebo-corrected treatment effect of -2.5 mg/year; 95% CI: -1.5, -3.7, P < 0.001).

Conclusion

In patients with heart failure with mildly reduced or preserved ejection fraction, the clinical benefits of dapagliflozin relative to placebo were consistent across a wide range of diuretic categories and doses with a similar safety profile. Treatment with dapagliflozin significantly reduced new loop diuretic requirement over time.

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Structured Graphical Abstract

Key Question

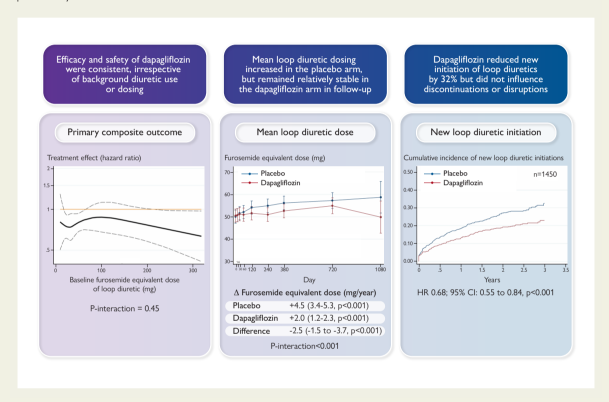
What is the safety and efficacy of the sodium–glucose cotransporter-2 inhibitor dapagliflozin according to background diuretic therapy and how does dapagliflozin influence longitudinal diuretic use in patients with heart failure?

Key Finding

In the DELIVER trial, treatment with dapagliflozin exhibited consistent safety and efficacy irrespective of background diuretic use or dose, and significantly reduced new loop diuretic requirement over time.

Take Home Message

Dapagliflozin exhibits a consistent treatment effect and safety profile irrespective of background diuretic therapy. Reduction in diuretic requirement over time represents an additional therapeutic benefit of dapagliflozin in patients with heart failure with mildly reduced or preserved ejection fraction.



The clinical benefits of dapagliflozin vs. placebo exhibit consistent safety and efficacy across a wide range of background diuretic use. Treatment with dapagliflozin reduces loop diuretic requirement over time. CI, confidence interval; HR, hazard ratio.

Keywords

heart failure with mildly reduced ejection fraction • heart failure with preserved ejection fraction • diuretics • SGLT2i

Introduction

Sodium–glucose co-transporter 2 (SGLT2) inhibitors now represent an important therapeutic pillar in the management of patients with chronic heart failure across the spectrum of left ventricular ejection fraction (LVEF), 1–3 demonstrating a significant and sustained reduction in cardiovascular death or worsening heart failure events. 4,5 Among other effects, SGLT2 inhibitors promote early natriuresis and diuresis, which may contribute to their clinical benefits in heart failure. Diuretic therapy is a cornerstone of the management of patients with heart failure, and their concomitant use with SGLT2 inhibitors will be a frequent occurrence. Understanding the interplay between SGLT2 inhibitors and conventional diuretics is therefore of great importance.

While loop diuretics are an essential facet of management of congestion in patients with heart failure, prolonged use, especially at higher doses, may result in neurohormonal activation, electrolyte abnormalities, and kidney dysfunction. The use of loop diuretics has been associated with worse clinical outcomes in heart failure in a dose-dependent manner. However, whether loop diuretic use is a mediator of risk or is merely a marker of more advanced disease remains uncertain. SGLT2 inhibitors may enhance diuretic efficiency, with a considerably more favourable neurohormonal, electrolyte, and safety profile than conventional diuretics. Whether SGLT2 inhibitors may translate to long-term diuretic sparing is thus important to characterize in the context of heart failure with mildly reduced or preserved ejection fraction. In this pre-specified analysis of the Dapagliflozin Evaluation to Improve the LIVEs of Patients With

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Randomized to dappgillocum 338 (493) 390 (507) 912 (604) Mean furozemide equivalent dose (mg) 704±10.3 715±9.2 718±9.5 Men 377 (55.2) 391 (50.8) 1162 (64.2) Men 377 (55.2) 391 (50.8) 1162 (64.2) Asian 377 (55.2) 532 (80.9) 1162 (64.2) Asian 377 (55.2) 532 (80.9) 1162 (64.2) Asian 443 119 (50.8) 1162 (64.2) Asian 47 (3.2) 24 (3.1) 48 (2.7) Annerican 39 (57.7) 24 (3.1) 48 (2.7) Other 26 (3.8) 38 (4.9) 18 (1.0) Region 114 (27.2) 24 (3.1) 18 (4.1) Orber 12 (3.8) 135 (4.2) 14 (3.1) North American 50 (3.3) 14 (2.7) 24 (3.1) North American 50 (3.3) 14 (6.2) 24 (1.3) Med al biotistorifuter 50 (3.3) 14 (2.25) 24 (3.1) North American 50 (3.3) 14 (6.2) 24 (3.1)		No diuretic $(n = 683)$	Non-loop diuretic $(n = 769)$	<40 mg ^{a,b} ($n = 1811$)	40 mg^{b} $(n = 1902)$	$>40 \text{ mg}^{\text{b}}$ (n = 1098)	P-value ^c
rrosemide equivalent dose (mg) ars) ars) ars) ars) ars) b ars) b ars) ars) ars) ars, ars,	Randomized to dapagliflozin	338 (49.5)	390 (50.7)	912 (50.4)	939 (49.4)	552 (50.3)	96:0
ars) 704±103 715±92 718±95 718±95 718±95 718±95 717552) 391 (508) 1061 (586)	Mean furosemide equivalent dose (mg)			17±6	40±3	116 ± 95	
e 377 (55.2) 391 (50.8) 1061 (88.6) e 377 (55.2) 5.23 (68.0) 1162 (44.2) or African 9 (13) 25 (3.3) 24 (1.3) nencican 9 (13) 25 (3.3) 24 (1.3) re African 99 (57) 24 (3.1) 48 (2.7) re African 26 (3.8) 38 (4.9) 18 (10) re African 26 (3.8) 352 (45.9) 48 (2.7) pe and Saudi Arabia 249 (36.5) 352 (45.9) 48 (2.7) America 154 (22.5) 244 (2.0) 542 (28.9) America 50 (7.3) 48 (6.2) 240 (13.3) Anterica 50 (7.3) 48 (6.2) 240 (13.3) America 50 (7.3) 48 (6.2) 240 (13.3) Amilians <	Age (years)	70.4 ± 10.3	71.5 ± 9.2	71.8 ± 9.5	72.2 ± 9.4	71.5 ± 9.7	<0.001
e 222 (340) 155 (800) 1162 (642) or African 9 (13) 25 (33) 24 (13) nercian 39 (57) 24 (31) 24 (13) reflaska Native 39 (57) 24 (31) 48 (27) reflaska Native 26 (38) 395 (459) 18 (10) pe and Saudi Arabia 249 (36.5) 353 (45.9) 864 (47.7) America 249 (36.5) 249 (36.5) 240 (13.3) hi America 260 (33.7) 154 (20.0) 542 (29.9) Anterica 50 (7.3) 46 (6.2) 240 (13.3) stroke 61 (8.9) 61 (7.9) 180 (9.9) ates Mellitus 61 (8.9) 61 (7.9) 180 (9.9) ates Mellitus 61 (8.9) 274 (25.0) 216 (31.6) mass index (kg/m²) 27.9 ± 5.2 29.2 ± 5.4 28.7 ± 5.8 g status 67 (9.8) 644 (8.3) 138 (7.6) reflect 211 (30.9) 464 (6.3) 1026 (56.7) class	Men	377 (55.2)	391 (50.8)	1061 (58.6)	1035 (54.4)	652 (59.4)	<0.001
e 522 (340) 159 (620) 1162 (642) 190 (72) (72) (72) (72) (72) (72) (72) (72)	Race						<0.001
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or African or African or African or African or African or African or Adaska Native 26 (38) r Alaska Native 26 (38) r B (4.9) 18 (1.0) 18 (1.0) 19 (1.1) 19 (1.1) 19 (1.1) 19 (1.1) 19 (1.2) 19 (1.2) 19 (1.2) 19 (1.2) 19 (1.2) 19 (1.2) 19 (1.3) 10 (1.3)	Asian	232 (34.0)	159 (20.7)	559 (30.9)	222 (11.7)	102 (9.3)	
re Alaska Native 39 (5.7) 24 (3.1) 48 (2.7) re Alaska Native 26 (3.8) 38 (4.9) 18 (1.0) re Alaska Native 26 (3.8) 38 (4.9) 18 (1.0) 18 (1.	Black or African American	9 (1.3)	25 (3.3)	24 (1.3)	57 (3.0)	44 (4.0)	
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pe and Saucil Arabia 249 (36.5) 353 (45.9) 864 (47.7) America 230 (33.7) 154 (20.0) 542 (29.9) America 154 (22.5) 214 (27.8) 165 (9.1) America 50 (7.3) 48 (6.2) 240 (13.3) history 333 (47.3) 315 (41.0) 1043 (57.6) stroke 61 (8.9) 61 (7.9) 180 (9.9) stroke 61 (8.9) 61 (7.9) 180 (9.9) retes Mellitus 269 (39.4) 336 (43.7) 738 (40.8) MI 218 (31.9) 218 (31.9) 218 (31.9) retes Mellitus 269 (39.4) 336 (43.7) 738 (40.8) MI 218 (31.9) 218 (31.9) 218 (31.9) HFH 216 (31.6) 217 (35.0) 218 (31.6) mass index (kg/m²) 27.9 ± 5.2 29.2 ± 5.4 28.7 ± 5.8 ent 67 (9.8) 64 (8.3) 138 (7.6) er 211 (30.9) 241 (31.3) 647 (35.7) dass 464 (60.3) 1026 (56.7)	Other	26 (3.8)	38 (4.9)	18 (1.0)	104 (5.5)	16 (1.5)	
and Arabia 249 (36.5) 353 (45.9) 864 (47.7) arica 230 (33.7) 154 (20.0) 542 (29.9) arica 154 (22.5) 214 (27.8) 165 (9.1) arica 50 (7.3) 48 (6.2) 240 (13.3) ary lation/flutter 323 (47.3) 315 (41.0) 1043 (57.6) ke 61 (8.9) 61 (7.9) 180 (9.9) Aellitus 269 (39.4) 336 (43.7) 738 (40.8) sion 591 (86.5) 982 (89.4) 591 (86.5) sion 591 (86.5) 27.9 ± 5.2 29.2 ± 5.4 28.7 ± 5.8 aridex (kg/m²) 27.9 ± 5.2 29.2 ± 5.4 28.7 ± 5.8 aridex (kg/m²) 241 (31.3) 641 (31.3) 647 (35.7) 405 (59.3) 464 (60.3) 1026 (56.7)	Region						<0.001
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rerica 154 (22.5) 214 (27.8) 165 (9.1) rerica 50 (7.3) 48 (6.2) 240 (13.3) Dry Suzy Re ed (18.9) 61 (7.9) 1043 (57.6) Rellitus 269 (39.4) 336 (43.7) 738 (40.8) Sion 591 (86.5) 982 (89.4) 591 (86.5) sion 591 (86.5) 615 (56.0) 216 (31.6) si ndex (kg/m²) 27.9 ±5.2 29.2 ±5.4 28.7 ±5.8 Cus 67 (98) 64 (8.3) 138 (75.7) 405 (59.3) 464 (60.3) 1026 (55.7)	Asia	230 (33.7)	154 (20.0)	542 (29.9)	210 (11.0)	90 (8.2)	
rerica 50 (7.3) 48 (6.2) 240 (13.3) pry llation/flutter 323 (47.3) 315 (41.0) 1043 (57.6) ke 61 (8.9) 61 (7.9) 180 (9.9) wellituss 269 (39.4) 336 (43.7) 738 (40.8) sion 591 (86.5) 982 (89.4) 591 (86.5) sion 591 (86.5) 615 (56.0) 216 (31.6) si index (kg/m²) 27.9 ± 5.2 29.2 ± 5.4 28.7 ± 5.8 cus 67 (9.8) 64 (8.3) 138 (7.6) 211 (30.9) 241 (31.3) 647 (35.7) 405 (59.3) 464 (60.3) 1026 (56.7)	Latin America	154 (22.5)	214 (27.8)	165 (9.1)	519 (27.3)	129 (11.7)	
hlation/flutter 323 (47.3) 315 (41.0) 1043 (57.6) (e1 (8.9) 61 (7.9) 180 (9.9) (e1 (8.9) 269 (39.4) 336 (43.7) 738 (40.8) (180 (9.9) 218 (31.9) 274 (25.0) 218 (31.9) 216 (31.6) 216 (31.6) 216 (31.6) 216 (31.6) 216 (31.6) 216 (31.6) 217.9 ± 5.2 29.2 ± 5.4 28.7 ± 5.8 211 (30.9) 211 (30.9	North America	50 (7.3)	48 (6.2)	240 (13.3)	244 (12.8)	269 (24.5)	
llation/flutter 323 (47.3) 315 (41.0) 1043 (57.6) ke 61 (8.9) 61 (7.9) 180 (9.9) 180 (9.9) 180 (9.9) 269 (39.4) 336 (43.7) 738 (40.8) 218 (31.9) 274 (25.0) 218 (31.9) 274 (25.0) 218 (31.9) 218 (31.9) 274 (25.0) 218 (31.9) 216 (31.6) 216 (31.6) 216 (31.6) 217.9 ± 5.2 29.2 ± 5.4 28.7 ± 5.8 211 (30.9) 211 (30.9) 241 (31.3) 647 (35.7) 405 (59.3) 464 (60.3) 1026 (56.7)	Medical history						
ke 61 (89) 61 (7.9) 180 (9.9) Mellitus 269 (39.4) 336 (43.7) 738 (40.8) 218 (31.9) 274 (25.0) 218 (31.9) sion 591 (86.5) 982 (89.4) 591 (86.5) 4 216 (31.6) 615 (56.0) 216 (31.6) s index (kg/m²) 27.9 ± 5.2 29.2 ± 5.4 28.7 ± 5.8 cus 67 (9.8) 64 (8.3) 138 (7.6) 211 (30.9) 241 (31.3) 647 (35.7) 405 (59.3) 464 (60.3) 1026 (56.7)	Atrial fibrillation/flutter	323 (47.3)	315 (41.0)	1043 (57.6)	1121 (58.9)	750 (68.3)	<0.001
Mellitus 269 (39.4) 336 (43.7) 738 (40.8) 218 (31.9) 218 (31.9) 274 (25.0) 218 (31.9) 218 (31.9) 218 (31.9) 218 (31.9) 218 (31.9) 216 (31.6) 216 (31.6) 216 (31.6) 21.9 ± 5.2 29.2 ± 5.4 28.7 ± 5.8 20.1 ± 5.8 29.2 ± 5.8 29.2 ± 5.8 29.2 ± 5.8 28.7 ± 5.8 29.2 ± 5.8 29.3 29.3 29.3 29.3 29.3 29.3 29.3 29.3	Prior stroke	61 (8.9)	61 (7.9)	180 (9.9)	178 (9.4)	117 (10.7)	0.33
sion 591 (8.5) $274 (25.0)$ $218 (31.9)$ $274 (25.0)$ $218 (31.9)$ $216 (31.6)$ $216 (31.6)$ $216 (31.6)$ $216 (31.6)$ 27.9 ± 5.2 29.2 ± 5.4 28.7 ± 5.8 27.9 ± 5.2 29.2 ± 5.4 28.7 ± 5.8 27.9 ± 5.2 29.2 ± 5.4 28.7 ± 5.8 $211 (30.9)$ $241 (31.3)$ $647 (35.7)$ $405 (59.3)$ $464 (60.3)$ $1026 (56.7)$	Diabetes Mellitus	269 (39.4)	336 (43.7)	738 (40.8)	895 (47.1)	568 (51.7)	<0.001
sion 591 (86.5) 982 (89.4) 591 (86.5) 14 (86.5) 15 (80.4) 15 (80.5) 15 (80.	Prior MI	218 (31.9)	274 (25.0)	218 (31.9)	511 (28.2)	465 (24.4)	<0.001
Here (kg/m^2) 216 (31.6) 615 (56.0) 216 (31.6) 27.9 ± 5.2 29.2 ± 5.4 28.7 ± 5.8 24.0 27.9 ± 5.2 27.9 ± 5.4 28.7 ± 5.8 24.0 24.0 24.0 24.0 24.0 24.0 24.0 24.0	Hypertension	591 (86.5)	982 (89.4)	591 (86.5)	1543 (85.2)	1723 (90.6)	<0.001
a sindex (kg/m^2) 27.9 ± 5.2 29.2 ± 5.4 28.7 ± 5.8 20.2 ± 5.4 28.7 ± 5.8 20.2 ± 5.4 28.7 ± 5.8 20.2 ± 5.4 20.2 ± 5.4 20.2 ± 5.4 20.2 ± 5.4 20.2 ± 5.4 20.2 ± 5.8 20.2 ± 5.4 20.2 ± 5.8 2	Prior HFH	216 (31.6)	615 (56.0)	216 (31.6)	775 (42.8)	797 (41.9)	<0.001
67 (9.8) 64 (8.3) 138 (7.6) 211 (30.9) 241 (31.3) 647 (35.7) 405 (59.3) 464 (60.3) 1026 (56.7)	Body mass index (kg/m²)	27.9 ± 5.2	29.2 ± 5.4	28.7 ± 5.8	30.5 ± 6.2	32.1 ± 6.6	<0.001
67 (9.8) 64 (8.3) 138 (7.6) 211 (30.9) 241 (31.3) 647 (35.7) 405 (59.3) 464 (60.3) 1026 (56.7)	Smoking status						<0.001
211 (30.9) 241 (31.3) 647 (35.7) 405 (59.3) 464 (60.3) 1026 (56.7)	Current	67 (9.8)	64 (8.3)	138 (7.6)	140 (7.4)	75 (6.8)	
405 (59.3) 464 (60.3) 1026 (56.7)	Former	211 (30.9)	241 (31.3)	647 (35.7)	679 (35.7)	483 (44.0)	
NYHA class	Never	405 (59.3)	464 (60.3)	1026 (56.7)	1083 (56.9)	540 (49.2)	
	NYHA class						<0.001
							Continued

Continued	
Table 1	

Table 1 Continued						
	No diuretic $(n = 683)$	Non-loop diuretic $(n = 769)$	<40 mg ^{a,b} (n = 1811)	40 mg ^b $(n = 1902)$	>40 mg ^b (n = 1098)	P-value ^c
=	566 (82.9)	665 (86.5)	1415 (78.1)	1385 (72.8)	682 (62.1)	
≡	116 (17.0)	101 (13.1)	392 (21.6)	513 (27.0)	409 (37.2)	
≥	1 (0.1)	3 (0.4)	3 (0.2)	4 (0.2)	7 (0.6)	
LVEF (%)	2±9	57±9	54±9	54±9	54±9	<0.001
NT-proBNP (ng/L)	875 (579–1503]	790 (497–1252)	993 (616–1665)	1084 (646–1942)	1252 (748–2227)	<0.001
Systolic blood pressure (mmHg)	127 ± 15	132 ± 15	128 ± 15	128 ± 15	126 ± 16	<0.001
Heart rate (bpm)	72 ± 12	71 ± 11	72 ± 12	72 ± 12	72±12	0.14
Creatinine (µmol/L)	95 ± 27	96 ± 29	100 ± 28	104 ± 32	115 ± 34	<0.001
eGFR (mL/min/1.73 m²)	66±19	65 ± 19	63 ± 19	60±19	55±19	<0.001
ACEi	223 (32.7)	249 (32.4)	718 (39.6)	728 (38.3)	377 (34.3)	<0.001
ARB	245 (36.0)	389 (50.6)	586 (32.4)	679 (35.7)	373 (34.0)	<0.001
ARNI	55 (8.1)	11 (1.4)	105 (5.8)	77 (4.0)	53 (4.8)	<0.001
Beta blocker	549 (80.6)	613 (79.7)	1508 (83.3)	1576 (82.9)	931 (84.8)	0.031
MRA	543 (79.7)	160 (20.8)	714 (39.4)	741 (39.0)	509 (46.4)	<0.001

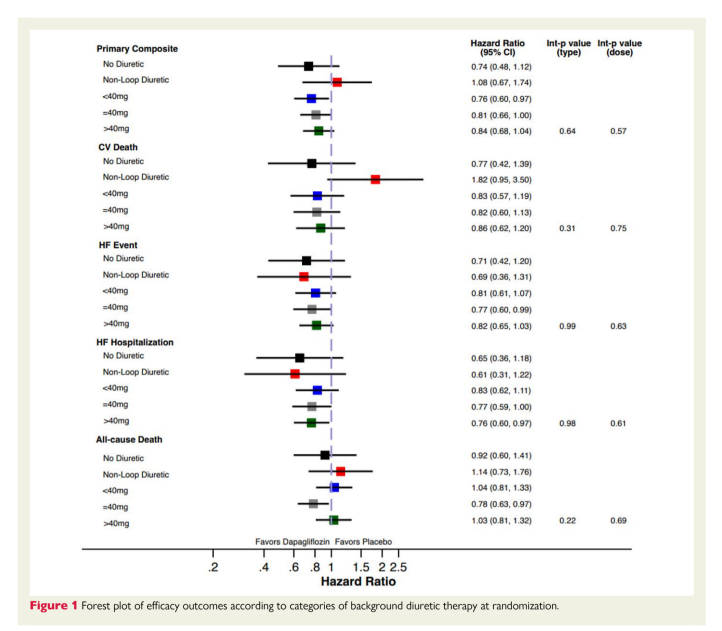
Data are presented as n (%), mean ± SD, or median (interquartile range).

^aA small number of patients (n = 185) on loop diuretic at baseline with missing or inadequate dose information were included in this group.

^bDose in furosemide equivalents.

^cPvalue for the comparison of no-diuretic, non-loop diuretic, and loop diuretic furosemide dose of <40, 40, and >40 mg.

HFH, heart failure hospitalization; NYHA, New York Heart Association; LVEF, left ventricular ejection fraction; ACE; angiotensin-converting enzyme inhibitor; ARB, angiotensin receptor blocker, ARNI, angiotensin receptor-neprilysin inhibitor; NT-proBNP, N-terminal prohormone of B-type natriuretic peptide; MRA, mineralocorticoid receptor antagonist; eGFR, estimated glomerular filtration rate; MI, myocardial infarction.



Preserved Ejection Fraction Heart Failure (DELIVER) trial, we assessed the efficacy and safety of dapagliflozin by background diuretic use and dosing and examined the effect of dapagliflozin on post-randomization changes in diuretic use.

Methods

Study design

The DELIVER trial study design and primary results have been previously reported.³ Briefly, DELIVER was a double-blind, randomized controlled trial which evaluated the effect of dapagliflozin 10 mg once daily vs. placebo among ambulatory or hospitalized patients age ≥40 years with symptomatic heart failure (New York Heart Association class II–IV) and at least intermittent diuretic requirement, LVEF >40%, evidence of structural heart disease, and elevated natriuretic peptides. Background diuretic use and dosing according to local standards of care were maintained throughout the study period. Patients were excluded if they received an SGLT2 inhibitor in the 4 weeks prior to randomization, experienced prior intolerance to SGLT2 inhibitor, had a history of Type 1 diabetes, screening estimated glomerular filtration

rate (eGFR) <25 mL/min/1.73 m 2 , or systolic blood pressure <95 mmHg. The study protocol was approved by the ethics committees at each study site and participants provided written informed consent.

Clinical endpoints

The primary endpoint was a composite of cardiovascular death or worsening heart failure event including hospitalization or urgent visits (requiring intravenous heart failure therapies). Secondary endpoints included total (first and recurrent) heart failure events and cardiovascular death, change in Kansas City Cardiomyopathy Questionnaire (KCCQ) total symptom score at 8 months, cardiovascular death, and all-cause death. The following safety outcomes were also evaluated: any serious adverse event, any adverse event leading to drug discontinuation or interruption, and select other adverse events of interest (including lower limb amputation, volume depletion, and renal adverse events).

Ascertainment of diuretic dose information and diuretic categorization

All patients with information on diuretic treatment were included in this analysis. Only records relating oral administration recorded in 'g', 'µg', 'mg' with

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Primary composite	S ON	No diuretic $(n = 683)$	Non-loc (n:	Non-loop diuretic (n = 769)	4 > = n)	<40 mg ^a (n = 1811)	40 (n =	40 mg ^a (n = 1902)	\	>40 mg ^a (n = 1098)	Pinteraction (diuretic use/tvpe)	Pinteraction (diuretic
	Placebo	Dapagliflozin	Placebo	Dapagliflozin	Placebo	Dapagliflozin	Placebo	Dapagliflozin	Placebo	Dapagliflozin		
Event rate	7.3	5.3	3.9	4.2	8.3	6.4	9.8	7.9	18.1	15.3		
HR (95% CI)	0.74 (C	0.74 (0.48–1.12)	1.08 (C	1.08 (0.67–1.74)	0.76 (0	0.76 (0.60–0.97)	0.81 (0.	0.81 (0.66–1.00)	0.84 ((0.84 (0.68, 1.04)	0.64	0.57
CV death												
Event rate	3.3	2.6	1.6	2.8	3.2	2.6	4.0	3.3	6.3	5.5		
HR (95% CI)	0.77 (0	0.77 (0.42–1.39)	1.82 (0	1.82 (0.95–3.50)	0.83 (0	0.83 (0.57–1.19)	0.82 (0.	0.82 (0.60–1.13)	HR = 0.8	HR = 0.86 (0.62, 1.20)	0.31	0.75
HF event												
Event rate	4.7	3.4	2.7	1.9	5.9	8.4	6.9	5.3	15.4	12.6		
HR (95% CI)	0.71 (0	0.71 (0.42–1.20)	0) 69:0	0.69 (0.36–1.31)	0.81 (0	0.81 (0.61–1.07)	0.77 (0.	0.77 (0.60–0.99)	0.82 ((0.82 (0.65, 1.03)	0.99	0.63
Ŧ												
Event rate	3.8	2.5	2.5	1.5	5.2	4.3	6.3	4.8	14.7	11.2		
HR (95% CI)	0) 59:0	0.65 (0.36–1.18)	0.61 (0	0.61 (0.31–1.22)	0.83 (0	0.83 (0.62–1.11)	0.77 (0.	0.77 (0.59–1.00)	0.76 ((0.76 (0.60, 0.97)	0.61	0.91
All-cause death												
Event rate	5.8	5.4	4.3	4.9	6.2	6.4	9.2	7.2	10.7	11.1		
HR (95% CI)	0.92 (0	0.92 (0.60–1.41)	1.14 (0	1.14 (0.73–1.76)	1.04 (0	1.04 (0.81–1.33)	0.78 (0.	0.78 (0.63–0.97)	1.03 ((1.03 (0.81, 1.32)	0.22	69:0
KCCQ-TSS 1 month	78.7 ± 20.2	77.5 ± 19.6	80.2 ± 17.9	80.4 ± 18.1	77.9 ± 19.0	80.5 ± 16.9	74.4 ± 19.7	75.2 ± 20.5	69.3 ± 22.5	71.9 ± 22.4		
Difference	1.7 (-0	1.7 (-0.7 TO 4.1)	1.4 (–(1.4 (-0.6 to 3.5)	2.0 ((2.0 (0.7–3.3)	0.6 (-1	0.6 (-1.0 to 2.2)	4.0 (4.0 (1.7, 6.3)	0.08	0.13
KCCQ-TSS 8 months	80.1 ± 18.8	82.0 ± 17.4	79.6 ± 19.7	81.5 ± 17.6	78.8 ± 19.7	81.6 ± 16.8	75.0 ± 21.3	77.1 ± 20.0	72.5 ± 21.8	74.7 ± 21.1		
Difference	3.5 (3.5 (0.9–6.2)	3.5 (1	3.5 (0.9–6.1)	2.0 ((2.0 (0.4–3.6)	1.7 (-0	1.7 (-0.3 to 3.7)	2.9 (2.9 (0.1, 5.7)	0.88	0.78

^aDose in furosemide equivalents.
CI, confidence interval; CV, cardiovascular; HF, heart failure, HFH, heart failure hospitalization; HR, hazard ratio; KCCQ-TSS, Kansas City Cardiomyopathy Questionnaire total symptom score.

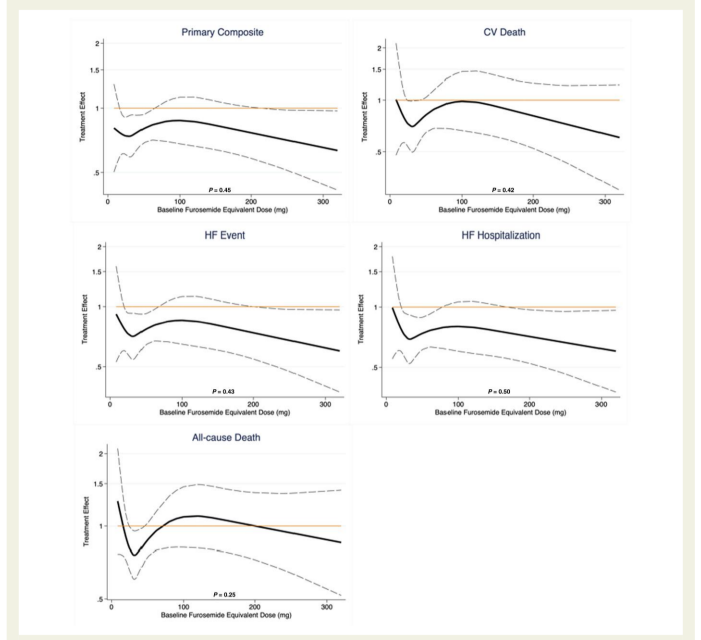


Figure 2 Treatment effect by baseline furosemide equivalent diuretic dose. The *P*-values displayed represent the *P*_{interaction} of treatment effects of dapagliflozin across a range of furosemide dose equivalents, evaluated as a continuous function. CV, cardiovascular; HF, heart failure.

standard dose frequency were included. Intermittent, as-needed diuretic therapy was not considered. Of the 13 357 diuretic records identified in DELIVER, 3586 records (27%) were excluded leaving 9771 records (73%) for analysis. For all patients on loop diuretic, a total daily furosemide dose equivalent was calculated. Bumetanide 1 mg, torsemide 20 mg, azosemide 60 mg, and ethacrynic acid 100 mg were considered equivalent to 80 mg of oral furosemide. Information on diuretic start and stop dates for any diuretic type or dose was collected throughout the study in an effort to capture between visit changes. In analyses of post-baseline changes in diuretic use, patients were excluded at time points with missing or insufficient diuretic dose data and included again at the next available time point. In the analysis of mean loop diuretic dose over time, only patients on a non-zero dose of loop diuretic were included. Where patients were on combination diuretic therapy, they were considered part of the group of the most potent diuretic in the combination for the purpose of analysis (i.e. analysed in the 'loop diuretic' subgroup

when treated with a combination of loop and thiazide diuretic). Patients treated with a mineralocorticoid receptor antagonist (MRA) alone were included in the 'no diuretic' category. The non-loop diuretic category most frequently included patients on thiazide and thiazide-like diuretics.

Statistical analysis

Baseline characteristics were compared according to the following categories defining baseline diuretic use: no diuretic, non-loop diuretic, and furosemide dose equivalents of >40, 40, and <40 mg, respectively. Data are reported as mean \pm standard deviation, median (interquartile range) for non-normal distributions, and frequency (percentage) for categorical variables. Student's *t*-test, Pearson χ^2 test, and ANOVA were used where appropriate.

The treatment effect of dapagliflozin on the primary composite outcome, its components, and all-cause death were evaluated across diuretic categories

				•								
Safety outcome	ž	No diuretic $(n = 683)$	Non-loo (n =	-loop diuretic $(n = 769)$	ν̈́ ε	<40 mg ^a (n = 1811)	4 u)	40 mg ^a $(n = 1902)$	λs	>40 mg ^a (n = 1098)	P _{interaction} (diuretic use/type)	P _{interaction} (diuretic dose)
	Placebo	Placebo Dapagliflozin	Placebo	Dapagliflozin	Placebo	Dapagliflozin	Placebo	Dapagliflozin	Placebo	Dapagliflozin		
Any SAE	143 (41.6%)	126 (37.4%)	135 (35.8%)	140 (36.0%)	380 (42.3%)	367 (40.3%)	457 (47.6%)	423 (45.1%)	308 (56.4%)	305 (55.4%)	0.84	0.51
Any AE leading to discontinuation of IP	15 (4.4%)	19 (5.6%)	11 (2.9%)	23 (5.9%)	57 (6.3%)	57 (6.3%)	54 (5.6%)	41 (4.4%)	44 (8.1%)	42 (7.6%)	0.65	0.43
Any AE leading to interruption of IP	50 (14.5%)	38 (11.3%)	54 (14.3%)	48 (12.3%)	131 (14.6%)	122 (13.4%)	148 (15.4%)	125 (13.3%)	111 (20.3%)	103 (18.7%)	96:0	0.50
Any AE that potentially placed a patient at increased risk for a lower limb amputation b	14 (4.1%)	(3.3%)	11 (2.9%)	(2.1%)	48 (5.3%)	46 (5.0%)	57 (5.9%)	60 (6.4%)	69 (12.6%)	63 (11.4%)	0.90	0.80
Any SAE or DAE suggestive of volume depletion	3 (0.9%)	(0.3%)	1 (0.3%)	2 (0.5%)	11 (1.2%)	13 (1.4%)	10 (1.0%)	11 (1.2%)	7 (1.3%)	15 (2.7%)	0.50	0.14
Any renal SAE or DAE	6 (1.7%)	5 (1.5%)	4 (1.1%)	6 (1.5%)	18 (2.0%)	15 (1.6%)	18 (1.9%)	20 (2.1%)	33 (6.0%)	27 (4.9%)	0.76	0.93

^aDose in furosemide equivalents.

^bPreceeding events were identified based on a predefined list of the European Medicines Agency's Pharmacovigilance Risk Assessment Committee preferred terms.

SAE, serious adverse event; AE, adverse event; IP, investigational product; DAE, adverse events leading to discontinuations.

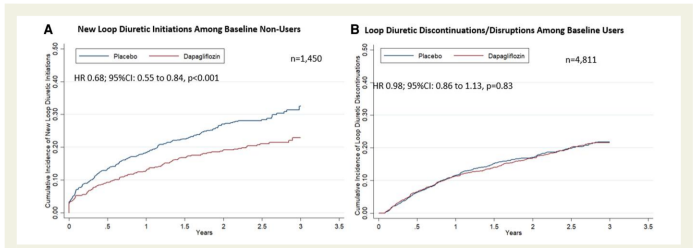


Figure 3 Treatment effect on loop diuretic use in study follow-up. Kaplan–Meier curves displaying the (A) time to loop diuretic initiation among patients not on loop diuretic at baseline and (B) time to discontinuation or disruption among patients being treated with loop diuretic at baseline are presented. Disruption was defined as any interruption in diuretic use of ≥30 days. CI, confidence interval; HR, hazard ratio.

in time-to-first event analyses using Cox regression and Kaplan–Meier curves. Treatment effects on the KCCQ total symptom score were assessed using repeated measures mixed-effect models, adjusted for baseline value, at 1 and 8 months. Total worsening heart failure events and cardiovascular death were analysed using semi-parametric proportional rates methods of Lin et al. Effects on safety outcomes were assessed with Cox regression in an ontreatment analysis. Interaction testing was carried out to evaluate the interaction between treatment and diuretic type as well as loop diuretic dose.

A repeated measures mixed-effect model was employed to assess temporal changes in mean furosemide equivalent dose over trial time among patients on loop diuretics. Treatment, time, and the interaction between assigned treatment and time were included as fixed effects. Cox-proportional hazards models were used to assess the time to new loop diuretic initiation (among baseline loop diuretic non-users) and discontinuation or disruption (among baseline loop diuretic users). A disruption in loop diuretic use was considered as any interruption in diuretic treatment ≥30 days. Results were displayed with Kaplan–Meier curves. The percentage of all patients experiencing a first sustained change (≥30 days) in loop diuretic dose was compared between treatment groups with regression analysis. All analyses were carried out using STATA version 17.0 (Stata Corp., College Station, TX, USA). A *P*-value of <0.05 was considered to be statistically significant.

Results

Baseline diuretic use in DELIVER

In DELIVER, of 6263 patients, 10.9% (n=683) were on no diuretic, 12.3% (n=769) were on a non-loop diuretic, and 76.8% (n=4811) were on a loop diuretic at the time of randomization (*Table 1*). Among those on loop diuretics at baseline, 37.7% (n=1811) were receiving a furosemide equivalent dose of <40 mg, 39.5% (n=1902) were on a dose equivalent 40 mg, and 22.8% (n=1098) were on a dose equivalent >40 mg (*Table 1*).

Patient profiles

Patients on the highest furosemide equivalent doses (>40 mg) at base-line were more often men, with a higher burden of comorbidities, higher body mass index, more severe heart failure symptoms, and more frequently had a prior history of heart failure hospitalization.

These patients also had lower baseline eGFR and were more likely to be treated with an MRA but less likely to be treated with sacubitril/valsartan (*Table 1*).

Efficacy and safety of dapagliflozin according to baseline diuretic use

The cumulative incidence of the primary composite outcome and its components as well as all-cause death were observed to be lowest in patients on non-loop diuretic or no diuretic and highest in patients on furosemide equivalent doses >40 mg (all P < 0.001; Supplementary data online, Figure S1). These findings were consistent when total daily furosemide equivalent dose was analysed as a continuous variable with increasing risk observed at higher furosemide equivalent doses for all evaluated outcomes (Supplementary data online, Figure S2).

The treatment benefit of dapagliflozin compared to placebo on the primary composite outcome did not significantly vary by baseline diuretic use/type ($P_{\text{interaction}} = 0.64$) or loop diuretic dose ($P_{\text{interaction}} = 0.57$, Figure 1, Table 2). Similarly, treatment effects on the components of the primary composite endpoint, all-cause death, and change in KCCQ total symptom score at 1 and 8 months was consistent across diuretic use categories (Table 2). Moreover, when baseline loop diuretic dose was analysed as a continuous variable, the treatment effect of dapagliflozin remained consistent across a broad range of doses for all evaluated outcomes (Figure 2). The safety profile of dapagliflozin was consistent across diuretic categories with similar risk of drug discontinuation or interruption due to adverse events between treatment groups. The occurrence of any serious adverse events or study drug discontinuation due to an adverse event suggestive of volume depletion or renal events were also similar between treatment groups, irrespective of diuretic use/type or dose (Table 3).

Effects of dapagliflozin on diuretic use over time

Among the 1450 patients not treated with a loop diuretic at baseline, new initiations occurred in 346 patients. Dapagliflozin reduced new initiations of loop diuretics by 32% [hazard ratio (HR) 0.68; 95% confidence interval (CI): 0.55–0.84, P < 0.001] (Figure 3). Among the 4811

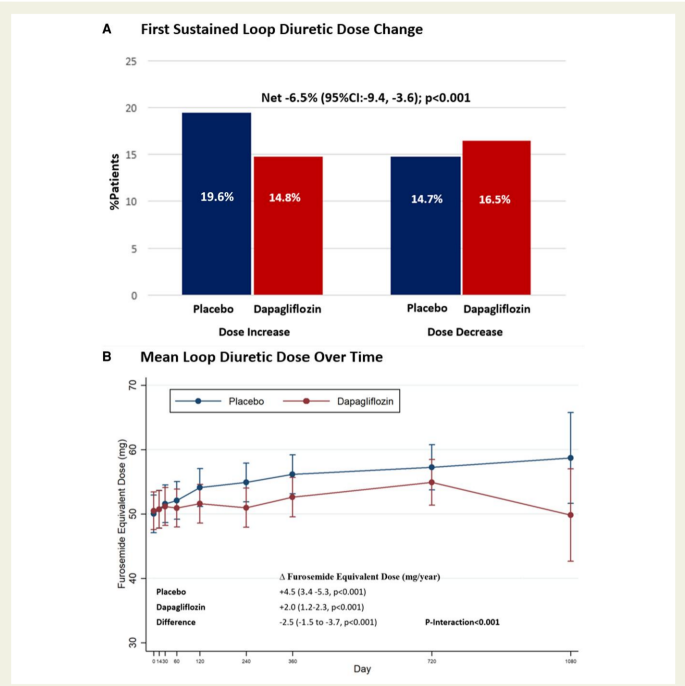


Figure 4 Effect of dapagliflozin on (A) first sustained loop diuretic dose change and (B) mean loop diuretic dose over time. The interaction between mean diuretic dosing, treatment allocation, and time was assessed.

participants on baseline loop diuretic, there was no significant difference in new loop diuretic discontinuations or disruptions (HR 0.98; 95% CI: 0.86-1.13, P=0.83) in follow-up.

The mean baseline furosemide equivalent dose, calculated among all patients on loop diuretics, was similar between treatment groups: 51 ± 59 mg in the dapagliflozin arm and 50 ± 61 mg in the placebo arm. Patients randomized to dapagliflozin compared to placebo less frequently experienced a loop diuretic initiation or dose increase (14.8% vs. 19.6%, P < 0.001) and more frequently experienced a loop diuretic discontinuation or dose decrease (14.7% vs. 16.5%, P < 0.001), with a significant net reduction with dapagliflozin of -6.5% (95% CI: -9.4,

-3.6, P < 0.001) (Figure 4). These results were consistent when considering dosing changes only among baseline loop diuretic users (Supplementary data online, Table S1). In follow-up, mean loop diuretic dose increased at a rate of 4.5 mg/year (95% CI: 3.4–5.3, P < 0.001) in the placebo group and a rate of 2.0 mg/year (95% CI: 1.2–2.3, P < 0.001) in the dapagliflozin group. Treatment with dapagliflozin significantly attenuated the rate of rise in loop diuretic dose relative to placebo resulting in a mean dose reduction over time of 2.5 mg/year (95% CI: -1.5, -3.7, P < 0.001). The difference in loop diuretic dose requirement between treatment groups emerged after day 60 and increased over time ($P_{\rm interaction} < 0.001$) (Figure 4).

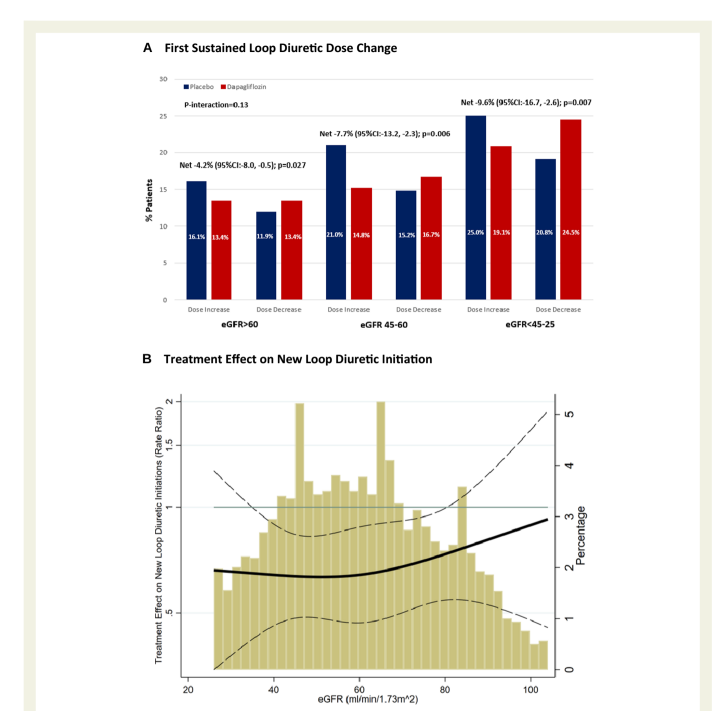


Figure 5 Effect of dapagliflozin on (A) first sustained loop diuretic dose change and (B) new loop diuretic initiation according to baseline kidney function.

Effect of dapagliflozin on diuretic use according to baseline kidney function

There was a significant net reduction in loop diuretic dose with dapagliflozin irrespective of baseline kidney function [eGFR >60 mL/min/ $1.73 \, \text{m}^2$: -4.2% (95% CI: -8.0, -0.5, P=0.027); eGFR 45–60 mL/min/ $1.73 \, \text{m}^2$: -7.7% (95% CI: -13.2, -2.3, P=0.006); eGFR <45– $25 \, \text{mL/min}/1.73 \, \text{m}^2$: -9.6% [-16.7, -2.6, P=0.007], $P_{\text{interaction}}=0.13$] (Figure 5A). When eGFR was analysed as a continuous variable, relative to placebo, dapagliflozin consistently reduced new

loop diuretic initiations across the spectrum of baseline kidney function (Figure 5B).

Discussion

In a pre-specified analysis of the DELIVER trial of patients with heart failure with mildly reduced or preserved ejection fraction, treatment with dapagliflozin compared to placebo resulted in consistent clinical benefits and tolerability across a broad range of diuretic use categories

and doses. Dapagliflozin significantly reduced new loop diuretic initiations, dose increases, and mean loop diuretic dose over time, compared to placebo (*Structured Graphical Abstract*).

Participants in DELIVER with the greatest diuretic requirement expectedly experienced the highest clinical risk across all evaluated outcomes, a finding that is well established in both patients with heart failure with preserved ejection fraction (HFpEF)^{10,11} and heart failure with reduced ejection fraction (HFrEF). 15,16 Nevertheless, the treatment effect of dapagliflozin vs. placebo remained consistent irrespective of background diuretic therapy. These observations are consistent with data on the use of SGLT2 inhibitors in patients with HFrEF in the DAPA-HF trial, where dapagliflozin resulted in similar improvement in clinical outcomes across diuretic subgroups. 13 Similarly, in the EMPEROR-Reduced trial, the benefits of empagliflozin did not differ among patients with and without recent volume overload. ¹⁷ Taken together, these data from DELIVER add to the growing body of literature suggesting the ease of use and practical implementation of SGLT2 inhibitors among patients with a broad spectrum of diuretic requirements, including among those not receiving a diuretic.

In DELIVER, dapagliflozin significantly reduced mean loop diuretic requirement over time in a population of well characterized patients with heart failure with mildly reduced or preserved ejection fraction. Similarly, other SGLT2i including empagliflozin have also been shown to reduce the need for diuretic intensification. In contrast, among patients with HFrEF in DAPA-HF, this longitudinal effect on diuretic requirement was less apparent. Notably, a higher proportion of patients were on background diuretic therapy in DELIVER compared to DAPA-HF (89% vs. 81%). Unlike DAPA-HF, DELIVER allowed enrolment of stabilized (no longer requiring intravenous diuretics) patients actively or recently hospitalized for heart failure, which may suggest greater degree of baseline congestion in the DELIVER population. The potential diuretic effect of SGLT2 inhibitors has been postulated to be more marked in the presence of greater volume overload. 18,19

Patients with worse kidney function incur greater levels of diuretic resistance and therefore require higher rates and doses of diuretics. Importantly, in this analysis, the diuretic sparing effects of dapagliflozin were consistent across the spectrum of baseline kidney function. These data suggest that such agents may reduce diuretic requirements over time, even in patients with significant kidney disease.

The reduction in mean loop diuretic dose with dapagliflozin appears to be driven by a reduction in new loop diuretic initiations and less need for sustained diuretic dose increase. Interestingly, rates of loop diuretic discontinuation or temporary interruption did not differ significantly between treatment groups. The effects of dapagliflozin to prevent diuretic intensification to a greater degree than effects on diuretic dose reduction or discontinuation is similar to previous observations from adjacent trials. The need for diuretic intensification in the ambulatory or inpatient setting represents an element of worsening heart failure and carries prognostic significance. Given that the reduced need for diuretic up-titration occurred in the context of an overall reduction in loop diuretic requirement that emerged late (day 60) and increased over time, this may reflect dapagliflozin's combined effect to prevent the progression of heart failure beyond initial decongestion.

Other contributing explanations for these observations should be noted. It is possible that the lack of impact on diuretic discontinuation may reflect clinical inertia to make alterations to diuretic regimens; thereby, resulting in fewer discontinuations even where volume status may have improved.²¹ The more favourable neurohormonal, renal, and

electrolyte profile of these drugs may also result in fewer clinical triggers for diuretic modification with improved volume status.¹²

In DELIVER, we observed a similar safety profile of dapagliflozin compared to placebo across a range of diuretic doses. Notably, no between-arm differences in the occurrence of renal adverse events or those related to volume depletion were detected even at the highest doses of diuretic. In a similar analysis of dapagliflozin in patients with HFrEF, ¹³ volume depletion was slightly more frequently observed in dapagliflozin treated patients on higher doses of diuretic. Traditionally, the concern for volume depletion has been of greater consequence in patients with HFpEF given the tendency to greater preload dependence in the setting of diastolic dysfunction. ²² In this light, the results of the present study are reassuring. Moreover, rates of study drug discontinuation were relatively low and did not differ between arms, highlighting the tolerability of the concomitant use of SGLT2 inhibitors and conventional diuretics.

Certain important limitations of this analysis should be noted. First, missing or inadequate dose information precluded analysis of all patients at certain time points. Second, specific clinical rationale motivating modifications to diuretic regimens was not available and may reflect issues other than volume status alone. Third, while information on diuretic regimens and start and stop dates were collected at study visits, these data were not cross-referenced against pharmacy claims data or other objective sources. Finally, DELIVER did not collect measures of diuresis and natriuresis such as urinary volumes or urinary electrolyte profiles.

In conclusion, in this pre-specified analysis of DELIVER in patients with heart failure and mildly reduced or preserved ejection fraction, dapagliflozin safely improved clinical outcomes across a wide range of diuretic use and doses. Treatment with dapagliflozin significantly reduced loop diuretic requirement over time, which appears to be driven by lower needs for diuretic intensification and underscores dapagliflozin's effect to prevent worsening heart failure. However, there were limited differences in the necessity for loop diuretic discontinuation or dose reduction between treatment arms. As such, these data may argue against anticipatory loop diuretic dose reduction at the time of SGLT2 inhibitor initiation.

Supplementary data

Supplementary data is available at European Heart Journal online.

Pre-registered clinical trial number

The pre-registered clinical trial number is NCT03619213.

Ethical approval

The study protocol was approved by the ethics committees at each study site and participants provided written informed consent.

Data availability

The trial sponsor is committed to Responsible Data Sharing Principles, including sharing of anonymized individual patient-level data and clinical documents related to DELIVER with qualified researchers. The trial data availability is according to the criteria and processes described at https://astrazenecagrouptrials.pharmacm.com/ST/Submission/Disclosure.

Conflict of interest

S.C is supported by the Canadian Arthur J.E. Child's Cardiology Fellowship. M.V. has received research grant support, served on advisory boards, or had speaker engagements with American Regent, Amgen, AstraZeneca, Bayer AG, Baxter Healthcare, Boehringer Ingelheim, Chiesi, Cytokinetics, Pharmaceuticals, Merck, Novartis, Novo Pharmacosmos, Relypsa, Roche Diagnostics, Sanofi, and Tricog Health and participates on clinical trial committees for studies sponsored by AstraZeneca, Galmed, Novartis, Bayer AG, Occlutech, and Impulse Dynamics. B.C. has been a consultant for Amgen, AO Biome, Biogen, Boehringer Ingelheim, Corvia, Gilead, Myokardia, Cardurion, and Novartis. O.V. has received either personal or institutional research support for DELIVER from AstraZeneca. A.S.D. has received research grant support from AstraZeneca, Alnylam, and Novartis; and has received consulting fees and/or honoraria from Abbott, AstraZeneca, Alnylam, Boehringer-Ingelheim, Boston Scientific, Biofourmis, Corvidia, DalCor Pharma, Novartis, Relypsa, and Regeneron. P.S.J. has received consulting fees, advisory board fees, and lecture fees from Novartis, advisory board fees from Cytokinetics, and grant support from Boehringer Ingelheim. R.A.D.'s institution, the UMCG, has received research grants and fees (outside the submitted work) from AstraZeneca, Abbott, Boehringer Ingelheim, Cardio Pharmaceuticals Gmbh, Ionis Pharmaceuticals, Inc, Novo Nordisk, and Roche. R.A.d.B. has received speaker fees from Abbott, AstraZeneca, Bayer, Novartis, and Roche (outside the submitted work). C.S.P.L. is supported by a Clinician Scientist Award from the National Medical Research Council of Singapore; has received research support from AstraZeneca, Bayer, Boston Scientific, and Roche Diagnostics; has served as a consultant or on the advisory board/steering committee/executive committee for Actelion, Amgen, Applied Therapeutics, AstraZeneca, Bayer, Boehringer Ingelheim, Boston Scientific, Cytokinetics, Darma Inc, Us2.ai, Janssen Research & Development LLC, Medscape, Merck, Novartis, Novo Nordisk, Radcliffe Group Ltd, Roche Diagnostics, Sanofi, and WebMD Global LLC; and serves as the cofounder and non-executive director of Us2.ai. M.N.K. has received research grant support from AstraZeneca, and Boehringer Ingelheim; has served as a consultant or on an advisory board for Amgen, Applied Therapeutics, AstraZeneca, Bayer, Boehringer Ingelheim, Eli Lilly, Esperion Therapeutics, Janssen, Merck (Diabetes and Cardiovascular), Novo Nordisk, Sanofi, and Vifor Pharma; has received other research support from AstraZeneca; and has received honorarium from AstraZeneca, Boehringer Ingelheim, and Novo Nordisk. S.J.S. has received either personal or institutional research support for DELIVER from AstraZeneca. F.M. has received personal fees from AstraZeneca. S.E.I. has served on clinical trial committees or as a consultant to AstraZeneca, Boehringer Ingelheim, Novo Nordisk, Merck, Pfizer and has given lectures sponsored by AstraZeneca and Boehringer Ingelheim. A.F.H. has received research grant support from American Regent, Amgen, AstraZeneca, Bayer, Boehringer Ingelheim, Merck, Novartis, Somo-logic, and Verily and has served as a consultant or on the Advisory Board for Amgen, AstraZeneca, Bayer, Boehringer Ingelheim, Boston Scientific, Bristol Myers Squibb, Cytokinetics, Eidos, Intercept, Merck, and Novartis. T.H. reports no disclosures. S.S.M. served on the advisory boards of BridgeBio and Alnylam Pharmaceuticals and also receives speaking fees from Alnylam Pharmaceuticals. Z.M.M. reports no disclosures. M.P. is an employee and shareholder of AstraZeneca. A.M.L. is an employee and shareholder of AstraZeneca. J.J.V.M. has received grants and his employer paid by AstraZeneca, Theracos, and GlaxoSmithKline during the conduct of the study and grants and his employer being paid by Novartis, Amgen, Bristol-Myers Squibb, Bayer, Abb-vie, Dal-Cor, Kidney Research UK, and Cardurion and grants from British Heart Foundation. S.D.S. has received research grants from Actelion, Alnylam, Amgen, AstraZeneca, Bellerophon, Bayer, BMS, Celladon, Cytokinetics, Eidos, Gilead, GSK, Ionis, Lilly, Mesoblast, MyoKardia, NIH/NHLBI, Neurotronik, Novartis, NovoNordisk, Respicardia, Sanofi Pasteur, Theracos, US2.Al and has consulted for Abbott, Action, Akros, Alnylam, Amgen, Arena, AstraZeneca, Bayer,

Boeringer-Ingelheim, BMS, Cardior, Cardurion, Corvia, Cytokinetics, Daiichi-Sankyo, GSK, Lilly, Merck, Myokardia, Novartis, Roche, Theracos, Quantum Genomics, Cardurion, Janssen, Cardiac Dimensions, Tenaya, Sanofi-Pasteur, Dinaqor, Tremeau, CellProThera, Moderna, American Regent, Sarepta, Lexicon, Anacardio, Akros, and Puretech Health. The remaining authors have nothing to disclose.

Funding

The DELIVER trial was funded by AstraZeneca.

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