## SYNOPSIS

Name of Sponsor/Company:		
Astellas Pharma Europe B.V		
Name of Finished Product:		
Name of Active Ingredient:		
Title of Study: A randomized double h	lind parallel group pla	asha and active controlled multi
center dose ranging study with the beta-	3 agonist VM178 in pat	ients with symptomatic overactive
bladder (DRAGON)	o agoinst 1 wi170 in pat	ients with symptomatic overactive
Coordinating Investigator:	BSc MD FRCS (U	rol) FEBU
	U.K.	,
Study Centers: In 14 countries, a total of	of 97 centers participate	d in the study.
Publication (reference): Not applicable	at the time of this repo	rt
Study Period:	•	Phase of Development: IIb
Date of First Enrollment: 12 April 200	6	-
Date of Last Evaluation: 26 March 200	)7	
Objectives: The primary objective of the	e study was to evaluate	the dose-response relationship of
YM178 OCAS on efficacy in patients w	ith OAB.	
The secondary objectives of the study w	ere:	
• To evaluate the safety and tolerability	of qd dosing with YM	178 OCAS in patients with OAB.
• To compare the efficacy of YM178 C	CAS with tolterodine 4	mg qd.
• To compare the safety and tolerability	y of YM178 OCAS with	n tolterodine 4 mg qd.
To collect population PK data in patients	ents with OAB.	
<b>Study Design:</b> This was a multinationa	l, multicenter, double-bl	lind, double-dummy, randomized,
parallel group, placebo- and active-conti	olled phase IIb study. P	atients were enrolled into a single-
blind, 2-week placebo run-in period follo	owed by a randomized,	double-blind, placebo-controlled,
12-week treatment period. There were o	VISITS IN TOTAL VISIT 1 at $1.5$ and 6 after 1 $4.8$ at	d 12 weeks of double blind treatment
respectively. At screening (Visit 1) patie	t, 5 allu 0 allel 1, 4, 6 al	on diary which had to be completed
during the 3 days preceding Visit 7 Pati	ents received medication	n for the placebo run-in period. In
addition the nationt received an ABPM	device for measuring n	ulse rate and blood pressure during the
3-day diary period. The patient was inst	ructed on how the measure	urements and documentation in the
diary had to be done. After the placebo r	un-in period, patients re	eturned to the clinic (Visit 2).
Micturition diary scores were checked a	gainst the inclusion crite	eria. Patients who met the inclusion
criteria and did not meet the exclusion c	riteria were randomized	to one of the treatments with YM178
OCAS 25 mg qd, 50 mg qd, 100 mg qd,	200 mg qd, placebo or	tolterodine 4 mg qd. Micturition
diaries had to be completed by the patient	nt during the 3 days pred	ceding Visits 3, 4, 5 and 6. During the
diary period preceding each visit, patien	ts measured vital signs (	(HR, SBP, DBP) 2 times per day in
triplicate by means of ambulatory blood	pressure monitoring. N	ature, frequency and severity of
reported or observed adverse events wer	e recorded at all visits. $\frac{1}{2}$	Symptom and Quality of Life
Questionnaires were completed at Visits	2, 3, 4, 5 and $6$ . Post-ve	bid residual volume was determined at
Visits 1 3 4 5 and 6 and ECGs were n	arformed at Visite 1 2	and 6
Diagnosis and Main Criteria for Inclu	sion: Men or women as	3, 4, and 0.
symptoms of OAB (including urinary fr	sion: Well of Wolliell ag	z = 18 years who had experienced
least 3 months prior to screening	equency, and digency w	ing without urge meontimence) for at
Number of Subjects (nlanned and and	lyzed): A total of 1070	patients were to be enrolled in order
to randomize 856 patients to obtain 770	evaluable patients. A to	tal of 1108 subjects were screened
and 927 were treated.		
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**Test Product, Dose and Mode of Administration:** Patients who met the inclusion criteria and did not meet the exclusion criteria were randomized to 12 weeks of double-blind treatment with YM178 OCAS 25 mg qd, 50 mg qd, 100 mg qd, 200 mg qd, placebo or tolterodine 4 mg qd. Each patient randomized to any group took 3 tablets and 1 capsule each morning after breakfast throughout the study. All treatments were taken orally with a glass of water and swallowed intact. YM178 tablets, tolterodine capsules and the corresponding placebo tablets and capsules were indistinguishable (double-dummy technique).

## Lot Numbers:

**Duration of Study and Treatment:** A single-blind, 2-week placebo run-in period followed by a randomized, double-blind, placebo-controlled, 12-week treatment period.

**Criteria for Evaluation:** The primary efficacy variable was the change from baseline to endpoint in mean number of micturitions per 24 hours.

Secondary efficacy variables were:

- Change from baseline in mean volume voided per micturition
- Change from baseline in mean number of urgency episodes (grade 3 and/or 4) per 24 hours
- Change from baseline in level of urgency
- Change from baseline in mean number of urge incontinence episodes per 24 hours
- Change from baseline in mean number of incontinence episodes per 24 hours
- Change from baseline in mean number of nocturia episodes per 24 hours
- Change from baseline in symptom scores as assessed by International Consultation on Incontinence Questionnaire-Overactive Bladder (ICIQ-OAB)
- Change from baseline in quality of life scores as assessed by International Consultation on Incontinence Questionnaire-Overactive Bladder-Quality of Life (ICIQ-OABqol)
- Change from baseline in patient's perception of treatment benefit.

Safety was assessed by monitoring of incidence and severity of adverse events (AEs), vital signs, laboratory tests, 12-lead-electrocardiogram (ECG), and post-void residual volume (PVR), as measured by ultrasonography or bladder scan.

**Statistical Methods**: The full analysis set (FAS) was the primary population for the efficacy analyses and comprised all patients who were randomized, who took at least 1 dose of double-blind study medication, and who provided primary efficacy data from the diary at baseline and endpoint. The following statistical hypotheses were applied:

 $H_0$ : The change in mean number of micturitions per 24 hours from baseline to endpoint is the same for all doses of YM178 OCAS and for placebo;

 $H_a$ : The change in mean number of micturitions per 24 hours from baseline to endpoint is not the same for all doses of YM178 OCAS and for placebo.

The hypothesis was evaluated using a 2-sided test with a significance level of 0.05.

The primary analysis was performed on the change from baseline (Visit 2) to endpoint in the mean number of micturitions per 24 hours as derived from the micturition diary.

Changes from baseline to endpoint in mean number of micturitions per 24 hours were subjected to a model including YM178 OCAS dose as a fixed factor taking on the values 0 mg (placebo), 25 mg, 50 mg, 100 mg and 200 mg. The tolterodine group was not part of this analysis. The baseline value of mean number of micturitions per 24 hours was included in the model as a covariate. Safety variables were reported descriptively.

## **RESULTS:**

**Analysis Sets and Subject Disposition**: The SAF comprised 927 patients who received at least 1 dose of double-blind study medication. The FAS comprised 919 randomized patients who received at least 1 dose of study medication and for whom primary efficacy data at the baseline visit and at least 1 on-treatment visit were available. The PPS consisted of a subset of 793 FAS patients who complied with the major protocol requirements through all applicable visits, i.e., those patients who had no major protocol violations. A total of 1108 patients were enrolled (screened) in the study of whom 928 were randomized of whom 927 received at least one dose of double-blind study medication.

Of these 927 subjects (SAF population), 169 patients received placebo, 169 patients were treated with YM178 OCAS 25 mg qd, 169 patients were treated with YM178 OCAS 50 mg qd, 168 patients were treated with YM178 OCAS 100 mg qd, 167 patients were treated with YM178 OCAS 200 mg qd, and 85 patients were treated with tolterodine 4 mg qd. Seventy subjects prematurely discontinued and 857 subjects completed the study.

**Demographics:** Patients were predominantly female (89.3%) and Caucasian (98.2%), between 18 and 91 years of age; the mean age was 57.2 years. Approximately half of the patients (42.2%) had urgency with incontinence. The mean time since the start of symptoms was 3.9 years. Overall, 45.5% of patients had had previous drug therapy for symptoms of OAB.

**Study Drug Exposure:** The mean treatment duration ranged between 79.8 and 84.4 days. The majority of the patients (60.9 to 68.9%) were treated between 84 and 90 days. The target exposure was 12 weeks (84 days).

**Efficacy Results:** Primary variable: The mean number of micturitions per 24 hours at endpoint and changes from baseline are presented below.

		YM178	YM178	YM178	YM178	
		OCAS	OCAS	OCAS	OCAS	Tolterodine
	Placebo	25 mg qd	50 mg qd	100 mg qd	200 mg qd	4 mg qd
	N = 166	N = 167	N = 167	N = 168	N = 166	N = 85
Micturitions/24 hrs	Mean (SD)	Mean (SD)	Mean (SD)	Mean (SD)	Mean (SD)	Mean (SD)
Baseline	11.67 (3.39)	11.87 (2.88)	11.85 (3.30)	11.81 (3.51)	11.34 (2.41)	12.31 (3.68)
Endpoint	10.25 (2.82)	9.84 (2.97)	9.71 (3.33)	9.67 (3.53)	9.27 (2.90)	10.07 (3.47)
CFB	-1.43 (3.24)	-2.03 (2.59)	-2.14 (2.47)	-2.14 (3.23)	-2.08 (2.67)	-2.23 (3.03)
% CFB	-9.79 (22.46)	-16.04 (19.90)	-17.45 (19.19)	-16.82 (22.37)	-17.34 (22.31)	-16.49 (20.04)
median	-11.11	-16.67	-17.50	-18.35	-20.00	-19.15

CFB = change from baseline

The results from the inferential analysis (FAS) are summarized in the following table.

				U	
		YM178 OCAS	YM178 OCAS	YM178 OCAS	YM178 OCAS
	Placebo	25 mg qd	50 mg qd	100 mg qd	200 mg qd
Adjusted mean CFB	-1.44	-1.88	-2.08	-2.12	-2.24
Estimated difference to		0.45	0.64	0.68	0.80
placebo		-0.43	-0.04	-0.08	-0.80
95% CI		-0.99; 0.10	-1.19; -0.10	-1.22; -0.13	-1.34; -0.25
P-value		0.1083	0.0205	0.0152	0.0041

The effect of YM178 OCAS on the mean number of micturitions per 24 hours increased with dose when compared to placebo. It was estimated from the model that at the endpoint there were 0.45, 0.64, 0.68 and 0.80 fewer micturitions per 24 hours in the YM178 OCAS 25 mg, 50 mg, 100 mg and 200 mg groups, respectively, compared to placebo. The YM178 OCAS 50 mg, 100 mg and 200 mg groups all had statistically significant larger increases when compared to placebo at endpoint. The overall comparison to placebo was statistically significant, indicating that YM178 OCAS is efficacious with respect to the mean number of micturitions per 24 hours ; the study met its primary objective.

The results of the inferential statistical analysis for the secondary variables are presented below. There was a statistically significant effect of YM178 over placebo for all secondary efficacy variables except 'mean number of nocturia episodes per 24 hours', 'mean number of incontinence episodes per 24 hours' and 'the sum of QoL questions 3 until 27 scores as assessed by ICIQ-OABqol questionnaire'. For the latter two, a statistically significant dose trend with YM178 was found in a secondary analysis. In general, differences versus placebo tended to become larger with increasing YM178 dose

Change in mean volume voided per micturition at endpoint (FAS)										
		YM178 OCAS	M178 OCAS   YM178 OCAS   YM178 OCAS   YM178							
	Placebo	25 mg qd	50 mg qd	100 mg qd	200 mg qd					
Adjusted mean CFB	7.29	15.32	27.34	25.56	33.34					
Estimated difference		8.03	20.05	18.28	26.06					
to placebo		8.03	20.03	10.20	20.00					
95% CI		-1.54; 17.60	10.48; 29.63	8.66; 27.89	16.49; 35.62					
P-value		0.0998	< 0.0001	0.0002	< 0.0001					
Change i	n mean num	ber of incontinen	ce episodes per 24	hours at endpoint	t (FAS)					
Adjusted mean CFB	-0.53	-1.36	-1.15	-1.06	-1.10					
Estimated difference		0.84	0.62	0.52	0.58					
to placebo		-0.84	-0.02	-0.33	-0.38					
95% CI		-1.45; -0.23	-1.22; -0.02	-1.12; 0.06	-1.16; 0.01					
P-value		0.0072	0.0416	0.0758	0.0551					
Change	e in mean nu	mber of nocturia	episodes per 24 h	ours at endpoint (	FAS)					
Adjusted mean CFB	-0.38	-0.52	-0.60	-0.42	-0.59					
Estimated difference										
to placebo		-0.15	-0.22	-0.04	-0.21					
95% CI		-0.36; 0.07	-0.44; -0.01	-0.26; 0.17	-0.43; 0.00					
P-value		0.1753	0.0426	0.6984	0.0523					
Change in r	nean numbe	r of urge incontin	ence episodes per	24 hours at endpo	oint (FAS)					
Adjusted mean CFB	-0.44	-1.31	-1.13	-1.18	-1.24					
Estimated difference		0.96	0.60	0.74	0.80					
to placebo		-0.80	-0.09	-0.74	-0.80					
95% CI		-1.38; -0.35	-1.18; -0.19	-1.23; -0.25	-1.29; -0.31					
P-value		0.0011	0.0068	0.0033	0.0014					
Chang	e in mean nu	umber of urgency	episodes (graded	≥3) at endpoint (F	FAS)					
Adjusted mean CFB	-1.07	-1.77	-1.67	-2.28	-2.48					
Estimated difference		0.70	0.60	1.21	1.42					
to placebo		-0.70	-0.60	-1.21	-1.42					
95% CI		-1.38; -0.01	-1.29; 0.08	-1.90; -0.52	-2.10; -0.73					
P-value		0.0456	0.0845	0.0006	0.0001					
(	Change in m	ean level of urgen	cy per 24 hours at	t endpoint (FAS)						
Adjusted mean CFB	-0.10	-0.21	-0.18	-0.29	-0.38					
Estimated difference		0.12	0.09	0.10	0.29					
to placebo		-0.12	-0.08	-0.19	-0.28					
95% CI		-0.25; 0.02	-0.22; 0.05	-0.33; -0.06	-0.41; -0.15					
P-value		0.0922	0.2189	0.0047	< 0.0001					
Change in sum of qu	ality of life s	cores as assessed	by ICIQ-OAB qu	estionnaire (quest	ions 3a, 4a, 5a, 6a)					
	·	at endp	oint (FAS)	× <b>*</b>						
Adjusted mean CFB	-1.82	-2.40	-2.51	-2.72	-3.02					
Estimated difference		0.58	0.60	0.00	1 20					
to placebo		-0.38	-0.09	-0.90	-1.20					
95% CI		-1.13; -0.02	-1.24; -0.13	-1.45; -0.34	-1.76; -0.65					
P-value		0.0410	0.0150	0.0016	< 0.0001					

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Change in sum of quality of life scores as assessed by ICIQ-OAB questionnaire (questions 3b, 4b, 5b, 6b)										
at endpoint (FAS)										
Adjusted mean	n CFB -6.	-7.83	-8.38	-8.47	-10.02					
Estimated diff to placebo	erence	-1.82	-2.37	-2.46	-4.01					
95% CI		-4.15; 0.52	-4.70; -0.03	-4.80; -0.12	-6.34; -1.68					
P-value		0.1273	0.0474	0.0396	0.0008					
Change in sum of quality of life scores as assessed by ICIQ-OABqol questionnaire (questions 3 to 27)										
endpoint (FAS)										
Adjusted mean	ean CFB -16.11 -17.09 -20.36 -20.57 -									
Estimated diff to placebo	erence	-0.98	-4.25	-4.46	-6.08					
95% CI		-5.88; 3.92	-9.13; 0.62	-9.37; 0.46	-11.0; -1.19					
P-value		0.6943	0.0872	0.0754	0.0149					
Change in	n sum of qualit	y of life scores as as	sessed by ICIQ-OA	<b>B</b> qol questionnaire	(question 28) at					
0	-	en	dpoint (FAS)							
Adjusted mean	n CFB -1.	50 -1.85	-2.12	-2.17	-2.52					
Estimated diff	erence	0.35	0.61	0.67	1.02					
to placebo		-0.33	-0.01	-0.07	-1.02					
95% CI		-0.96; 0.27	-1.23; 0.00	-1.29; -0.05	-1.63; -0.40					
P-value		0.2709	0.0518	0.0330	0.0012					
CFB = change	e from baseline									
The efficacy	of YM178 OC	CAS and tolterodin	e 4 mg qd were in t	the same range.						
<b>Pharmacoki</b>	i <b>netic Results</b> he table below	: The summary star	tistics of the individ	dual values of the F	PK parameters are					
Dose	(mg)	VM178 OCAS	VM178 OCAS	VM178 OCAS	VM178 OCAS					
Dust	(iiig)	25  mg ad	50 mg ad	100 mg ad	200 mg ad					
	Statistic		00 mg qu	100 mg qu	200 mg qu					
	Mean	6.8	16.6	47.6	137.8					
	SD CV (%)	2.7 40.0	5 0 30 3	16.9.35.5	40 2 29 2					
C <sub>max</sub>	Min - max	3 3-22 1	7 4-35 0	10.0-121.3	40.2, 29.2 43 2_273 A					
(ng/mL)	Median	6.0	15.9	46.1	43.2-273.4					
	N	135	138	146	146					
	Mean	1.8	4.5	12.0	34.8					
	SD CV (%)	0 7 37 1	17370	4 6 36 0	96 27 6					
Ctrough	Min - max	0.8-4.8	1 7-14 2	2 5-29 2	12 0-71 5					
(ng/mL)	Median	17	4 3	12.7	34.7					
	N	135	138	146	146					
	Mean	77 9	191.8	548.0	1547.4					
	SD CV (%)	298 382	63 5 33 1	194 8 35 5	4267 276					
AUC	50,0 (70)	27.0, 50.2	05.5, 55.1	177.0, 55.5	-r20.7, 27.0					

Median 69.8 Ν 135 138 146 146 Increasing the dose from 25 to 200 mg qd (an 8-fold increase) resulted in a more than doseproportional increase in each of the parameters. The mean values of C<sub>max</sub>, C<sub>trough</sub> and AUC<sub>tau</sub> increased approximately 21, 19, and 20-fold, respectively.

77.6-498.2

184.0

110.0-1319.8

524.0

34.9-203.2

Safety Results: Overall, 45.2% of all patients had 1 or more treatment-emergent AEs during the study. Patients from placebo treatment had the lowest incidence of AEs (43.2%), and patients from 4 mg tolterodine qd had the highest incidence of AEs (48.2%). The incidence of AEs in the YM178 OCAS treatment groups slightly increased from 43.8% for the 25 mg qd YM178 OCAS group to 47.9% for the 200 mg qd YM178 OCAS treatment group.

There was no relevant difference in the type of AEs or the number of patients reporting AEs between YM178 OCAS treatment and placebo or tolterodine treatment and there was no dose relationship.

(ng.h/mL)

Min - max

508.1-2778.5

1543.3

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summary	table of tr	eatme	ent-eme	ergen	t AEs i	s pre	sented.									
		Placebo		YM178 OCAS		Y 0	YM178 OCAS		YM178 OCAS		YM178 OCAS		Toltero- dine		Total	
Number (%) of patients		(N =	= 169)	25 (N	mg qd = 169)	50 (N	mg qd = 169)	100 (N	mg qd = 168)	200 (N	mg qd = 167)	4 1 (N	mg qd = 85)	(N :	= 927)	
		n	(%)	n	(%)	n	(%)	n	(%)	n	(%)	n	(%)	n	(%)	
With AEs <sup>1</sup>	1	73	(43.2)	74	(43.8)	74	(43.8)	77	(45.8)	80	(47.9)	41	(48.2)	419	(45.2)	
Total numb	oer of AEs	132		148		171		148		164		78		841		
With SAE	s <sup>1</sup>	1	(0.6)	1	(0.6)	1	(0.6)	2	(1.2)	3	(1.8)	1	(1.2)	9	(1.0)	
Total numb	per of SAEs	2		2		1		2		3		1		11		
With AEs	Mild	34	(20.1)	38	(22.5)	37	(21.9)	40	(23.8)	46	(27.5)	19	(22.4)	214	(23.1)	
by	Moderate	37	(21.9)	29	(17.2	33	(19.5)	32	(19.0)	30	(18.0)	19	(22.4)	180	(19.4)	
severity <sup>1</sup>	Severe	2	(1.2)	7	(4.1)	4	(2.4)	5	(3.0)	4	(2.4)	2	(2.4)	24	(2.6)	
	Missing	0	-	0	-	0	-	0	-	0	-	1	(1.2)	1	(0.1)	
Who disco because of	ntinued AEs <sup>1, 2</sup>	5	(3.0)	9	(5.3)	4	(2.4)	4	(2.4)	7	(4.2)	1	(1.2)	30	(3.2)	
With treat related AE	<b>ment-</b>	26	(15.4)	34	(20.1)	38	(22.5)	36	(21.4)	37	(22.2)	13	(15.3)	184	(19.8)	
Who died		0	-	0	-	0	-	0	-	0	-	0	-	0	-	

The majority of AEs were of mild or moderate intensity. There were no deaths during the study. A summary table of treatment-emergent AEs is presented.

<sup>1</sup> Only treatment-emergent AEs are taken into account

<sup>2</sup> Only AEs that were the primary reason for discontinuation are taken into account

<sup>3</sup> AEs that are possibly or probably treatment-related, or for which the relationship is missing

The most commonly reported AEs during the treatment period were infections (bronchitis, influenza, nasopharyngitis, pharyngitis and urinary tract infection) in the class of infections and infestations, followed by gastrointestinal disorders (constipation, diarrhea, dry mouth, dyspepsia, nausea and vomiting). The majority of infection and infestation AEs were considered not treatment-related by the investigator. Approximately two-thirds of the gastrointestinal AEs were considered treatment-related by the investigator. The AEs dry mouth, constipation, dyspepsia and nausea were always (dry mouth) or mostly (constipation, dyspepsia and nausea) considered related to treatment by the investigator. The AEs diarrhea and vomiting were mostly considered unrelated to treatment by the investigator. Nine (1.0%) patients had treatment-emergent serious adverse events (SAE). These SAEs were experienced by 3 patients in the 200 mg qd YM178 OCAS group, 2 patients in the 100 mg qd YM178 OCAS group and by 1 patient in each of the remaining groups. The SAEs reported were unstable angina and cardiac failure (both reported by the same patient), hypothyroidism, gastritis, pneumonia (reported by 3 patients), complex regional pain syndrome, multiple sclerosis and pulmonary edema and hypertensive crisis (also both reported by the same patient).

The proportion of patients that discontinued because of treatment-emergent AEs was 5 (3.0%) in the placebo group, 4-9 (2.4-5.3%) in the YM178 OCAS treatment groups and 1 (1.2%) in the tolterodine 4 mg treatment group. Within the YM178 OCAS treatment groups, there was no dose-relationship. None of the changes in hematology parameters or serum chemistry parameters in YM178 OCAS-treated patients showed a dose related trend. The incidence of patients with abnormal values at the end of the 12-week treatment period was low and similar across treatment groups. An analysis of glucose levels in patients with diabetes revealed no tendency towards abnormalities in glucose levels. The results of the inferential statistical analyses of the vital signs parameters are presented below:

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Change from baseline to endpoint in mean systolic blood pressure (morning)									
		YM178 OCAS	78 OCAS   YM178 OCAS   YM178 OCAS   YM						
	Placebo	25 mg qd	50 mg qd	100 mg qd	200 mg qd				
Adjusted mean CFB	0.35	-0.70	-1.14	0.79	0.83				
Estimated difference		-1.05	-1 49	0.43	0.47				
to placebo		-1.05	-1.47	0.45	0.47				
95% CI		-3.00; 0.90	-3.44; 0.45	-1.52; 2.39	-1.48; 2.43				
P-value		0.2911	0.1318	0.6628	0.6334				
Change from baseline to endpoint in mean systolic blood pressure (afternoon)									
Adjusted mean CFB	0.38	0.57	0.39	0.65	2.24				
Estimated difference to placebo		0.20	0.01	0.28	1.86				
95% CI		-1.67; 2.06	-1.85; 1.87	-1.59; 2.15	-0.00; 3.73				
P-value		0.8367	0.9907	0.7719	0.0502				
Change	from baseline	e to endpoint in m	ean diastolic blood	l pressure (mornin	ng)				
Adjusted mean CFB	0.24	-0.77	-0.22	1.06	0.90				
Estimated difference		1.01	0.46	0.82	0.66				
to placebo		-1.01	-0.40	0.82	0.00				
95% CI		-2.15; 0.13	-1.60; 0.68	-0.32; 1.97	-0.48; 1.80				
P-value		0.0823	0.4281	0.1589	0.2550				
Change f	rom baseline	to endpoint in me	an diastolic blood	pressure (afterno	on)				
Adjusted mean CFB	0.67	0.07	0.50	1.32	0.94				
Estimated difference to placebo		-0.60	-0.17	0.65	0.28				
95% CI		-1.82; 0.61	-1.38; 1.05	-0.57; 1.87	-0.94; 1.49				
P-value		0.3313	0.7889	0.2955	0.6569				
С	hange from b	aseline to endpoin	nt in mean pulse r	ate (morning)					
Adjusted mean CFB	0.51	0.34	1.64	2.15	4.66				
Estimated difference		0.17	1.12	1.64	4.14				
to placebo		-0.17	1.15	1.04	4.14				
95% CI		-1.42; 1.08	-0.11; 2.38	0.39; 2.89	2.90; 5.39				
P-value		0.7901	0.0747	0.0103	< 0.0001				
Change from baseline to endpoint in mean pulse rate (afternoon)									
Adjusted mean CFB	-0.04	0.44	1.12	2.71	4.63				
Estimated difference to placebo		0.48	1.15	2.74	4.67				
95% CI	T	-0.92; 1.87	-0.24; 2.55	1.34; 4.14	3.27; 6.06				
P-value	T	0.5025	0.1044	0.0001	< 0.0001				
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CFB = change from baseline

The effect of YM178 OCAS on blood pressure was not statistically significant when compared to placebo at any dose level. The overall comparison for pulse rate was statistically significant, meaning that YM178 OCAS has an effect on morning and afternoon pulse rate.

Heart rate (HR) at baseline was comparable between treatment groups. No changes over time were observed, except for the 100 mg and 200 mg YM178 OCAS groups, where a small, dose-related, increase was observed. In the 100 mg YM178 OCAS group, mean HR increased by 1.7, 3.2, and 2.6 bpm at Visits 3, 4, and 6 (Weeks 1, 4 and 12), respectively. In the 200 mg YM178 OCAS group, mean HR increased by 4.5, 5.1, and 3.8 bpm at Visits 3, 4, and 6 (Weeks 1, 4 and 12), respectively. QTcF at baseline was comparable between YM178 OCAS treatment groups. No consistent changes over time were observed.

There was a mean decrease in PVR in all treatment groups, except for the YM178 OCAS 100 mg group, where a small increase was observed. However, the mean changes were small, and the interpatient variability was very high.

## CONCLUSIONS:

Based on the results of this study, it is concluded that:

- YM178 was efficacious in the treatment of patients with OAB at doses of 50 mg, 100 mg, and 200 mg in a dose-dependent manner.
- YM178 was well tolerated in patients with OAB.
- A small dose-dependent increase in pulse rate was observed across the dose range studied (25-200mg YM178 OCAS once daily). This pulse rate change was not associated with an overt increased incidence of cardiovascular adverse events.

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