# Validation of the OMRON M6 AC (HEM-7322-E) upper arm blood pressure monitor, in oscillometry mode, for clinic use and self measurement in a general population, according to the European Society of Hypertension International Protocol revision 2010

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#### Abstract

The OMRON M6 AC (HEM-7322-E), an upper arm blood pressure monitor, in oscillometry mode, for clinical use and self measurement, was validated, in a general population, according to the European Society of Hypertension International Protocol revision 2010. The protocol requirements were followed precisely. The device passed all of the requirements and, fulfilling the standards of the protocol, is recommended for clinical use.

#### **Device Details**

Jevice Details					
Brand	OMRON				
Model	M6 AC (HEM-7322-E)				
Manufacturer	OMRON HEALTHCARE Co., Ltd				
Location	Upper Arm				
Method	Oscillometry				
Purpose	Clinic Measurement, Self/ Home Measurement				
Operation	Fully Automatic				
Arm Cuff	Standard Adult: 22.0 cm to 42.0 cm				
Other Features	The function to guide cuff wrapping,				
	Memory capacity for 100 readings and 2users, 3 readings average value within 10 minutes, morning/evening				
	average, The indicator for blood pressure level, The function to detect body motion, The function to detect irregula				
	heatbeat.				

# Methodology

# Familiarisation

Hundreds of test-measurements were carried out. No problems were encountered.

#### Recruitment

Hypertesive subjects were recruited from outpatients clinic in the Department of Cardiology in Kansai Medical University, Hirakata Hospital (Osaka, Japan). Some participated immediately without appointment. Normotensive subjects were recruited from outpatients and volunteers. There were some difficulties in recruiting subjects with DBP in the high range.

# Screening and Recruitment Details

Screening and Recruitment			Recruitment Ranges				
Total Screened	40		mmHg All		All	On Rx	
Total Excluded	7			Laur	< 90	0	4
Ranges Complete	0			Low	90 - 129	11	1
Ranges Adjustment	0		SBP	Medium	130 - 160	11	2
Arrhythmias	4			Llinda	161 - 180	8	1
Device Failure	0			High	> 180	3	
Poor Quality Sounds	3						
Cuff Size Unavailable	0			< 40		0	0
Observer Disagreement	0			Low	40 - 79	10	2
Distribution	0		DBP	Medium	80 - 100	12	2
Other Reasons	0			Lliab	101 - 130	10	0
Total Recruited		33		High	> 130	1	0

#### Procedure

The European Society of Hypertension International Protocol revision 2010 for the validation of blood pressure measuring devices in adults was followed precisely.[1] Overseen by an independent supervisor, measurements were recorded by two observers blinded from both each other's readings and from the device readings.

# Results

# Subject Details

Sex			
Male : Female	20 : 13		
Age (years)			
Range (Low : High)	28 : 71		
Mean (SD)	48.7 (11.5)		
Arm Circumference (cm)			
Range (Low : High)	22.2 : 41.8		
Mean (SD)	31.4 (5.7)		
Cuff for test device			
Standard	33	(22.0 - 42.0 cm)	
	SBP	DBP	
Recruitment BP (mmHg)			
Range (Low : High)	92 : 188	46 : 133	
Mean (SD)	144.9 (26.7)	90.2 (19.0)	

## Observer Measurements in each Recruitment Range

SBP (mmHg)		DBP (mmHg)	
Overall Range (Low : High)	84 : 188	Overall Range (Low : High)	48 : 128
Low (< 130)	34	Low (< 80)	24
Medium (130 – 160)	42	Medium (80 – 100)	40
High (> 160)	23	High (> 100)	35
Maximum Difference	19	Maximum Difference	16

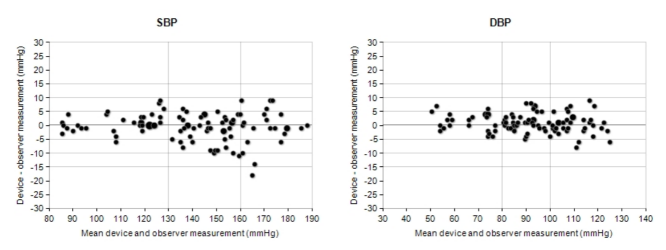
# **Observer Differences**

	SBP (mmHg)	DBP (mmHg)	Repeated measurements	
Observer 2 – Observer 1				
Range (Low : High)	-2:+4	-4 : +4		
Mean (SD)	0.3 (1.3)	0.0 (1.4)	0	

Part 1	≤ 5 mmHg	≤ 10 mmHg	≤ 15 mmHg	Grade 1	Mean (mmHg)	SD (mmHg
Pass Requirements						
Two of	73	87	96			
All of	65	81	93			
Achieved						
SBP	75	96	98	Pass	-0.5	5.0
DBP	85	99	99	Pass	1.1	3.4
Part 2	2/3 ≤ 5 mm	nHg 0/	′3 ≤ 10 mmHg	Grade 2		Grade 3
Pass Requirements	≥ 24		≤ 3			
Achieved						
SBP	26		1	Pass		Pass
DBP	28		1	Pass		Pass
Part 3						Result
						PASS

#### Validation Results

#### Plots



### Discussion

No specific problems were encountered during validation and distribution conditions were fulfilled. But recruitment of subjects with high BP, particularly high DBP, was difficult.

As the device has reached the required standards, it is recommended for clinical and personal use in a general population.

## Conclusion

As the device has reached the required standards, it is recommended for clinical and personal use in a general population.

# Acknowledgements and Conflict of Interest

The monitor was supplied for the purposes of the study by the manufacturer OMRON Healthcare Co.,LTD. who also funded the study. None of the authors has any association with OMRON Healthcare Co.,LTD. or has received any personal benefit from OMRON Healthcare Co.,LTD.

# References

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