# Hypertensive emergencies and urgencies: a preliminary report of the ongoing Italian multicentric study ERIDANO 

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#### Abstract

Hypertensive urgencies (HU) and hypertensive emergencies (HE) are challenges for the Emergency Department (ED). A prospective multicentre study is ongoing to characterize patients with acute hypertensive disorders, prevalence of subclinical hypertension-mediated organ damage (HMOD), short- and long-term prognosis; this is a preliminary report. Patients admitted to the ED with symptomatic blood pressure (BP) $\geq 180 / 110 \mathrm{mmHg}$ were enrolled. They were managed by ED personnel according to their clinical presentations. Subsequently they underwent clinical evaluation and subclinical HMOD assessment at a Hypertension Centre within 72 h from enrolment. 122 patients were included in this report. Mean age was $60.7 \pm 13.9$ years, $52.5 \%$ were females. $18(14.8 \%)$ patients were diagnosed with HE, $108(88.5 \%)$ with HU . There were no differences in gender, BMI, and cardiovascular comorbidities between groups. At ED discharge, $66.7 \%$ and $93.6 \%$ ( $p=0.003$ ) of HE and HU patients, respectively, had $\mathrm{BP}<180 / 110 \mathrm{mmHg}$. After $72 \mathrm{~h}, 34.4 \%$ of patients resulted normotensive; $35.2 \%, 22.1 \%$, and $8.2 \%$ had hypertension grade 1,2 , and 3 , respectively. Patients with uncontrolled BP at office evaluation had higher vascular HMOD ( 49.1 vs. $25.9 \%, p=0.045$ ). Cardiac ( 60 vs. $34 \%, p=0.049$ ), renal ( 27.8 vs. $9.6 \%, p=0.010$ ) and cerebral ( $100 \mathrm{vs} .21 \%, p<0.001$ ) HMOD was more frequent in HE compared to HU group. HE showed greater cardiac, renal, and cerebral subclinical HMOD, compared to HU. 72-hours BP control is not associated with different HMOD, except for vascular HMOD; therefore, proper comprehensive examination after discharge from the ED could provide added value in cardiovascular risk stratification of such patients.


Keywords emergency department • hypertensive emergencies • hypertensive urgencies • short-term blood pressure control $\cdot$ hypertension mediated organ damage

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## Graphical Abstract

One third of patients with acute blood pressure rise evaluated to the ED resulted normotensive at office evaluation ( $<72$ hours after discharge). Patients with hypertensive emergency showed greater cardiac, renal, and cerebral subclinical HMOD, compared to the patients with hypertensive urgency. BP: blood pressure; HMOD: hypertension-mediated organ damage; y.o.: years old; mo.: months.

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## METHODS

RESULTS

| Time 0 | Enrolment in |
| :---: | :---: |
|  | Emergency Department |
|  | Symptomatic BP $\geq 180 / 110 \mathrm{mmHg}$ |
|  | $\rightarrow$ Acute BP control |
|  | $\rightarrow$ Acute HMOD assessment |
|  | - Hypertensive Emergency (HE) |
|  | Hypertensive Urgency (HU) |

Admission BP
Systolic $201 \pm 20 \mathrm{mmHg}$ Diastolic $113 \pm 13 \mathrm{mmHg}$ Discharge BP
Systolic $152 \pm 21 \mathrm{mmHg}$
Diastolic $88 \pm 12 \mathrm{mmHg}$

$3^{\text {rd }}$ and $12^{\text {th }}$ mo. Ongoing follow-up
(Data not yet available)



## Introduction

Acute blood pressure (BP) disorders are a major challenge for the Emergency Department (ED). The prevalence of acute BP disorders considerably differs among studies, even depending on the definition used, but it ranges from $0.24 \%$ to $2.4 \%$ of ED admissions for hypertensive urgencies (HU) and from $0.08 \%$ to $0.76 \%$ for hypertensive emergencies (HE) [1]. These prevalences seem comparable across continents [2], although with some differences probably due to ethnic disparities, medication adherence, and insurance status [3]. Although HU do not appear to be associated with short-term adverse outcomes [4, 5], or at least have significantly lower in-hospital mortality compared to HE [6],
long-term implications, such as risk of stroke and fatal or non-fatal cardiovascular events, are relevant [7-9].

Despite the significant clinical and epidemiological impact, the management of patients with acute BP disorders is still very uneven among professionals of critical areas, as pointed out by a recent Italian surveys [10, 11]. The lack of good-quality evidence makes it difficult to propose strong recommendations for clinical practice. The therapeutic management is very uneven, especially for HU. The timing of follow-up, when present, is heterogeneous and it is not clear whether a referral to a Hypertension Centre could have a prognostic role compared to standard care.

In order to obtain more and more accurate information on this category of patients, we are conducting the ERIDANO
prospective multicenter cohort study on behalf of the Italian Society of Hypertension (SIIA: Società Italiana dell'Ipertensione Arteriosa). The aim of the ERIDANO study is to characterize patients with acute hypertensive disorders, their prevalence of subclinical organ damage and secondary hypertension and their short- and long-term prognosis, by providing referral to a Hypertension Centre immediately after discharge from the ED, as described below.

The present study is intended to be a preliminary, mainly descriptive, report of the first hundred patients enrolled, focusing on the clinical and demographic characteristics, on the management in the ED, on BP control within 72 hours of discharge, and on the prevalence of hypertensionmediated subclinical organ damage (HMOD).

## Methods

The current enrolment has involved 6 Italian hospitals, officially starting in Turin, the main center, in January 2020. Enrollment is not to be considered consecutive, as there were many months of interruption, caused by the closure of Hypertension Centers during the four Italian epidemic wave and the commitment of internal medicine, emergency medicine, and cardiology specialists to COVID wards; in addition, the other recruiting centers became active in the first months of the year 2021 or 2022.

Consecutive patients, aged 18 years and over, admitted to the ED with a symptomatic BP rise, defined as systolic BP $\geq$ 180 mmHg and/or diastolic BP $\geq 110 \mathrm{mmHg}$ associated to at least one symptom consistent with suspected HE as defined by latest guidelines [12], were enrolled. BP measurements were performed according to the current European Society of Hypertension/European Society of Cardiology (ESH/ESC) recommendations [13], with validated automatic sphygmomanometers (e.g., Omron, M10-IT models, Matsusaka, Kyoto, Japan), with patients in the sitting position whenever possible. Three BP measurement were performed, and the mean value was used for subsequent analysis.

Patients with BP rise due to traumatic causes or known neoplastic pain, or with BP rise without any associated symptoms were excluded, as were those who withheld their informed consent.

Enrolled patients were managed by the emergency physicians in the ED, according to their clinical presentations, as suggested in the current European position paper [12]. After appropriate work-up, in the presence of acute organ damage (coronary ischemia, acute cardiogenic pulmonary oedema, acute ischemic or hemorrhagic stroke, hypertensive encephalopathy, acute aortic disease, acute kidney injury, relative to known creatinine values in the previous 12 months) as defined by current guidelines [12] (HE),
patients were admitted to an appropriate hospital specialist setting; in the absence of acute organ damage (HU), they were discharged after a period of observation. In any case, an evaluation at an ESH Hypertension Excellence Centre was performed within 72 hours of enrolment; this visit was carried out on an outpatient-basis for discharged patients (HU), and on an inpatient-basis for HE patients, still hospitalized in the appropriate specialist setting. Subsequent therapeutic modifications, or indications for further diagnostic investigations, related to the detection of subclinical organ damage (which may be present independently of the acute organ damage), have been left to the discretion of the hypertension specialist, always guided by current guidelines [13]. The presence of subclinical HMOD does not reclassify patients into HE or HU , the definition of which is based on acute clinical damage.

Figure 1 summarizes the study protocol, although data from visit 2 and visit 3 have not yet been considered in the present report.

## Subclinical HMOD criteria

## Subclinical cardiac HMOD—Echocardiography

Standard two-dimensional transthoracic echocardiographic (TTE) images were acquired by expert accredited staff with commercially available ultrasound machines (e.g., IE33, Phillips Medical Systems, Andover, Massachusetts, USA). Conventional parameters were assessed according to the current guidelines [14]. Left ventricular (LV) mass was estimated. Dubois’ formula was used to calculate body surface area (BSA) and LV mass values were indexed for BSA (LVMi). LV volumes and ejection fraction, and left atrial volume were assessed using Simpson's Biplane technique from apical two and four-chamber views. LV diastolic function was estimated through the evaluation of left atrial volume, mitral inflow peak systolic velocities of early (E) and late (A) diastolic filling on pulsed-wave Doppler, color-tissue Doppler imaging of the septal and lateral mitral annulus ( $\mathrm{E}^{\prime}$ ), according to current international recommendations [15].

Alterations of LV mass and geometry, increased left atrial volume, and diastolic dysfunction were considered subclinical cardiac HMOD [13, 16]. LV hypertrophy (LVH) was defined by LVMi $>115 \mathrm{~g} / \mathrm{m}^{2}$ in men and $>95 \mathrm{~g} / \mathrm{m}^{2}$ in women [13, 14]. Relative wall thickness (RWT) was defined as two-times inferolateral wall thickness divided by the LV diastolic diameter and was used to classify LV remodeling as either concentric ( $\mathrm{RWT}>0.42$ ) or eccentric ( $\mathrm{RWT} \leq 0.42$ ). Left atrial enlargement (LAe) was considered as left atrial volume indexed to BSA (LAVi) > $34 \mathrm{ml} / \mathrm{m}^{2}$ [14].


Fig. 1 Summary of Eridano Study protocol. ED emergency department, HMOD hypertension mediated organ damage

## Subclinical vascular HMOD

Arterial stiffness was quantified using carotid-femoral pulse wave velocity (PWV). Pressure waveforms at the carotid and femoral artery were obtained non-invasively by applanation tonometry with validated instruments (e.g., Sphygmocor, AtCor Medical-Sydney, Australia) [17].

Carotid artery imaging assessment was performed by experienced staff using available ultrasound machines, equipped with $4-12 \mathrm{MHz}$ linear-array ultrasound transducer. The common carotid artery (CCA) intima-media thickness (IMT) was detected by validated software (e.g., Q-lab, Philips) on longitudinal bidimensional imaging. When clinically indicated patients underwent further imaging investigation.
$\mathrm{PWV}>10 \mathrm{~m} / \mathrm{s}$ and CCA $\mathrm{IMT}>0.9 \mathrm{~mm}$ or the presence of carotid plaques (identified by an IMT $\geq 1.5 \mathrm{~mm}$, or by a focal increase in thickness of 0.5 mm or $50 \%$ of the surrounding carotid IMT value) were considered subclinical vascular HMOD [13, 17].

## Subclinical renal HMOD

Estimated glomerular filtration rate (eGFR) was assessed with Chronic Kidney Disease Epidemiology Collaboration (CKD-EPI) formula based on serum creatinine measured within 3 months from ED presentation [18]. Moreover,
patients underwent microalbuminuria assessment. eGFR $<60 \mathrm{ml} / \mathrm{min} / 1.73 \mathrm{~m}^{2}$, urinary albumin/creatinine ratio $>30 \mathrm{mg} / \mathrm{g}$, and albuminuria $>30 \mathrm{mg} / 24 \mathrm{~h}$ were considered endpoints of significant renal HMOD [13, 19].

## Subclinical cerebral HMOD

When clinically indicated, according to ED presentations, patients underwent brain imaging, either by computed tomography or magnetic resonance imaging. The presence of white matter lesions, microinfarcts (e.g., lacunar infarctions), microbleeds, and brain atrophy identified by experienced radiologists were considered cerebral HMOD [13, 20, 21].

## Statistical analysis

Statistical analysis was performed by a dedicated software (R: A Language and Environment for Statistical Computing, v4.0.0 for Mac OSX, R Core Team., Vienna, Austria). Continuous variables were expressed as mean $\pm$ standard deviation. Qualitative variables were expressed as absolute values of frequency and percentage values. Normal distribution of variables was tested using the KolmogorovSmirnov and residual analysis tests. Differences between independent groups were evaluated using a $t$-test for continuous variables with normal distribution and the Mann-

Table 1 Demographic and clinical characteristics of study population

|  | Total $N=122$ | HE $N=18$ | HU $N=104$ | $p$ value |
| :--- | :--- | :--- | :--- | :--- |
| Male Sex [no. (\%)] | $58(47.5 \%)$ | $9(50.0 \%)$ | $49(47.1 \%)$ | 0.821 |
| Age (y) | $60.7 \pm 13.9$ | $66.5 \pm 15.9$ | $60.0 \pm 13.5$ | 0.134 |
| Height (cm) | $165 \pm 10$ | $166 \pm 9$ | $165 \pm 11$ | 0.644 |
| Weight (kg) | $79.6 \pm 19.4$ | $78 \pm 25$ | $79 \pm 19$ | 0.883 |
| BMI (kg/m²) | $28.9 \pm 5.78$ | $28.6 \pm 7.2$ | $28.9 \pm 5.6$ | 0.826 |
| ED SBP (mmHg) | $201 \pm 20$ | $205 \pm 18$ | $200 \pm 20$ | 0.372 |
| ED DBP (mmHg) | $113 \pm 13$ | $110 \pm 14$ | $113 \pm 13$ | 0.357 |
| Discharge SBP (mmHg) | $152 \pm 21$ | $155 \pm 25$ | $151 \pm 20$ | 0.669 |
| Discharge DBP (mmHg) | $88 \pm 12$ | $87 \pm 14$ | $88 \pm 12$ | 0.820 |
| ED Stay (h) [IQ range] | $7.2[4.7 ; 12.8]$ | $5.6[4.7 ; 18.7]$ | $7.2[4.7 ; 12.2]$ | 0.900 |
| BP <180/110 at ED discharge [no. (\%)] | $96(78.7 \%)$ | $8(44.4 \%)$ | $88(84.6 \%)$ | 0.003 |
| Office SBP (mmHg) | $147 \pm 22$ | $149 \pm 22$ | $147 \pm 23$ | 0.680 |
| Office DBP (mmHg) | $87 \pm 15$ | $88 \pm 15$ | $87 \pm 16$ | 0.746 |
| Difference ED-Office SBP (mmHg) | $54 \pm 28$ | $56 \pm 34$ | $53 \pm 27$ | 0.770 |
| Difference ED-Office DBP (mmHg) | $26 \pm 17$ | $22 \pm 19$ | $26 \pm 17$ | 0.322 |
| Silent medical history [no. (\%)] | $20(16.4 \%)$ | $1(5.6 \%)$ | $19(18.3 \%)$ | 0.179 |
| Arterial Hypertension [no. (\%)] | $94(77.0 \%)$ | $17(94.4 \%)$ | $77(74.0 \%)$ | 0.057 |
| Hypertension duration (y) [IQ range] | $10.0[5.0 ; 18.0]$ | $15.5[10.0 ; 28.5]$ | $10.0[5.0 ; 16.0]$ | 0.066 |
| Diabetes [no. (\%)] | $24(19.7 \%)$ | $4(22.2 \%)$ | $20(19.2 \%)$ | 0.768 |
| Dyslipidemia [no. (\%)] | $36(29.5 \%)$ | $8(44.4 \%)$ | $28(26.9 \%)$ | 0.132 |
| CAD [no. (\%)] | $15(12.3 \%)$ | $4(22.2 \%)$ | $11(10.6 \%)$ | 0.165 |
| Heart failure [no. (\%)] | $5(4.1 \%)$ | $2(11.1 \%)$ | $3(2.9 \%)$ | 0.104 |
| Atrial fibrillation [no. (\%)] | $7(5.7 \%)$ | $1(5.6 \%)$ | $6(5.8 \%)$ | 0.971 |
| Previous stroke [no. (\%)] | $5(4.1 \%)$ | $2(11.1 \%)$ | $3(2.9 \%)$ | 0.104 |
| CKD [no. (\%)] | $8(6.6 \%)$ | $2(11.1 \%)$ | $6(5.8 \%)$ | 0.398 |

$B M I$ body mass index, $B P$ blood pressure, $C A D$ coronary artery disease, $C K D$ chronic kidney disease, $E D$ emergency department, $H E$ hypertensive emergencies, $H U$ hypertensive urgencies, $S B P$ systolic blood pressure, $D B P$ diastolic blood pressure

Whitney or Kruskal-Wallis test for continuous variables with non-normal distribution. Categorical variables were compared using the chi-square test or Fisher's exact test, as appropriate. Statistical significance was considered for $p$ values $<0.05$.

The present study was firstly approved by the Institutional Review Committee of Turin (Comitato Etico Interaziendale A.O.U. Città della Salute e della Scienza di Torino - A.O. Ordine Mauriziano, CS2/1075), as well as by the local ethics committees of each participating center. All subjects gave their written informed consent.

## Results

A total of 113,694 patients were registered in the ED during the months of active enrollment; 1910 (1.7\%) admissions were due to acute BP elevation, but only $122(0.1 \%$ of the total) met the inclusion/exclusion criteria. Therefore, a total of 122 patients ( $52.5 \%$ female) with a mean age of $60.7 \pm 13.9$ years were enrolled until May 2022 and thus
included in the present report. A total of 18 patients ( $14.8 \%$ ) had acute organ damage at ED presentation (HE), whereas the remaining 104 (85.2\%) patients were diagnosed as HU. These data correspond to a prevalence of $0.09 \%$ (104/ $113,694)$ for HU and $0.02 \%(18 / 113,694)$ for HE.

The acute organ damages detected were heart failure (no. $7,39 \%$ ), stroke (no. 6, 33\%), acute coronary syndrome (no. $2,11 \%$ ), hypertensive encephalopathy (no. $2,11 \%$ ), aortic dissection (no. $1,6 \%$ ). An eGFR $<60 \mathrm{ml} / \mathrm{min} / 1.73 \mathrm{~m}^{2}$ was found in 5 patients at ED evaluation, but the presence of similar creatinine values in the previous 12 months (available in the informatic system), did not make them categorize as acute kidney injury.

No significant difference emerged between HE and HU groups in terms of gender, BMI, cardiovascular comorbidities (Table 1). Hypertensive therapy ongoing at ED admission is listed in Table 2.

At ED presentation mean systolic BP was $201 \pm 20 \mathrm{mmHg}$ and mean diastolic BP was $113 \pm 13 \mathrm{mmHg}$, without significant difference between HE and HU patients. The most common clinical presentation was headache (46.7\%),

Table 2 Ongoing hypertensive therapy and medications of study population at ED admission

| Previous Hypertensive Therapy | Total $N=122$ | HE $N=18$ | HU $N=104$ | $p$ value |
| :--- | :---: | :--- | :--- | :--- |
| Previous Hyp therapy [no. (\%)] | $85(69.7 \%)$ | $15(83.3 \%)$ | $70(67.3 \%)$ | 0.172 |
| Nr. Previous Hyp drugs [IQ range] | $1.0[0.0 ; 2.0]$ | $1.0[1.0 ; 2.0]$ | $1.0[0.0 ; 2.0]$ | 0.471 |
| Previous Hyp drugs $\geq 3$ [no. (\%)] | $23(18.9 \%)$ | $2(11.1 \%)$ | $21(20.2 \%)$ | 0.363 |
| ACE-Inhibitors [no. (\%)] | $33(27.0 \%)$ | $8(44.4 \%)$ | $25(24.0 \%)$ | 0.072 |
| ARB [no. (\%)] | $30(24.6 \%)$ | $1(5.6 \%)$ | $29(27.9 \%)$ | 0.042 |
| CCB [no. (\%)] | $27(22.1 \%)$ | $6(33.3 \%)$ | $21(20.2 \%)$ | 0.215 |
| CCB NDH [no. (\%)] | $0(0.0 \%)$ | $0(0.0 \%)$ | $0(0.0 \%)$ | - |
| Beta-blockers [no. (\%)] | $44(36.1 \%)$ | $7(38.9 \%)$ | $37(35.6 \%)$ | 0.787 |
| Alfa-blockers [no. (\%)] | $9(7.4 \%)$ | $0(0.0 \%)$ | $9(8.7 \%)$ | 0.195 |
| Alfa2-agonist [no. (\%)] | $2(1.6 \%)$ | $0(0.0 \%)$ | $2(1.9 \%)$ | 0.553 |
| MRA [no. (\%)] | $1(0.8 \%)$ | $0(0.0 \%)$ | $1(1.0 \%)$ | 0.676 |
| Thiazides [no. (\%)] | $14(11.5 \%)$ | $2(11.1 \%)$ | $12(11.5 \%)$ | 0.958 |
| Loop diuretics [no. (\%)] | $8(6.6 \%)$ | $1(5.6 \%)$ | $7(6.7 \%)$ | 0.852 |
| Potassium sparing [no. (\%)] | $1(0.8 \%)$ | $0(0.0 \%)$ | $1(1.0 \%)$ | 0.676 |
| Nitrates [no. (\%)] | $3(2.5 \%)$ | $1(5.6 \%)$ | $2(1.9 \%)$ | 0.358 |
| Others hyp drugs [no. (\%)] | $2(1.6 \%)$ | $1(5.6 \%)$ | $1(1.0 \%)$ | 0.156 |
| Benzodiazepines [no. (\%)] | $7(5.7 \%)$ | $1(5.6 \%)$ | $6(5.8 \%)$ | 0.971 |

ACE-Inhibitors inhibitors of angiotensin-converting enzyme, $A R B$ angiotensin II receptor blockers, $C C B$ calcium channel blockers, $C C B-N D H$ non-dihydropyridine $C C B, E D$ emergency department, $H E$ hypertensive emergencies, $H U$ hypertensive urgencies, $H y p$ hypertension, $M R A$ mineralocorticoid receptor antagonists
followed by chest pain (23.8\%), dyspnea (14.8\%), and neurological symptoms ( $6.6 \%$ ), while other non-specific symptoms were present in $68.9 \%$ of patients.

A silent medical history was present in 20 patients (16.4\%). Moreover, 94 patients (77\%) had previously known arterial hypertension and 85 (69.7\%) were on antihypertensive medical therapy, with a median number of medications of 1.0 [IQ range 0.0;2.0]; 23 patients ( $18.9 \%$ ) were on $\geq 3$ hypertensive drugs.

## Hypertensive therapy and BP control during ED stay

Among patients enrolled, $61.1 \%$ and $94.2 \%$ of HE and HU group ( $p<0.001$ ) received antihypertensive therapy during ED stay ( $89.3 \%$ of total population), with more drugs administered in the latter group (1.0 [0.0;2.0] vs. 2.0 [1.0;2.0] in HE and HU patients, respectively, $p=0.003$ ). A total of 25 patients ( $24 \%$ ) of HU group received 3 or more antihypertensive medications. Intravenous antihypertensive drugs were given to $27.8 \%$ and $15.4 \%$ of patients in HE and HU group ( $p=0.198$ ).

The most used class of medication was calcium channel blockers (CCB), administered to 74 patients (60.7\%) ( $22.2 \%$ vs. $67.3 \%$ in HE and HU group, respectively, $p<0.001$ ), followed by benzodiazepines, administered to 57 patients $(46.7 \%)(16.7 \%$ vs. $51.9 \%$, in HE and HU group, respectively, $p=0.006$ ) and ACE-Inhibitors, given to 53 patients $(43.4 \%)(16.7 \%$ vs. $48.1 \%$, in HE and HU group, respectively, $p=0.013$ ). The remaining
classes of drugs administered during ED stay are listed in Table 3.

After 1 hour from ED admission, $50.0 \%$ of HE patients and $76.7 \%$ of HU patients had BP values $<180 / 110 \mathrm{mmHg}$. At the time of ED discharge, these percentages increased to $66.7 \%$ of and $93.6 \%$, respectively $(p=0.003)$ ( $90.6 \%$ of total population), with a median ED stay of 7.2 hours [IQ range $4.7 ; 12.8$ ]. At ED discharge mean systolic BP was $152 \pm 21 \mathrm{mmHg}$ and diastolic BP was $88 \pm 12 \mathrm{mmHg}$.

No drugs were significantly associated with the achievement of BP values $<180 / 110 \mathrm{mmHg}$ during ED stay (data not shown).

## Office blood pressure control (72 hours after ED discharge)

At 72 hours visit patients had mean systolic BP of $148 \pm 22 \mathrm{mmHg}$ ( $p=0.037$, compared to BP at ED discharge) and diastolic BP of $88 \pm 16(p=0.944)$.

BP values $<140 / 90 \mathrm{mmHg}$ were achieved in 42 patients ( $34.4 \%$ ) who resulted normotensive at 72 hours visit ( $22.2 \%$ and $36.5 \%$ of HE and HU patients, $p=0.238$ ). 43 patients ( $35.2 \%$ ) had grade 1 hypertension, 27 ( $22.1 \%$ ) had grade 2 hypertension, and $10(8.2 \%)$ had grade 3 hypertension, with no differences between HE and HU patients ( $p=0.592$ ).

Patients with uncontrolled BP were more frequently males ( $56.3 \%$ vs. $31.0 \%, p=0.008$ ), but there were no other significant differences in terms of age, body size,

Table 3 Hypertensive therapy and medications administered during ED stay

Table 4 Hypertensive therapy and medications prescribed at 72 hours visit

| Hypertensive therapy administered in ED | Total $N=122$ | HE $N=18$ | HU $N=104$ | $p$ value |
| :--- | :---: | :--- | :--- | ---: |
| Hyp therapy in ED [no. (\%)] | $109(89.3 \%)$ | $11(61.1 \%)$ | $98(94.2 \%)$ | $<0.001$ |
| Nr. Hyp drugs in ED [IQ range] | $2.0[1.0 ; 2.0]$ | $1.0[0.0 ; 2.0]$ | $2.0[1.0 ; 2.0]$ | 0.003 |
| Hyp drugs in ED $\geq 3$ [no. (\%)] | $25(20.5 \%)$ | $0(0.0 \%)$ | $25(24.0 \%)$ | 0.020 |
| IV Hyp drugs in ED [no. (\%)] | $21(17.2 \%)$ | $5(27.8 \%)$ | $16(15.4 \%)$ | 0.198 |
| ACE-Inhibitors [no. (\%)] | $53(43.4 \%)$ | $3(16.7 \%)$ | $50(48.1 \%)$ | 0.013 |
| ARB [no. (\%)] | $13(10.7 \%)$ | $2(11.1 \%)$ | $11(10.6 \%)$ | 0.946 |
| CCB [no. (\%)] | $74(60.7 \%)$ | $4(22.2 \%)$ | $70(67.3 \%)$ | $<0.001$ |
| CCB NDH [no. (\%)] | $0(0.0 \%)$ | $0(0.0 \%)$ | $0(0.0 \%)$ | - |
| Beta-blockers [no. (\%)] | $26(21.3 \%)$ | $2(11.1 \%)$ | $24(23.1 \%)$ | 0.252 |
| Alfa-blockers [no. (\%)] | $14(11.5 \%)$ | $0(0.0 \%)$ | $14(13.5 \%)$ | 0.098 |
| Alfa2-agonist [no. (\%)] | $11(9.0 \%)$ | $1(5.6 \%)$ | $10(9.6 \%)$ | 0.579 |
| MRA [no. (\%)] | $0(0.0 \%)$ | $0(0.0 \%)$ | $0(0.0 \%)$ | - |
| Thiazides [no. (\%)] | $6(4.9 \%)$ | $0(0.0 \%)$ | $6(5.8 \%)$ | 0.296 |
| Loop diuretics [no. (\%)] | $11(9.0 \%)$ | $2(11.1 \%)$ | $9(8.7 \%)$ | 0.737 |
| Potassium sparing [no. (\%)] | $1(0.8 \%)$ | $0(0.0 \%)$ | $1(1.0 \%)$ | 0.676 |
| Nitrates [no. (\%)] | $8(6.6 \%)$ | $2(11.1 \%)$ | $6(5.8 \%)$ | 0.398 |
| Other vasodilators [no. (\%)] | $1(0.8 \%)$ | $0(0.0 \%)$ | $1(1.0 \%)$ | 0.676 |
| Others hyp drugs [no. (\%)] | $2(1.6 \%)$ | $1(5.6 \%)$ | $1(1.0 \%)$ | 0.156 |
| Benzodiazepines [no. (\%)] | $57(46.7 \%)$ | $3(16.7 \%)$ | $54(51.9 \%)$ | 0.006 |

Abbreviations as in Table 2

| Hypertensive therapy prescribed at 72 h visit | Total $N=122$ | HE $N=18$ | HU $N=104$ | $p$ value |
| :--- | :--- | :--- | :--- | ---: |
| Hyp therapy at 72 h [no. (\%)] | $105(86.0 \%)$ | $10(55.6 \%)$ | $95(91.3 \%)$ | $<0.001$ |
| Hyp drugs at 72 h (no) [IQ range] | $3.0[2.0 ; 4.0]$ | $2.0[0.0 ; 2.75]$ | $3.0[2.0 ; 4.0]$ | 0.023 |
| Hyp drugs at $72 \mathrm{~h} \geq 3$ [no. (\%)] | $65(53.3 \%)$ | $5(27.8 \%)$ | $60(57.7 \%)$ | 0.019 |
| ACE-Inhibitors [no. (\%)] | $30(24.6 \%)$ | $5(27.8 \%)$ | $25(24.0 \%)$ | 0.734 |
| ARB [no. (\%)] | $56(45.9 \%)$ | $4(22.2 \%)$ | $52(50.0 \%)$ | 0.029 |
| CCB [no. (\%)] | $85(69.7 \%)$ | $7(38.9 \%)$ | $78(75 \%)$ | 0.002 |
| CCB NDH [no. (\%)] | $15(12.3 \%)$ | $0(0.0 \%)$ | $15(14.4 \%)$ | 0.085 |
| Beta-blockers [no. (\%)] | $39(32.0 \%)$ | $6(33.3 \%)$ | $33(31.7 \%)$ | 0.893 |
| Alfa-blockers [no. (\%)] | $30(24.6 \%)$ | $1(5.6 \%)$ | $29(27.9 \%)$ | 0.042 |
| Alfa2-agonist [no. (\%)] | $2(1.6 \%)$ | $0(0.0 \%)$ | $2(1.9 \%)$ | 0.553 |
| MRA [no. (\%)] | $8(6.6 \%)$ | $2(11.1 \%)$ | $6(5.8 \%)$ | 0.398 |
| Thiazides [no. (\%)] | $25(20.5 \%)$ | $2(11.1 \%)$ | $23(22.1 \%)$ | 0.286 |
| Loop diuretics [no. (\%)] | $9(7.4 \%)$ | $3(16.7 \%)$ | $6(5.8 \%)$ | 0.102 |
| Potassium sparing [no. (\%)] | $2(1.6 \%)$ | $0(0.0 \%)$ | $2(1.9 \%)$ | 0.553 |
| Nitrates [no. (\%)] | $10(8.2 \%)$ | $1(5.6 \%)$ | $9(8.7 \%)$ | 0.658 |
| Others hyp drugs [no. (\%)] | $1(0.8 \%)$ | $1(5.6 \%)$ | $0(0.0 \%)$ | 0.016 |
| Benzodiazepines [no. (\%)] | $2(1.6 \%)$ | $1(5.6 \%)$ | $1(1.0 \%)$ | 0.156 |

Abbreviations as in Table 2
and cardiovascular comorbidities. Moreover, patients with uncontrolled BP had higher mean PWV ( $10.1 \pm 2.3$ vs. $8.9 \pm 2.2 \mathrm{~m} / \mathrm{s}, \quad p=0.017$ ) and higher prevalence of PWV $>10 \mathrm{~m} / \mathrm{s}$ ( 49.1 vs. $25.9 \%, p=0.045$ ), even after adjusting for heart rate and mean BP (data not shown). Hypertensive therapy prescribed at 72 hours visit is depicted in Table 4.

## Hypertension-mediated subclinical organ damage (HMOD) at 72 hours visit

LVH was present in 41 patients ( $33.6 \%$ of total population; $50 \%$ and $30.8 \%$ of HE and HU patients, respectively, $p=0.0 .54$ ). HE group showed higher LVMi compared to HU group ( $110.9 \pm 36.0$ vs. $93.0 \pm 26.4 \mathrm{~g} / \mathrm{m}^{2}, p=0.023$ ).

Table 5 Subclinical hypertension mediated organ damage characteristics of study population

|  | Total $N=122$ | HE $N=18$ | HU $N=104$ | $p$ value |
| :--- | :--- | :--- | :--- | ---: |
| LVMi $\left(\mathrm{g} / \mathrm{m}^{2}\right)$ | $95.5 \pm 28.4$ | $110.9 \pm 36.0$ | $93.0 \pm 26.4$ | 0.023 |
| LVH [no. $(\%)$ ] | $41(33.6 \%)$ | $9(50.0 \%)$ | $32(30.8 \%)$ | 0.054 |
| EF $(\%)$ | $61.3 \pm 7.9$ | $57.9 \pm 5.0$ | $61.9 \pm 7.0$ | 0.067 |
| LAVi $\left(\mathrm{ml} / \mathrm{m}^{2}\right)$ | $29.2 \pm 11.2$ | $37.8 \pm 17.4$ | $28.2 \pm 10.0$ | 0.014 |
| LAe [no. $(\%)$ ] | $26(21.3 \%)$ | $4(22.2 \%)$ | $22(21.2 \%)$ | 0.836 |
| Ascending aorta (mm) | $34.4 \pm 4.9$ | $36.2 \pm 5.0$ | $34.1 \pm 4.8$ | 0.171 |
| E/E' ratio | $9.28 \pm 4.57$ | $9.91 \pm 3.34$ | $9.21 \pm 4.71$ | 0.634 |
| E/E' ratio > 14 [no. (\%)] | $12(9.8 \%)$ | $3(16.7 \%)$ | $9(8.7 \%)$ | 0.081 |
| TR max vel (m/s) | $2.32 \pm 0.43$ | $2.51 \pm 0.33$ | $2.31 \pm 0.44$ | 0.380 |
| PWV (m/s) | $9.71 \pm 2.30$ | $9.83 \pm 1.54$ | $9.68 \pm 2.41$ | 0.847 |
| PWV > 10 m/s [no. (\%)] ${ }^{\text {a }}$ | $34(39.5 \%)^{\mathrm{a}}$ | $4(33.3 \%)^{\mathrm{a}}$ | $30(40.5 \%)^{\mathrm{a}}$ | 0.536 |
| Abnormal carotid US [no. $(\%)]^{\mathrm{b}}$ | $25(43.1 \%)^{\mathrm{b}}$ | $4(66.7 \%)^{\mathrm{b}}$ | $21(40.4 \%)^{\mathrm{b}}$ | 0.218 |
| Vascular HMOD [no. (\%)] | $49(53.8 \%)^{\mathrm{c}}$ | $7(46.7 \%)^{\mathrm{c}}$ | $42(55.3 \%)^{\mathrm{c}}$ | 0.542 |
| Renal HMOD [no. (\%)] | $15(12.3 \%)$ | $5(27.8 \%)$ | $10(9.6 \%)$ | 0.010 |
| Cerebral HMOD [no. (\%)] ${ }^{\mathrm{d}}$ | $16(34.8 \%)^{\mathrm{d}}$ | $8(100 \%)^{\mathrm{d}}$ | $8(21.1 \%)^{\mathrm{d}}$ | $<0.001$ |

$E / E$ ' ratio mean transmitral inflow early wave on pulsed-wave Doppler to mitral annulus (lateral/septal) early wave on tissue-doppler imaging ratio, $E F$ ejection fraction, $H M O D$ hypertension mediated organ damage, $L A e$ left atrial enlargement, LAVi left atrial volume indexed for body surface area, LVH left ventricular hypertrophy, $L V M i$ left ventricular mass indexed for body surface area, $P W V$ pulse wave velocity
${ }^{\text {a }}$ Data available for 86 patients ( 12 patients among HE, 74 patients among HU)
${ }^{\mathrm{b}}$ Data available for 58 patients ( 6 patients among HE, 52 patients among HU)
${ }^{\mathrm{c}}$ Data available for 91 patients ( 15 patients among HE, 76 patients among HU)
${ }^{\mathrm{d}}$ Data available for 46 patients ( 8 patients among HE, 38 patients among HU)

LAe was detected in 26 patients (21.3\%); no difference in LAe prevalence was found between HE and HU group ( $22.2 \%$ vs. $21.2 \%, p=0.836$ ), but the former group had significant higher LAVi ( $37.8 \pm 17.4$ vs. $28.2 \pm 10.0$, $p=0.014$ ). Systolic and diastolic function was similar between the two groups.

Subclinical vascular HMOD was assessed in 91 patients and was detected in 49 patients ( $53.9 \%$ ). Of the 82 patients whose arterial stiffness was tested, 34 (41.5\%) had PWV $>10 \mathrm{~m} / \mathrm{s}$, and of the 58 patients tested with carotid ultrasound, $25(43.1 \%)$ had CCA IMT $>0.9 \mathrm{~mm}$ or carotid plaques. Indices of subclinical vascular HMOD were proved to be comparable between the two groups (Table 5).

Subclinical renal HMOD was observed in 15 patients (12.3\%). HE patients had higher prevalence of renal damage than HU patients ( $27.8 \%$ vs. $9.6 \%, p=0.010$ ).

Brain damage was detected in 16 patients ( $34.8 \%$ of 46 patients who underwent brain imaging during ED evaluation), and it was detected in all HE patients who underwent brain imaging ( $100 \%$ vs. $21.1 \%, p<0.001$ ).

In summary, subclinical HMOD was detected in 82 patients ( $67.2 \%$ of total population), $100 \%$ of HE patients and $61.5 \%$ of HU patients $(p=0.001)$. Patients with detected subclinical HMOD were older than patients without HMOD ( $64.4 \pm 13$ vs. $53.3 \pm 12$ years, $p<0.001$ ), and had more likely history of diabetes ( $p<0.001$ ), dyslipidemia ( $p=0.042$ ), coronary artery disease ( $p=0.021$ ), and
chronic kidney disease ( $p=0.041$ ). Patients with detected subclinical HMOD were also taking higher median number of hypertensive drugs at ED admission (1.0 [0.0; 1.0] vs. 1.0 [0.0; 2.0], $p=0.004$ ), and had higher mean systolic BP values at ED admission ( $204 \pm 18$ vs. $194 \pm 20 \mathrm{mmHg}$, $p=0.007)$ and at 72 h visit $(150 \pm 23 \mathrm{vs} .140 \pm 19 \mathrm{mmHg}$, $p=0.016$ ).

## Discussion

This report described around the first hundred patients with acute hypertensive disorders enrolled within the Italian multicenter prospective study called Eridano. This study has an ambitious prognostic aim, but, at present, only descriptive data from the first visits have been presented, specifically the ED enrolment and the office evaluation within 72 hours of ED discharge.

Acute hypertensive disorders are serious medical conditions, with a combined prevalence of $1.2 \%$ of total admission in the ED, in the most recent meta-analysis on the topic [1]. In the present prospective study, it is difficult to estimate the true prevalence of these conditions, considering the changes in the ED admissions dictated by the COVID19 pandemic [22,23]. Our data correspond to a prevalence of $0.09 \%$ for HU and $0.02 \%$ for HE; much lower prevalences compared to literature data, which could partly
account for the truth regarding the lower number of ED admissions during the pandemic, but could also be the result of underestimation of data due to enrollment issue for logistical difficulties that have affected all italian hospitals in recent years. In fact, the overall prevalence of admission for acute BP disorders ( $1.7 \%$ ), not considering the inclusion/exclusion criteria of the current study, is similar to the literature data. Also, this large difference could be dictated by the high attention to the presence of symptoms at ED presentation. Most data regarding acute BP disorders consider both symptomatic and asymptomatic patients, while we focused on patients having symptoms consistent with possible acute hypertensive organ damage. This could have led to a low proportion of patients actually included in the study, among the overall patients being registered at ED presentation as having acute BP disorders.

The ratio between HE and HU is similar to those of previous studies [24-30]. Some differences are at least in part explained by the different HE/HU definitions, in terms of BP cut-off or diagnostic coding; in a large retrospective study, the prevalence of HE in the United States between 2006 and 2013 was lower, probably due to the strict definition, based on acute BP elevation together with a diagnosis of acute organ damage based on the ICD-9 code [2]. In a recent review, the prevalence of HE in the Asian population ranges from 0.1 to $1.5 \%$ [31].

Our population is younger than the previous Italian multicenter study, whose enrolment was held in 2009, by about 10 years [27], but with similar age of an Asian study from the most recent recruitment [32]. Although we need to increase the sample size to confirm these data, no differences in age, sex and cardiovascular comorbidities are currently present between HE and HU. This seems to disagree with previous findings, in which HE was associated with male sex [24, 27], older age, and comorbidities [29].

Pharmacological management in ED confirms for the umpteenth time the great inconsistency among professionals concerning the treatment of acute BP disorders, as well pointed out by the GEAR project [10]. Frequently, antihypertensive drugs are used with the goal of acutely reducing BP in HU, while there is no benefits to support this practice [4, 33]. In contrast, there are data on the possible damage from rapid BP reduction in patients without organ damage [34].

Although mostly based on expert opinion, there are official recommendations on the treatment of HE [12]; moreover, a reasoned pharmacological approach has recently been proposed, starting from the pathophysiology of HE [35, 36]. Indeed, the major problem seems to be represented by patients with HU , where the greatest discrepancies in treatment approach are found, including the use of intravenous drugs, absolutely not recommended in this context. The current European position paper [12]
suggests that HU should be treated in the same way as asymptomatic uncontrolled hypertension, by modifying home therapy without claiming rapid BP reduction in the emergency room. In these patients, oral administration of antihypertensive drugs, aimed at gradual BP reduction over the following days, is the best approach [37-39].

In our cohort, CCBs were the most widely used class; in particular, amlodipine, the most available drug in the class in Italian ED, was used in $99 \%$ of cases ( 73 out of 74 patients); nifedipine was used in only one case. These data are fortunately a marked improvement from the frequent use shown in the survey cited above [10], where $22 \%$ of participants (and $23 \%$ of those working in the ED) were inclined to use sublingual nifedipine to reduce BP, although its use has been discouraged for years because of possible deleterious effects [40]. Long-acting CCBs are also encouraged in this context because they do not interfere with diagnostics, and consequently allow the search for secondary causes of hypertension when indicated [35].

Captopril remains by far the most widely used drug within the class of ACE inhibitors (31 out of 53 patients treated with this class in our cohort). Compared to nifedipine, captopril has been shown to be equally effective in terms of BP reduction, but with fewer side effects [41]; however, considerations must be taken even with this drug due to the possible sudden hypotension [42].

A special consideration should be given to benzodiazepines, class not officially suggested but widely used in clinical practice, as evidenced by previous studies [10, 24, 29]. Administered in almost half of the cases in our cohort, benzodiazepines are definitely recommended medication in adrenergic hyperactivity BP disorders, such as cocaine abuse [38, 43], but their use outside this context would merit more in-depth studies. Patients with HU treated with benzodiazepines demonstrated greater reductions in systolic BP values, than patients not treated with anxiolytic therapy [44]. In a randomized clinical trial, diazepam demonstrated the same pressor effect as nifedipine and propranolol [45]; in another trial, the same pressor effect of captopril [46]. In the present report, $70 \%$ of prescribers considered administering benzodiazepines to reduce an obvious anxiety symptomatology associated with the BP rise, while $30 \%$ of prescribers used benzodiazepines for an expected stand-alone antihypertensive effect, independent of anxiety. The marked difference in the proportion of benzodiazepines administration between ED and ambulatory visit ( 52 vs. $1 \%$ ) is probably due to the need to counter the anxiety effect on BP during symptomatic ED presentation, not so markedly present at the office evaluation. Likely, patients with acute BP disorders, especially those without acute clinical organ damage, suffer from an overlap of true BP elevation and anxiety effect, that leads to very high BP values. This effect is also probably enhanced by the presence of symptoms. During office
evaluation, once the symptoms and the fear for lifethreatening situation are over, BP values are less influenced by anxiety, and benzodiazepines prescription is not required anymore.

The fact that not all patients with HE were treated in our cohort is surprising, but this data could be distorted by rapid admission to the intensive or semi-intensive units with treatment initiated outside the ED (indeed, the median ED stay of 7.2 hours is mainly due to HU , in a situation of Italian ED currently characterized by overcrowding and boarding problems). Furthermore, in ischemic strokes (no. 5 in our cohort), the cut-off for starting acute antihypertensive treatment is higher than that of HE diagnosis.

To our knowledge, our study is currently the only one that prospectively and systematically assesses short-term (72 hours) BP control in office setting after ED discharge, except for a small study on 21 hypertensive patients in which 24h-ABPM immediately after discharge from the ED [47]. Approximately $90 \%$ of patients in our study were discharged from the ED with $\mathrm{BP}<180 / 110 \mathrm{mmHg}$, thus no longer meeting the criteria for HU , for those without organ damage; a similar rate has been described in recent studies [48, 49]. In about one-third of the cases, normal office BP was present at 72 hours after ED discharge; similar outcome than that reported, of about $20 \%$ at 2 weeks after discharge, in a retrospective study conducted in the Thai population [49], but very different from the previously cited Israeli report in which 17 out of 21 patients remained with a SBP $>180 \mathrm{mmHg} 24$ hours after ED discharge [47].

The median number of hypertensive drugs prescribed increased from 1.0 [IQR 0.0; 2.0] before ED admission, to 2.0 [IQR 1.0; 2.0] during ED stay, and eventually to 3.0 [IQR 2.0; 4.0] at 72 h visit. These data confirm both the high BP variability in this population and the need for aggressive treatment.

Finally, we presented some data on subclinical HMOD: to our knowledge this is the first study to assess subclinical HMOD in HE and HU patients immediately after ED discharge.

In general, HE patients had worse subclinical HMOD profile than HU patients, particularly cardiac, renal, and cerebral HMOD, while vascular HMOD was comparable. At 72 h visit, patients with uncontrolled BP had worse PWV, suggesting a possible role of aortic stiffness in impeding proper BP control, or possibly grater vascular damage in patients with short-term uncontrolled BP. A recent study showed that HU patients had subclinical HMOD profile midway between patients with asymptomatic grade 3 hypertension and patients with various grade hypertension, matched for office BP [50]. The higher prevalence of subclinical HMOD in HE patients found in the present study underlines that HE patients have worse baseline CV risk profile than HU patients, leading to more
severe manifestations of acute BP rise. Moreover, this difference in subclinical HMOD was not observed when comparing patients with controlled and uncontrolled BP at 72 h visit, somehow indicating that some patients could represent a special high-risk population, irrespectively of acute and short-term BP control. Ongoing follow-up is needed to better define this aspect.

## Study limitations and future perspectives

It should be stressed first of all that the prevalence data are the result of an estimation calculated on the basis of the months of active enrollment in the various centers and the average visits to the EDs; therefore, these are numbers to be taken with caution because they could represent an underestimation of reality even if a reduction in prevalences could have been expected during COVID-19 pandemic.

As specified, complete follow-up is needed to add prognostic value to the subclinical organ damage in this category of patients. It may be interesting to assess whether early in-depth evaluation at specialized hypertension centers could improve prognosis compared to standard management especially for the category of patients without acute clinical damage.

The present study has a purely descriptive nature, impaired by the small total number and the numerical discrepancy between the two groups analyzed (HE and HU); this must make comparisons interpreted with caution. At the same time, it has the advantages of describing short-term BP control and the investigation of subclinical HMOD immediately after discharge from the ED.

Ad hoc designed studies are needed to suggest appropriate management of HU in the ED, as well as targeted education to ED physicians by hypertension experts. Finally, in addition to cardiovascular drugs, benzodiazepines may be powerful weapons, already long used in clinical practice, to treat these disorders. It would be intriguing to evaluate their real hypotensive potential, perhaps considering the psychological characteristics of each patient. The use of benzodiazepines in these cases may be beneficial for both hypotensive and stress releasing effects, while significant harm is unlikely to result, being careful not to overestimate anxiety/stress effect over BP. Anyway, at present, pending stronger evidence, the results presented do not allow benzodiazepines to be recommended in acute hypertensive disorders, except in cases of associated overt anxiety, which is itself an indication for such therapy.

## Conclusions

Acute BP disorders are a major challenge for the ED. The lack of good-quality evidence makes it difficult to propose
strong recommendations for clinical practice. In this first report about the ongoing prospective Italian multicenter study ERIDANO, we showed that great inconsistency is present in acute BP disorder management. Up to one third of patients resulted normotensive after 72 h after ED discharge. HE patients showed greater cardiac, renal, and cerebral subclinical HMOD, compared to HU patients. 72 h BP control is not associated with different subclinical HMOD, except for vascular HMOD; therefore, proper comprehensive examination after discharge from the ED could provide added value in cardiovascular risk stratification of such patients.

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## Compliance with ethical standards

Conflict of interest The authors declare no competing interests.

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