

MOLECULAR TEST REQUISITION FORM EVALUATION OF MOLECULAR RISK MARKERS FOR ARTERIAL HYPERTENSION FULL PANEL

INDEX PATIENT/ FAMILIAR INFORMATION (obligatory field, delete as applicable)

Name: Ethnicity and geographical or - from the mother Consultancy Referral Number	igin: - from index patient _ , - from the f	ather	Place here the identification label
SPECIMEN SOURCE (obt	igatory field)		
☐ Whole bloo	od 🗆 DNA [Saliva	URGENT Reason:
PHYSICIAN INFORMA	TION ^(obligatory field)		
Physician			
Telephoner	1 dx1		
MOLECULAR TEST REQ	QUESTED (obligatory field)		
Evaluation of molecu	ılar risk markers fo	r arterial hypertens	sion - Full panel 🗌
in particular genetic variant system, 2) vascular endothe	s that are related with the elium 3) renal tubule, 4) iseases associated with an	ne regulation and / or dys signal transduction syste	k markers predisposing for arterial hypertension, function of the 1) renin-angiotensin-aldosterone m, 5) sodium channels, 6) autonomous nervous ation of genetic variants that contribute to anti-
PREVIOUS GENETIC CO	NSULTANCY: Date	/; AGE O	F DIAGNOSTIC:

FAMILIAR INFORMATION

Previously studied familial members: identification in genealogical tree. Point out the individual in the present study with an arrow (/) .

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FULL PANEL

Name:	
Consultancy Referral Number:	

101	102	103 1	04 105	106	107	108 1	09 110	111	112
201 2	202 20	3 204	205	20	7 208	209	210	211 21	12
301	302	303 30	305	306	307	308 30	9 310	311	312
401 4	02 40	3 404	405 4	06 407	7 408	409	410	411 41	2

Position in the tree	Name / Consultancy Referral Number	Clinical information and age of diagnostic *

^{*} Legend: age (A) arterial hypertension (AHTN), AHTN in pregnancy, acute myocardial infarction (AMI), congestive heart failure (CHF), stroke (S), acute pulmonary edema (APE), peripheral artery disease (PAD), retinopathy (R), sudden death (SD)

CLINICAL INFORMATION: COMPLEMENTARY DIAGNOSTIC EXAMS

By filling these fields you are contributing to improve your future patient diagnostics as we are developing tools to model clinical and genetic data regarding. Arterial Hypertension risk factors.

Clinical information	Data
AHTN diagnostic date	(day) /(month) /(year), years
Age of AHTN diagnostic	
AP (previous to therapeutics)	(systolic)(diastolic) mmHg
Cardiac frequency (bpm)	
Physical activity (hours / week)	
AP (subsequent to therapeutics)	(systolic)(diastolic) mmHg
Antihypertensive therapeutics	
(dose frequency)	
Secondary AHTN - disease	renal \Box , endocrinology \Box , conjunctive tissue \Box , arterial vessels \Box
associated	

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FULL PANEL

Name:	
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Associated risk factors	poor diet with high sodium intake \Box , fast food \Box , lack of physical activity \Box , smoking
	□, nº cigarettes /day, nº packs /day, stop smoking at years,
	ethanolism \Box , stress \Box , anxiety \Box
Associated diseases	metabolic syndrome \Box , diabetes mellitus \Box , dyslipidemia \Box , renal artery disease \Box ,
	aortic diseases □
	ischemic stroke \square $^{(A)}$, hemorrhagic stroke \square $^{(A)}$; AMI \square $^{(A)}$, APE \square $^{(A)}$,
Target organs *	angina \square $^{(A)}$, CHF \square $^{(A)}$, renal disease \square $^{(A)}$, , PAD \square $^{(A)}$, AHTN crisis \square $^{(A)}$
AH pregnancy	preeclampsia \Box , eclampsia \Box , fetal losses \Box
Chronic diseases	
Long life therapeutics	
Diagnostic Exams	
Diagnostic Exams	
	, urea, creatinine, uric acid, Na, K, Cl, Ca, P,
	DL, LDL, APOB, VLDL, protein /albumin, vit.D, ACTH, T4, PTH, renin, angiotensin, aldoesterone
	, 14, 14, blood glomerular flitration rate,
	(serum)
- urea, creatinin, uric acid	d, urine II, Na, K, Cl, Ca, P
	, urine metanephrines 24h
- Echocardiography (alterations)	
- Radiography of chest (alterations)	
- Ultrasound (alterations)	
- Doppler ultrasound of carotid (alteration	is)
- Doppler ultrasound of lower limb art	eries (alterations)
- Ambulatory blood pressure (alterations)	
- Cardiac exercise stress test (alterations)	
- TAC/Angio-TAC	
- Arterial pulse wave velocity (alterations)	
- Others	

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^{*} age (A), acute myocardial infarction (AMI), congestive heart failure (CHF), acute pulmonary edema (APE), peripheral artery disease (PAD).



Physician signature ___

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FULL PANEL

	Name:
	Consultancy Referral Number:
ANNEX	
collection tube DNA (Date obtained: / ; Conditions: minimum 3 Saliva (Recommended kit: Oragene	btained: / /), Conditions: 4mL or 2 X 3mL in K ₂ EDTA or K ₃ EDTA /); VolumeμL; Concentration μg/mL; Purification Method 300ng of 35ng/μL,
for the genetic test specified in this req	/ my child 's [name] biological sample quest. I declare that I have been informed about genetic testing features and that I of the cardiovascular genetic test regarding genetic analysis of molecular risk markers giving permission.
, , , , , , , , , , , , , , , , , , , ,	processing of the obtained digital data: yes \square no \square imen and clinical information to be anonymously used in research studies: yes \square no \square
	;// 20 Signature

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