

### Rasha 123

20/08/2025 05:56:18

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Primary registry identifying number

LBCTR2018090151

MOH registration number

20555/2018

Study registered at the country of origin

Yes

Type of registration

Prospective

Date of registration in national regulatory

agency 15/09/2018

**Primary sponsor** 

Rasha

Date of registration in primary registry

22/04/2020

**Public title** 

Rasha 123

Scientific title

Rasha 123

Brief summary of the study: English

**Great Job** 

Brief summary of the study: Arabic

السجل الوطني للابحاث السريرية الاول من نوعه

Health conditions/problem studied: Specify

Heart Failure

Interventions: Specify

New medication

Key inclusion and exclusion criteria: Inclusion criteria

all adults

Key inclusion and exclusion criteria: Gender

Key inclusion and exclusion criteria: Age minimum

**Protocol number** 

Rasha123

Study registered at the country of origin: Specify

Type of registration: Justify

N/A

Primary sponsor: Country of origin

Lebanon

Date of registration in national regulatory agency

15/09/2018

Acronym

RH

Acronym

RH

Key inclusion and exclusion criteria: Specify gender

Key inclusion and exclusion criteria: Age maximum

75

Key inclusion and exclusion criteria: Exclusion criteria

less than 18

Type of study

Observational

Type of intervention

Pharmaceutical

Trial scope

Therapy

Study design: Allocation

Non-randomized controlled trial

Study design: Control

Uncontrolled

Study design: Purpose

Treatment

Study design: Assignment

Other

IMP has market authorization

No

Name of IMP

In Love

Type of IMP

Others

Pharmaceutical class

heart diseases

Therapeutic indication

to treat heart failure

Therapeutic benefit

increase ejection fraction

Study model Case-Control

Study model: Specify model

N/A

Time perspective

Prospective

Time perspective: Specify perspective

N/A

Type of intervention: Specify type

pharmaceutical with behavioral

Trial scope: Specify scope

N/A

Study design: Masking

N/A

Study phase

1 to 2

Study design: Specify purpose

Study design: Specify assignment

single arm

IMP has market authorization: Specify

USA

Year of authorization Month of authorization

2014

Study model: Explain model

OK

Time perspective: Explain time perspective



OK

Months

Target follow-up duration

30

Number of groups/cohorts

4

Biospecimen retention

Samples without DNA

Biospecimen description

Target follow-up duration: Unit

DNA saliva samples

Target sample size

1000

Date of first enrollment: Type

Anticipated

Date of study closure: Type

Anticipated

Recruitment status

Complete

Date of completion

24/04/2020

IPD sharing statement plan

No

Actual enrollment target size

1

Date of first enrollment: Date

20/10/2018

Date of study closure: Date

20/10/2020

**Recruitment status: Specify** 

IPD sharing statement description

I do not want to share

Additional data URL

later

**Admin comments** 

**Trial status** 

Approved



Secondary Identifying Numbers
No Numbers
Sources of Monetary or Material Support
No Sources
Secondary Sponsors
No Sponsors
Contact for Dublic/Scientific Queries
Contact for Public/Scientific Queries  No Contacts
The Contacts
Centers/Hospitals Involved in the Study
No Centers/Hospitals
Ethics Review
No Reviews
<u> </u>
Countries of Recruitment
No Countries



Health Conditions or Problems Studied	
o Problems Studied	
Interventions	
o Interventions	
Primary Outcomes	
o Outcomes	
Outcomes	
Key Secondary Outcomes	
o Outcomes	



Trial Results				
Summary results				
Study results globally				
Date of posting of results summaries	Date of first journal publication of results			
Results URL link				
Baseline characteristics				
Participant flow				
Adverse events				
Outcome measures				
URL to protocol files				