

Rasha 123

20/08/2025 08:56:05

ain Information	
Primary registry identifying number	Protocol number
_BCTR2018090151	Rasha123
MOH registration number 20555/2018	
Study registered at the country of origin Yes	Study registered at the country of origin: Specify
Type of registration	Type of registration: Justify
Prospective	N/A
Date of registration in national regulatory agency 15/09/2018	
Primary sponsor	Primary sponsor: Country of origin
Rasha	Lebanon
Date of registration in primary registry	Date of registration in national regulatory agency
16/04/2020	15/09/2018
Public title	Acronym
Rasha 123	RH
Scientific title	Acronym
Rasha 123	RH
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: Specify gender
Both	
Key inclusion and exclusion criteria: Age minimum	Key inclusion and exclusion criteria: Age maximum
18	75

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Key inclusion and exclusion criteria: Exclusion criteria less than 18

Type of study Interventional

Type of intervention	Type of intervention: Specify	type
Pharmaceutical	pharmaceutical with behavioral	
Trial scope	Trial scope: Specify scope	
Therapy	N/A	
Study design: Allocation	Study design: Masking	
Non-randomized controlled trial	N/A	
Study design: Control	Study phase	
Uncontrolled	Study phase 1 to 2	
Study design: Purpose	Study design: Specify purpos	e
Treatment	N/A	
Study design: Assignment	Study design: Specify assign	ment
Other	single arm	
IMP has market authorization	IMP has market authorization	: Specify
No	USA	
Name of IMP	Year of authorization	Month of authorization
In Love	2014	4
Type of IMP Others		
Pharmaceutical class		
heart diseases		
Therapeutic indication		
to treat heart failure		
Therapeutic benefit		
increase ejection fraction		
Study model	Study model: Explain model	
N/A	N/A	
Study model: Specify model N/A		
N/74		
Time perspective	Time perspective: Explain tim	e perspective
N/A		
Time perspective: Specify perspective		
N/A		

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	N/A
Target follow-up duration	Target follow-up duration: Unit
Number of groups/cohorts	
Biospecimen retention	Biospecimen description
Samples without DNA	DNA saliva samples
Target sample size	Actual enrollment target size
1000	Actual enronment target size
Date of first enrollment: Type	Date of first enrollment: Date
Anticipated	20/10/2018
Date of study closure: Type	Date of study closure: Date
Anticipated	20/10/2020
Recruitment status	Recruitment status: Specify
Complete	
Date of completion	
IPD sharing statement plan No	IPD sharing statement description I do not want to share
	r do not want to snare
Additional data URL	
Admin comments	
Trial status	
Approved	



Secondary Identifying Numbers		
Full name of issuing authority	Secondary identifying number	
KSA	25325	

Sources of Monetary or Material Support

Name

King faisal hospital

Secondary Sponsors

Name

None

Contact for Public/Scientific Queries						
Contact type	Contact full name	Address	Country	Telephone	Email	Affiliation
Public	Rasha Hamra	Beirut	Iceland	0255.0000	rasha@gmail	МОН
Scientific	Marie Ok	Beirut	Lebanon	565540000 0	marie@yahoo	МОН

Centers/Hospitals Involved in the Study			
Center/Hospital name	Name of principles investigator	Principles investigator speciality	Ethical approval
AUB	RH	Oncologist	Approved
USJ	RH	Oncologist	NA

Ethics Review				
Ethics approval obtained	Approval date	Contact name	Contact email	Contact phone
American University of Beirut Medical Center King faisal hospital	11/10/2018	RH	@gmail	565458
Makassed General Hospital	11/12/2019	RH	@yahoo	3336661

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Countries of Recruitment

No Countries

Health Conditions or Problems Studied		
Condition Code Keyword		Keyword
heart failure	Heart failure (I50)	RF

Interventions		
Intervention	Description	Keyword
pharmaceutical and behavioral	new medication with lifestyle changes	RH and forever

Primary Outcomes		
Name	Time Points	Measure
increast EF	in 6 months	EF

Key Secondary Outcomes		
Name	Time Points	Measure
I dont have	later	Ok



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