

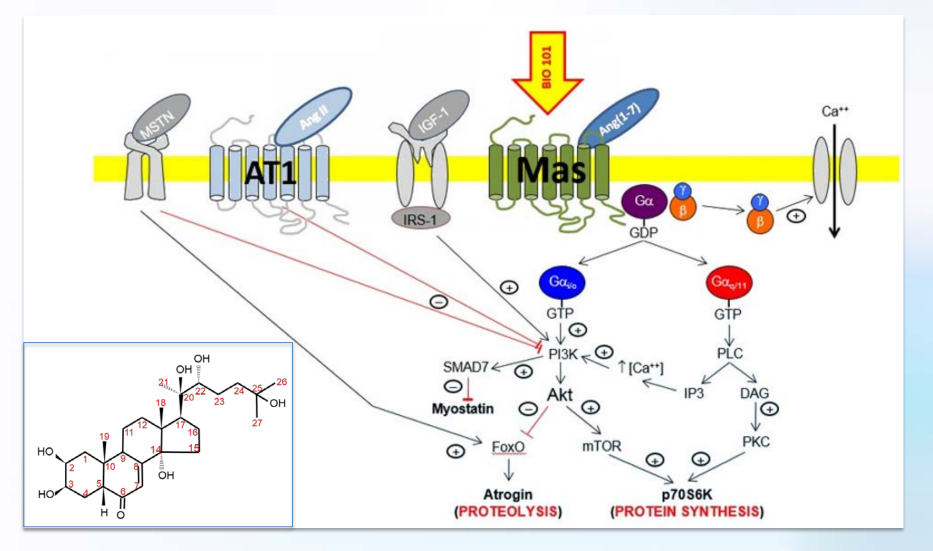
SARA PROGRAM: PRELIMINARY FINDINGS & IMPLICATIONS FROM SARA-OBS STUDY AND ITS IMPACT ON SARA-INT STUDY

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biophytis

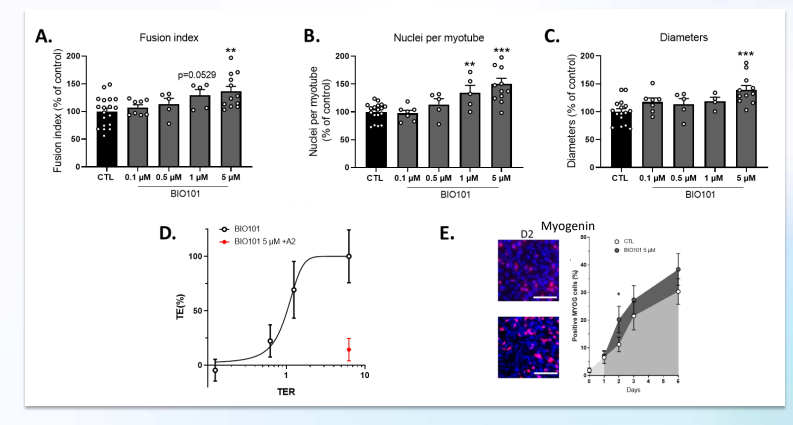
BIO101: MECHANISM OF ACTION

MAS receptor activation





Myoblast differentiation

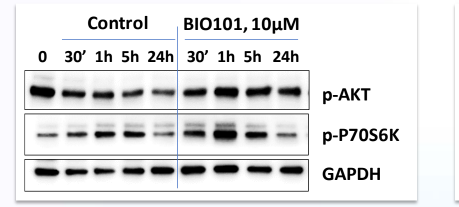


⇒ Dose-dependent effect of BIO101 on myoblast differentiation, based on effects observed on fusion index, number of nuclei per myotube and myotube diameter, as well as Myogenin protein expression.

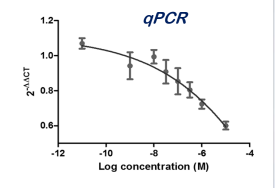


TARGET ENGAGEMENT STUDIES IN-VITRO

AKT signaling pathway

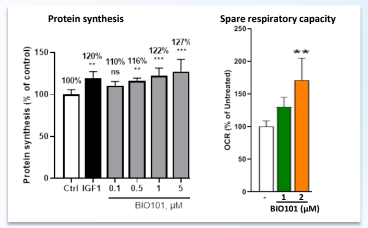


Myostatin gene expression



=> BIO101 stimulates AKT signaling pathway and inhibits myostatin gene expression

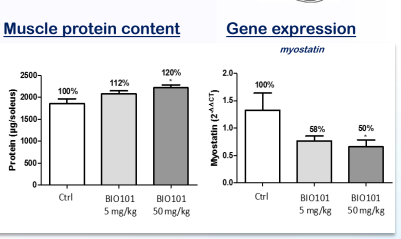
Myoblast protein synthesis and mitochondrial respiration



- ⇒ BIO101 is responsible for a significant increase in protein synthesis in differentiated muscle cells.
 ⇒ BIO101 increases significantly the
 - mitochondrial spare respiratory capacity in differentiated C2C12

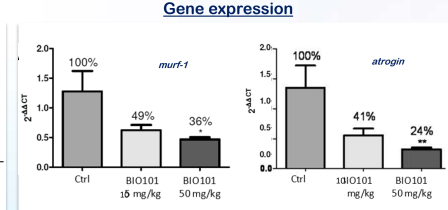
TARGET ENGAGEMENT STUDIES IN-VIVO

Mouse soleus, 6 weeks



Rat tibialis anterior, 12 weeks oral treatment





BIO101 causes an increase in muscle protein content in treated mice.

BIO101 administration cause a partial inhibition of myostatin gene expression.

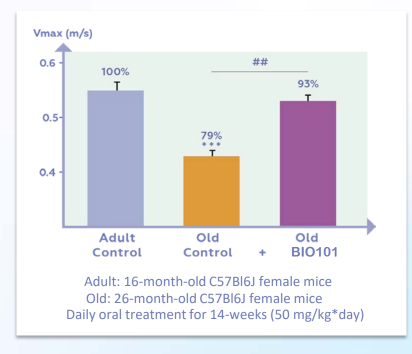
Chronic BIO101 treatment causes a dosedependent gene expression inhibition of two important regulators of ubiquitinmediated protein degradation in skeletal muscle (Murf-1 and Atrogin).

=> BIO101 has a dose-dependent effect in the target tissue with a chronic treatment



PROOF OF CONCEPT

Age-related muscle wasting and loss of function



 \Rightarrow Significant protection against muscle function loss during ageing

THE SARA CLINICAL PROGRAM



SARA = SARcopenia and sarcopenic obesity in patients Aged ≥ 65 years

SARA-PK

- Phase 1 study
- Safety and pharmacokinetic profiles of BIO101 in older adults
- Selection of doses for SARA-INT

SARA-OBS

 An observational study to characterize the target population and main parameters of SARA-INT

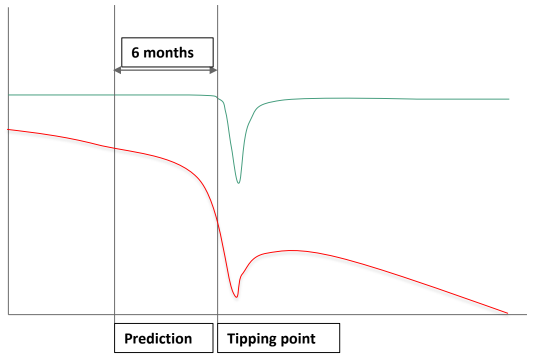
SARA-INT

• A phase 2 interventional trial to evaluate the safety and efficacy of BIO101 after 6-month administration in sarcopenic patients on mobility function

SARA PROGRAM: IDENTIFICATION OF THE TARGET POPULATION

Aiming to identify individuals who are at a high risk for mobility disability

- In standard care, individuals with a chronic disease are often diagnosed when a bad thing happens. This is the **Tipping Point**
- The tipping point is an untoward medical event, from which the affected individual is not able to recover to their pre-event function
- Resilience indicates the ability of individuals to recover from untoward events. The opposite of resilience is frailty
- Sarcopenia is a significant cause for increased frailty and losing the ability to complete the 400 meter gait task.
 400MW test is a measure that is associated with loss of resilience.



Exclusion of individuals with SPPB = 9 is the main difference between the SARA program and previous sarcopenia studies

SARA-OBS: AN OBSERVATIONAL CLINICAL TRIAL

Characterizing Sarcopenia and sarcopenic obesity in patients Aged 65 years and over, at risk of mobility disability

Objectives:

- To characterize sarcopenia, including sarcopenic obesity, in older patients (<u>>65</u> years) living in the community and at risk of mobility disability
- Evaluate physical performance and body composition for the design of a phase 2 interventional study on the efficacy and safety of BIO101
- Estimate the relative prevalence and recruitment rate in sarcopenia

Main endpoints:

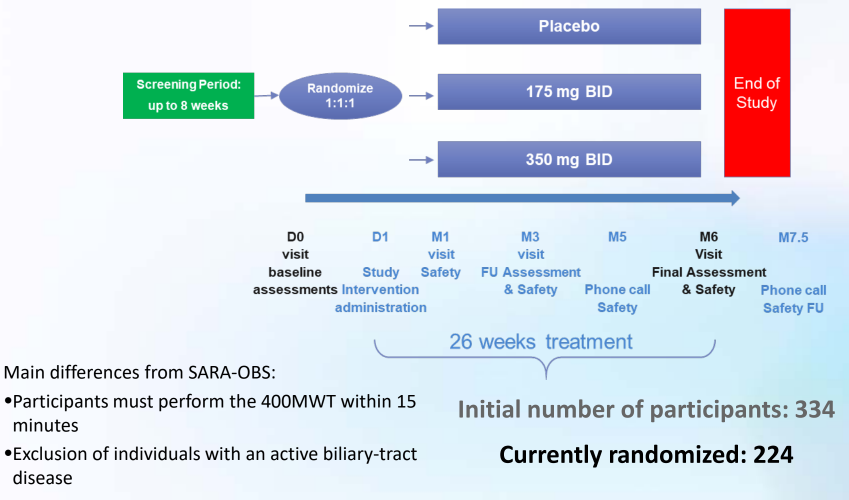
- Primary endpoint: gait speed of the 400MW test
- Co-primary endpoint: The Physical Function Domain (PF-10) of the Short Form Health Survey (SF-36)
- Key secondary endpoint: Hand grip strength
- Other secondary endpoints: 6 MWD, completion of 400MW Test, SPPB, SPPB sub-score of sit-stand, stair climbing Power Test, PROs (SF36; SarQoL; TSD-OC) knee extension strength, DXA measurements
- Safety and tolerability

Main inclusion criteria:

- Men and women aged ≥ 65 years, living in the community, and reporting loss of physical function
- Short Physical Performance Battery (SPPB) score ≤ 8
- ALM/BMI < 0.789 in men and 0.512 in women, or ALM <19.75 kg in men and <15.02 kg in women by DXA

SARA-INT: A PHASE 2 INTERVENTIONAL STUDY

Following the same outline as that of the SARA-OBS study





WHAT WE LEARNED FROM SARA-OBS (PRELIMINARY ANALYSIS ON 106 COMPLETERS)

Disease progression and its impact on the sample size

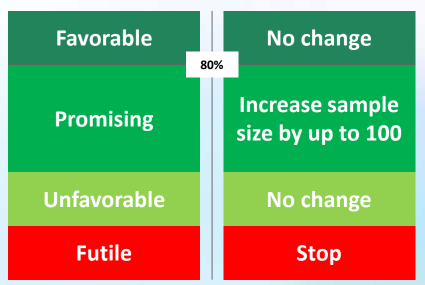
		E	SL	M6	Change	Р	
400WT (0.	90	0.84	-0.05	0.0036	
SPPB score 6		6.	67	6.99	0.32	0.0915	
6MWT 2		298	3.84	279.56	-19.28	0.0012	
Chair-stand		19.63		20.04	0.41	0.61	
Handgrip 2		22	.73	22.29	-0.43	0.36	
Effect size assumption: 0.05m/sec above baseline							
Std div.	Expected diff from placebo		N / arr	n N tota	Expected change from BL (%)		
	0.05		253	759	5.5%		
	0.08				8.9%		
0.20	0.08		111	334	8.9	%	

64

0.10

A 'promising zone' interim analysis

- Re-confirm the statistical power
- Re-adjust the sample size if needed



Population of the 2 studies have similar baseline characteristics

192

⇒ Exclusion of SPPB=9 led to the selection of a more vulnerable population, which is closer to the 'tipping point'

11.0%

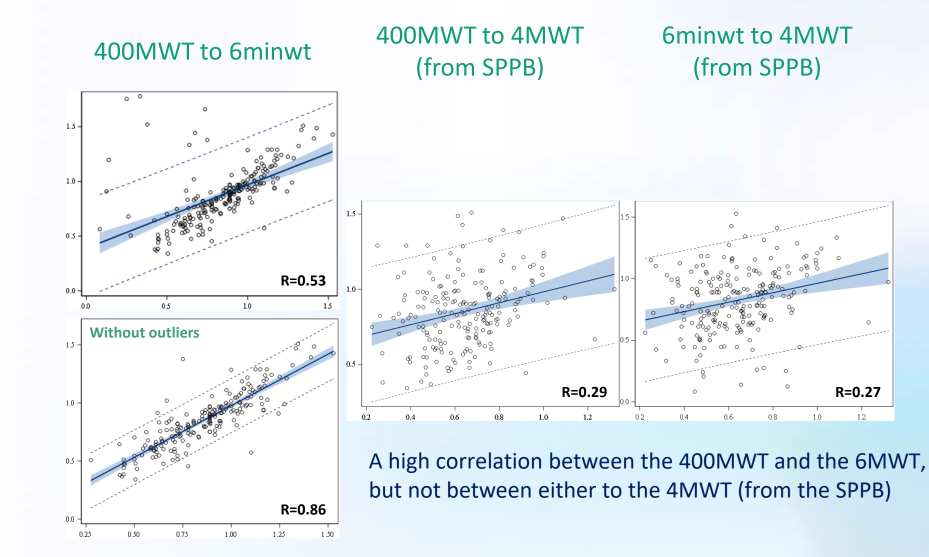
 \Rightarrow Sample size re-estimation to 192 participants (231 including 20% premature termination).

CORRELATIONS



R=0.27

1.2





For more details on SARA-OBS study, please come and visit our poster **CLINICAL TRIALS**

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Design and Population Baseline Features of the SARA-OBS Study: Characterizing SARcopenia and Sarcopenic Obesity in Patients Aged65 years and over, at Risk of Mobility Disability

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