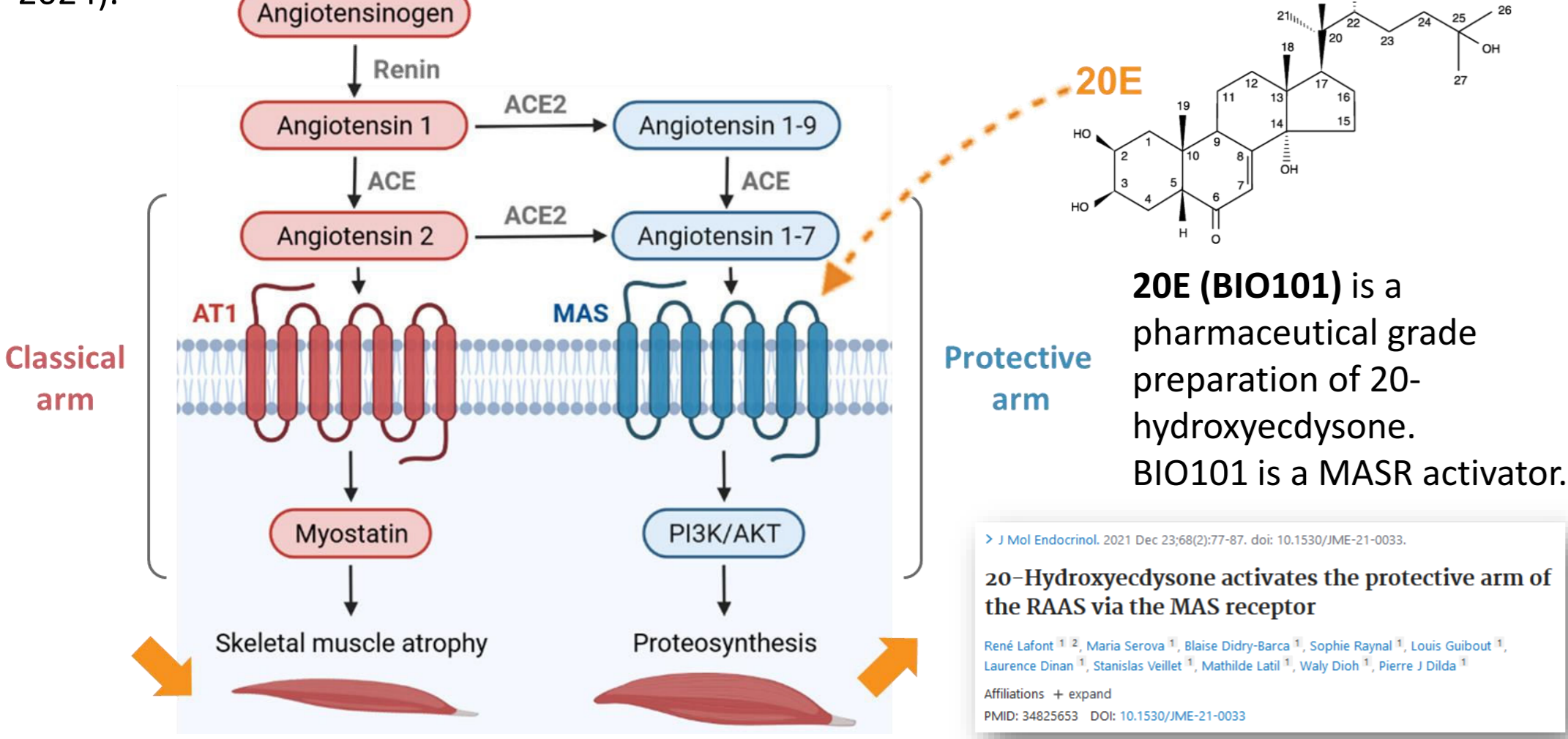


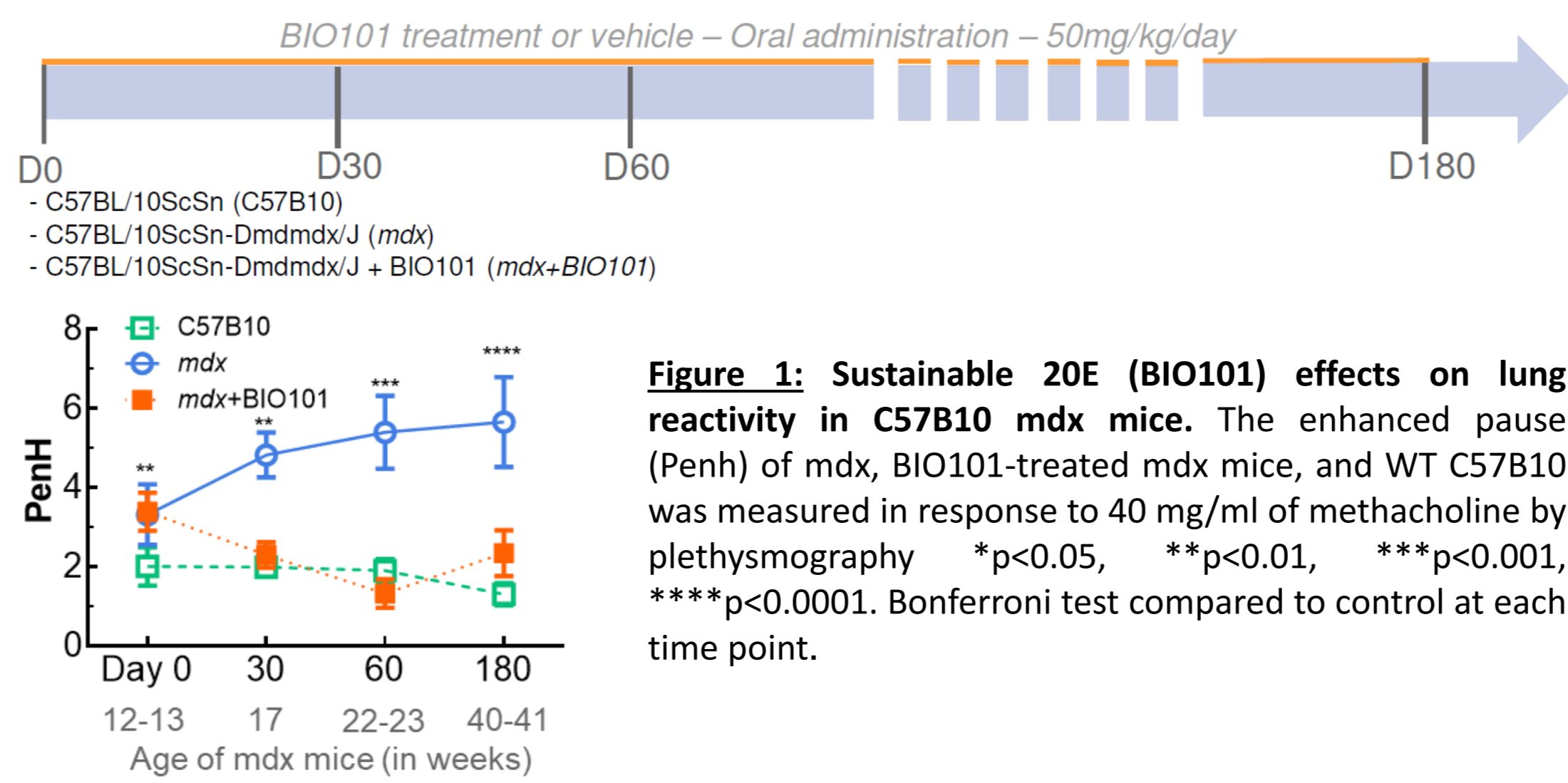
## 20-Hydroxyecdysone (20E-BIO101)

20E (BIO101) is a new oral drug candidate activating the MAS receptor (a major receptor in the renin-angiotensin system) leading to muscle anabolism, which already demonstrated meaningful activity in animal models of muscular dystrophies (Lafont et al., 2021) and respiratory function. Furthermore, 20E (BIO101) has also demonstrated beneficial effects in two vulnerable populations in clinical settings, in the COVA (Lobo et al., 2024) and SARA-INT trials (Dih et al., 2024).

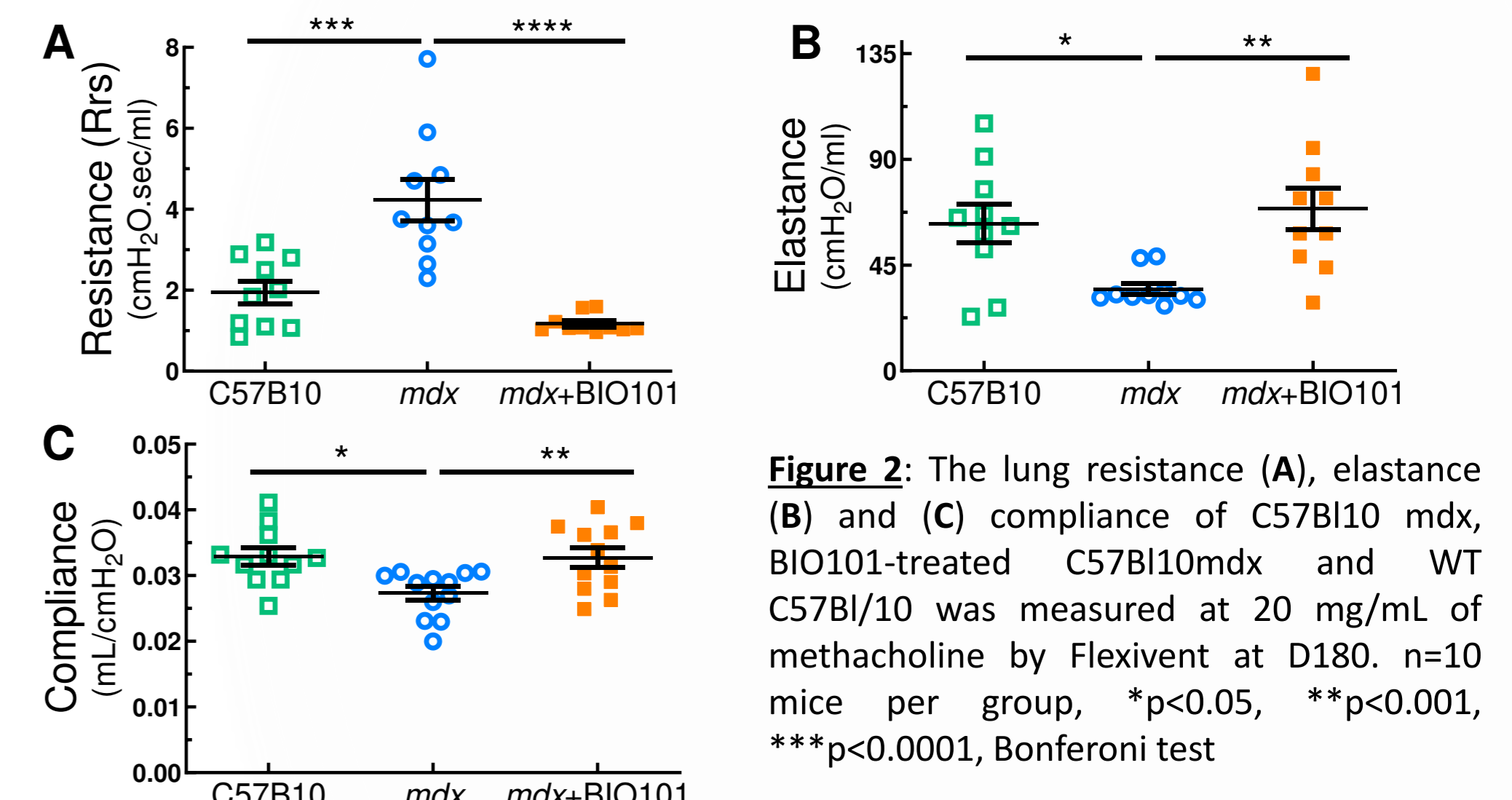


## Duchenne Muscular Dystrophy preclinical data on respiratory function in mdx mice

### - Whole body Plethysmography



### - Flexivent



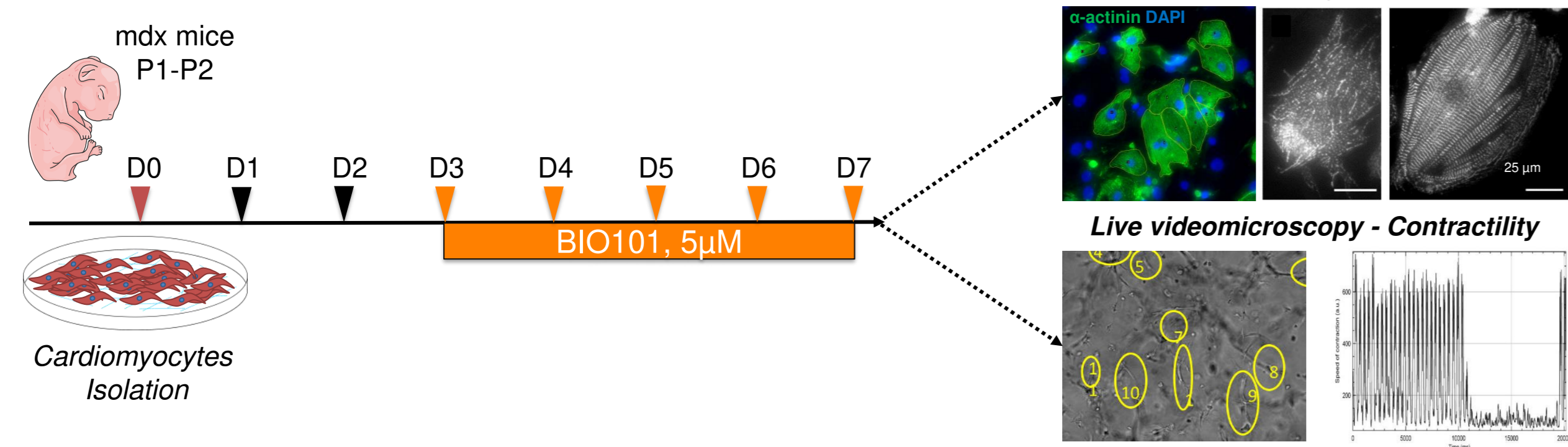
➤ **BIO101 treatment significantly improves airway responsiveness as measured by PenH (normalization of Penh values vs control mice).**

➤ **BIO101 effects are sustained for 6 months.**

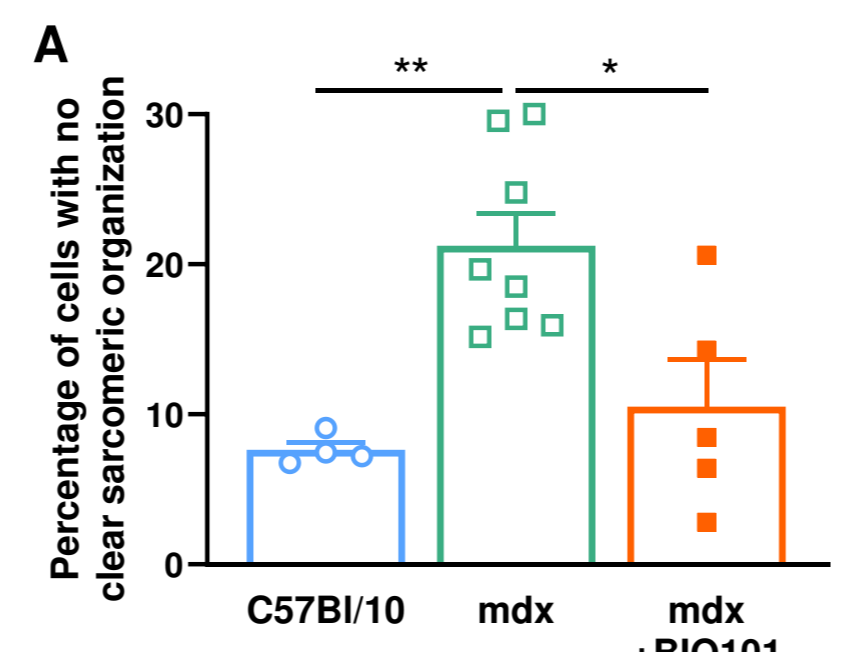
➤ **BIO101 treatment normalized C57BL/10 mdx mice lung mechanical properties such as resistance, elastance and compliance.**

## Duchenne Muscular Dystrophy preclinical data on neonatal mdx-derived cardiomyocytes phenotype and function

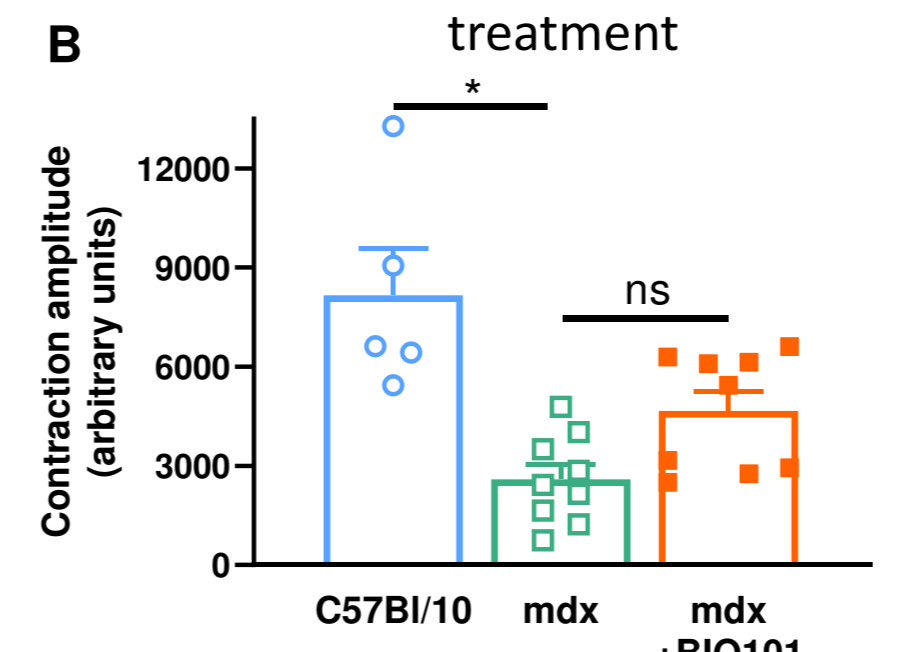
### Cardiomyocytes cultures



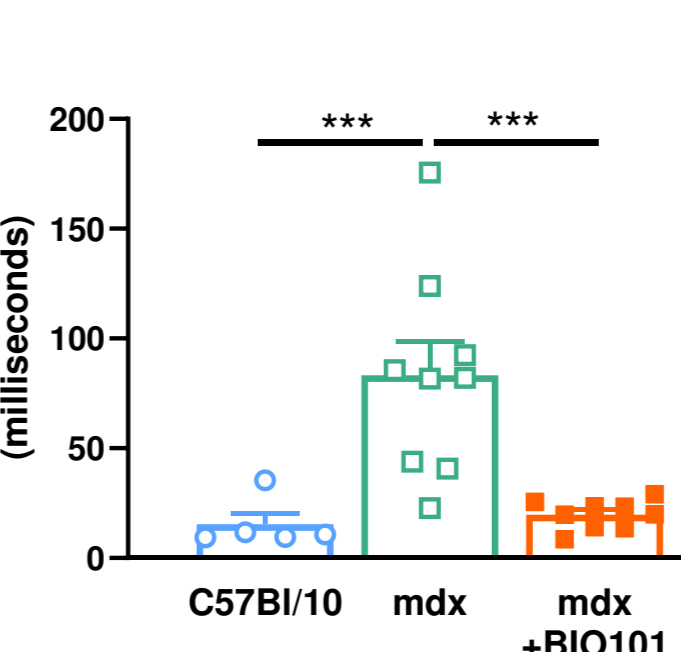
✓ Sarcomeres organization is improved with BIO101 treatment



✓ Contraction amplitude is not significantly improved with BIO101 treatment



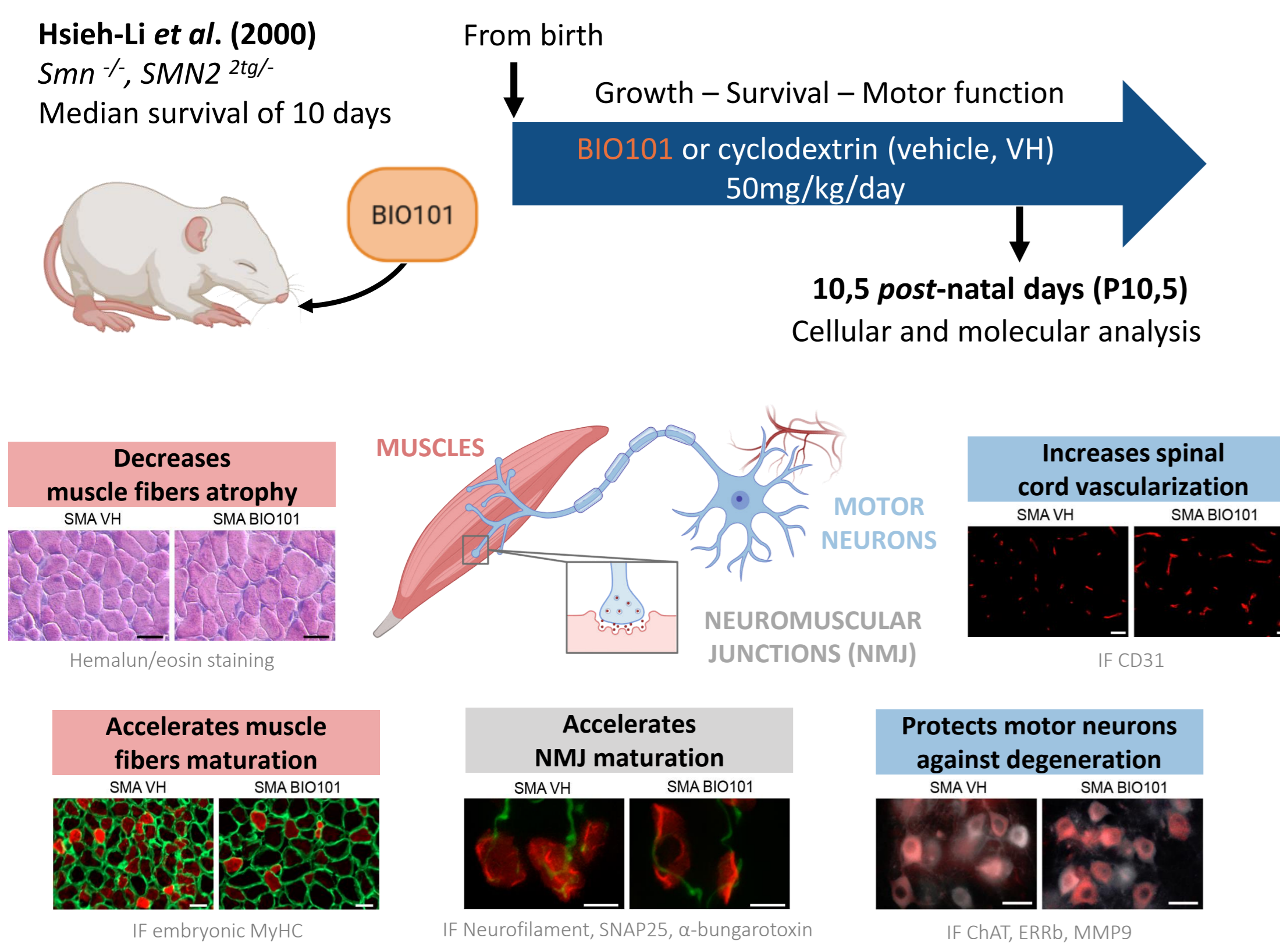
✓ Arrhythmias are abolished with BIO101 treatment



➤ **BIO101 treatment significantly improves mdx cardiomyocytes functionality**

## Spinal Muscular Atrophy preclinical data

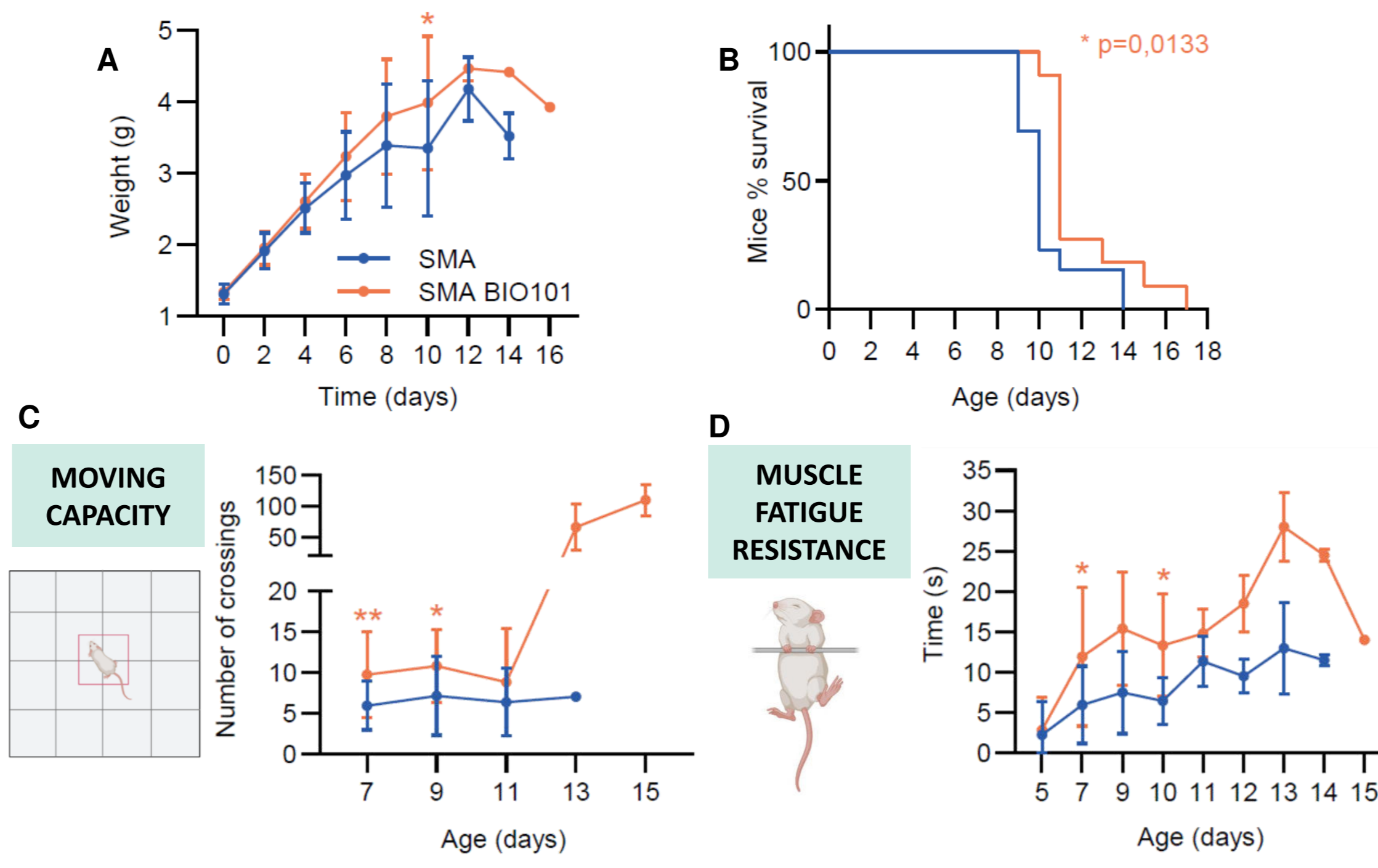
### BIO101 monotherapy in severe SMA-like mice



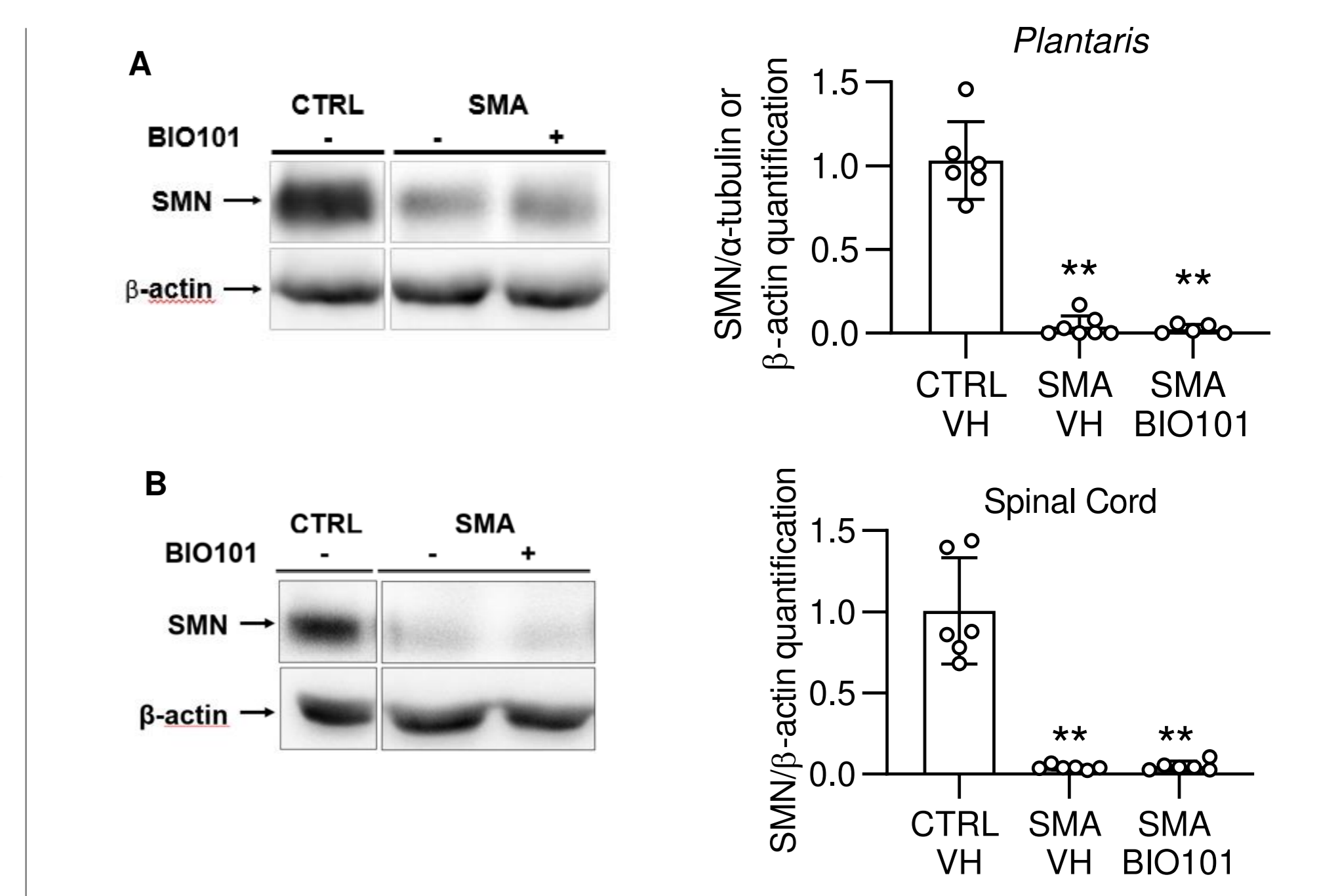
**Figure 4: Illustration of the different beneficial effects of BIO101 monotherapy in severe SMA-like mice on the entire motor unit (muscles, motor neurons and neuromuscular junctions)**

➤ **BIO101 induces beneficial effects on the entire motor unit and delays the weight loss, increases survival, improves moving capacity and muscle fatigue resistance in severe SMA-like mice. BIO101 does not increase SMN protein expression in the plantaris and in the lumbar spinal cord of SMA-like mice. Similar results are found in the tibialis and the soleus (not shown).**

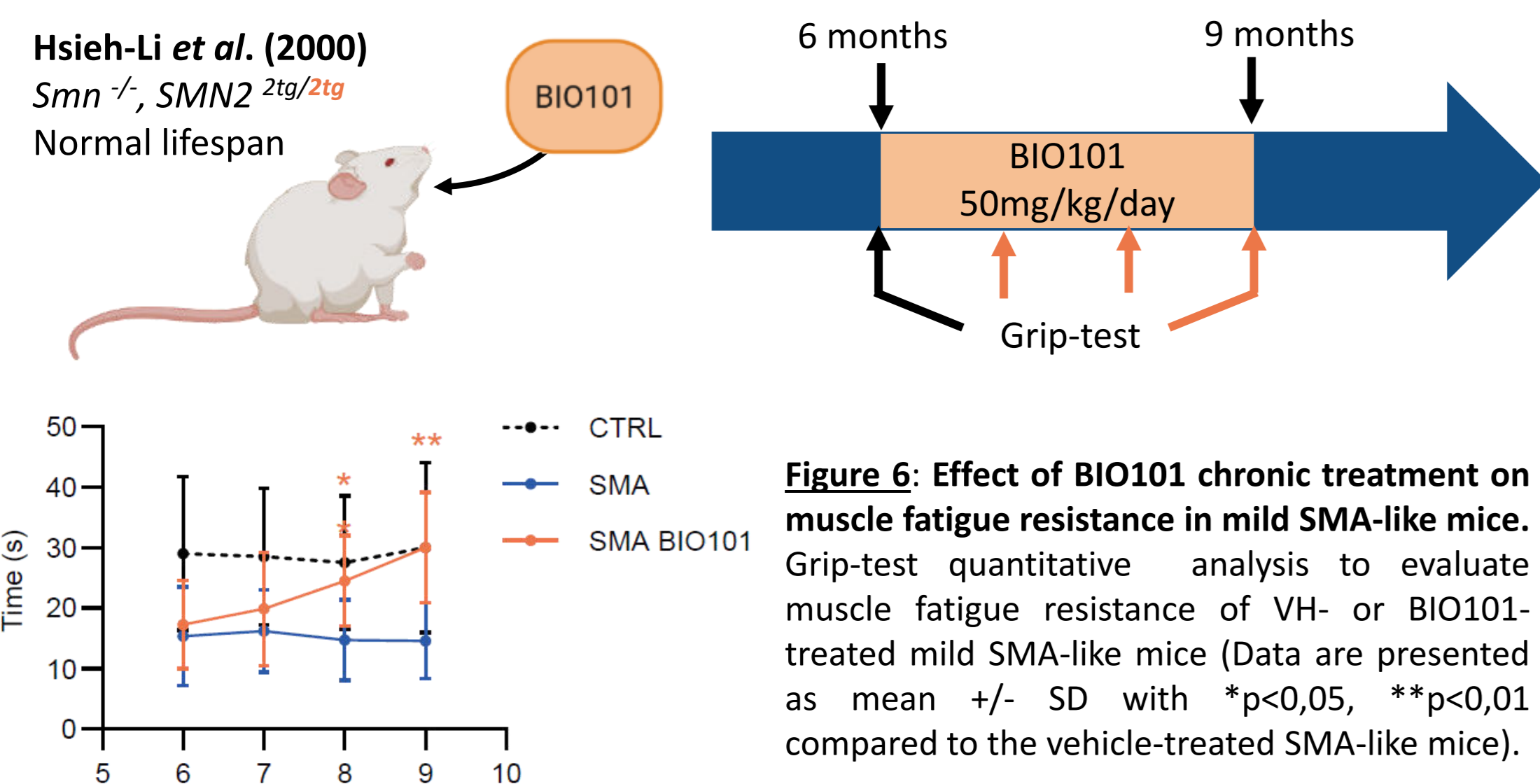
### - Effects on growth, survival and motor function



### - Effects on SMN protein expression in skeletal muscles and spinal cord

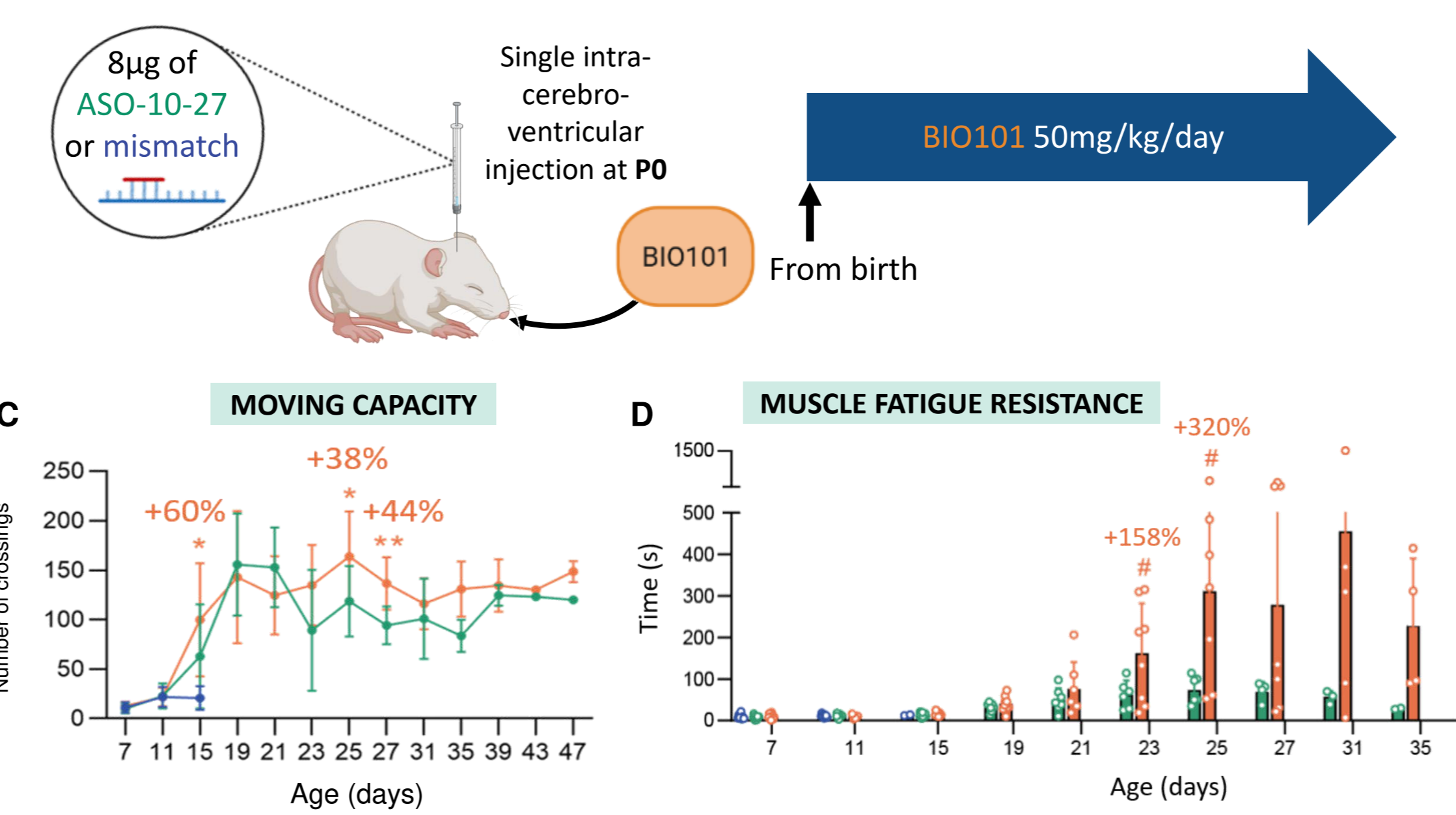


### BIO101 monotherapy in mild SMA-like mice



➤ **BIO101 restores muscle fatigue resistance of symptomatic mild SMA-like mice to control level after 3 months of treatment.**

### Combinatorial therapy with BIO101 and ASO-10-27 in severe SMA-like mice



➤ **BIO101 increases lifespan, improves growth, enhances moving capacity and resistance to muscle fatigue in SMA-like mice treated with ASO-10-27.**

## Conclusion

- **20E (BIO101) is a promising oral treatment for DMD patients with respiratory deterioration and SMA patients.**
- Favorable safety profile (SARA-PK phase 1, good safety data on 149 SARA-INT participants with at least 6 Months of dosing)
- Beneficial effects on motor function in sarcopenic patients (SARA-INT phase 2b) and beneficial effects on COVID-19 patients with respiratory failure (COVA phase 2/3)
- ODD granted in Europe and US, Biophytis intends to start the Phase 1/2 MYODA clinical trial in the upcoming months (see Poster P360).
- For SMA patients, alone or in combination with approved therapies, BIO101 may improve survival, growth and motor function, especially resistance to muscle fatigue which plays a key role in the quality of life of SMA patients.