# **Characterization of pre-existing immunity to AAV5**

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## **Objective**

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- The objective of this study was to further characterize pre-existing antibody responses to AAV serotype 5 (AAV5) in severe hemophilia A patients (n=540) who are naïve to gene therapy. Characteristics evaluated included:
- AAV5 neutralizing factors (transduction inhibition, (TI) using a cell-based assay
- AAV5 total binding antibodies (TAb) using a bridging immunoassay
- AAV serotype specificity using a bridging immunoassay
- AAV5 in vitro complement activation
- AAV5 antibody isotypes

### **Methods**

### Assay methodologies for assessing AAV5 TI and TAb pre-existing immunity

### **Global seroprevalence of AAV varies by serotype**

We previously assessed the seroprevalence of pre-existing immunity to AAV5 and four additional AAV serotypes<sup>3</sup>: AAV2, AAV6, AAV8, and AAVrh10



### **Complement activation to AAV5 >200% in vitro** was observed only in subjects that were positive for all five serotypes (AAV5/2/6/8/rh10)



### **AAVx Positivity (out of 5 total)**

**Complement Activation** 

**Cell-based Transduction Inhibition (TI) Assay** 



- Measures ability of plasma to block transduction of a cell line by AAV
- Differing amounts of capsid needed for different serotypes can make TI titer comparisons across capsids more difficult

### Anti-AAV Total Antibody (TAb) Assay



- Clinical TAb "Capsid Bridging" Non-clinical TAb Ru-Protein A/G/L
- Electrochemiluminescent assay (ECLA) on the MSD platform
- Ease of qualifying divergent capsid serotype assays in identical format

### broader AAV serotype cross-reactivity

We assigned subjects from the global seroprevalence study to AAVx positivity groups (Negative, 1, 2, 3, 4, 5) based on the number of serotypes for which they were positive

Higher AAV5 TAb titers were not correlated with





There was no significant association between higher AAV5 TAb titers and the presence of antibodies reactive with other serotypes, suggesting that AAV5-specific TAb titers were not a result of broad previous AAV exposures (and subsequent cross-reactivity)



		(wean % increase)				
AAVx Positivity	N	Bb	C5a	C3a	C4a	Average
Negative	9	102%	98%	96%	115%	103%
1	9	101%	104%	99%	109%	103%
2	6	110%	105%	121%	155%	123%
3	0	_	_	_	_	_
4	5	105%	114%	106%	131%	114%
5	14	260%	257%	271%	407%	299%

C3a: central point of activation Bb: alternative pathway activation

C4a: classical/lectin pathway activation C5a: terminal pathway activation

### AAV5 TAb responses were predominantly the IgG Isotype

- In a subset of subjects (n=78) with pre-existing AAV5 TAb positivity, we characterized the isotype and subtype of the anti-AAV5 response (IgM, IgG1, IgG2, IgG3, IgG4, IgA)
- Different isotypes reflect maturation of antibody response, and can have variable effector functions (neutralization and Fc receptor affinities)
- IgG represents a more affinity mature and likely neutralizing response
- IgM is the first to arise and reflects an immature response



### Results

### The majority of subjects with pre-existing immunity to AAV5 were positive for both TAb and TI<sup>1</sup>

- Majority of Hemophilia A subjects are negative or positive in both TAb and TI assays
- 62% TAb and TI negative
- 25% TAb and TI positive
- A small subset of subjects have discordant results
- 4% TAb+ and TI-
- 9% TAb- and TI+



#### **Pre-existing AAV5 TAb titers were much lower**

### **Some AAV5 TAb positive subjects demonstrated** complement activation in response to AAV5

AAV5-induced complement activation was assessed in vitro by measuring complement split products. Sera was incubated with either AAV5-bCG or control buffer



In some AAV5 TAb+ subjects, a >200% increase in activated complement was observed; however, this was not strictly correlated with the magnitude of AAV5 TAb titers as complement activation was not observed in all subjects with higher titers

N = 7 with IgA titers of 100-200, randomly distributed with respect to other isotypes

### Subjects that were AAV5 TAb+ and TI negative or had low TI titers were predominantly AAV5 IgG1 negative

- Higher AAV5 TAb titers (~ >100) and TI titers (~ >60) were typically associated with detection of IgG1, but higher AAV5 TAb titers that were TI negative or low TI titer ( $\sim$  <60) did not reveal a similar trend
- Consequently, highly neutralizing antibody responses were associated with detection of AAV5-specific IgG1



than AAV5 gene therapy post-treatment titers



#### References

1. Hayes et al. ISTH Presentation. 2019. 2. Greg Hayes, Global Seroprevalence of pre-existing immunity against various AAV serotypes in the hemophilia A population, ISTH 2019. 3. Klamroth et al. Human Gene Therapy. 2021.

#### **Disclosures**

Kelly Lau, Christian Vettermann, Greg de Hart, Benjamin Hock, Brian Long, and Soumi Gupta are employees of BioMarin Pharmaceutical Inc.

### Summary

- The presence of pre-existing AAV antibodies may impact the successful administration and efficacy of gene therapy
- The majority of subjects with pre-existing immunity to AAV5 were positive for both TAb and TI, with a small subset showing discordant results. Pre-existing AAV5 TAb titers were much lower than AAV5 gene therapy post-treatment titers
- There was no association between higher AAV5 TAb titers and the presence of antibodies reactive with other serotypes, suggesting that AAV5-specific TAb titers were not a result of broad previous AAV exposures (and subsequent cross-reactivity)
- In some AAV5 TAb+ subjects, a >200% increase in activated complement was observed; however, this was not strictly correlated with the magnitude of AAV5 TAb titers. Complement activation was not observed in TAb-negative subjects, and not all TAb-positive subjects exhibited complement activation
- Complement activation to AAV5 >200% in vitro was observed only in subjects that were positive for all five serotypes (AAV5/2/6/8/rh10)
- The isotype and subtype of the anti-AAV5 response (IgM, IgG1, IgG2, IgG3, IgG4, IgA) was predominantly the IgG isotype. Higher AAV5 TAb titers and TI titers were typically associated with detection of IgG1
- Understanding the most important immunological determinants of pre-existing immunity that impact efficacy could enable more patients to gain access to gene therapy