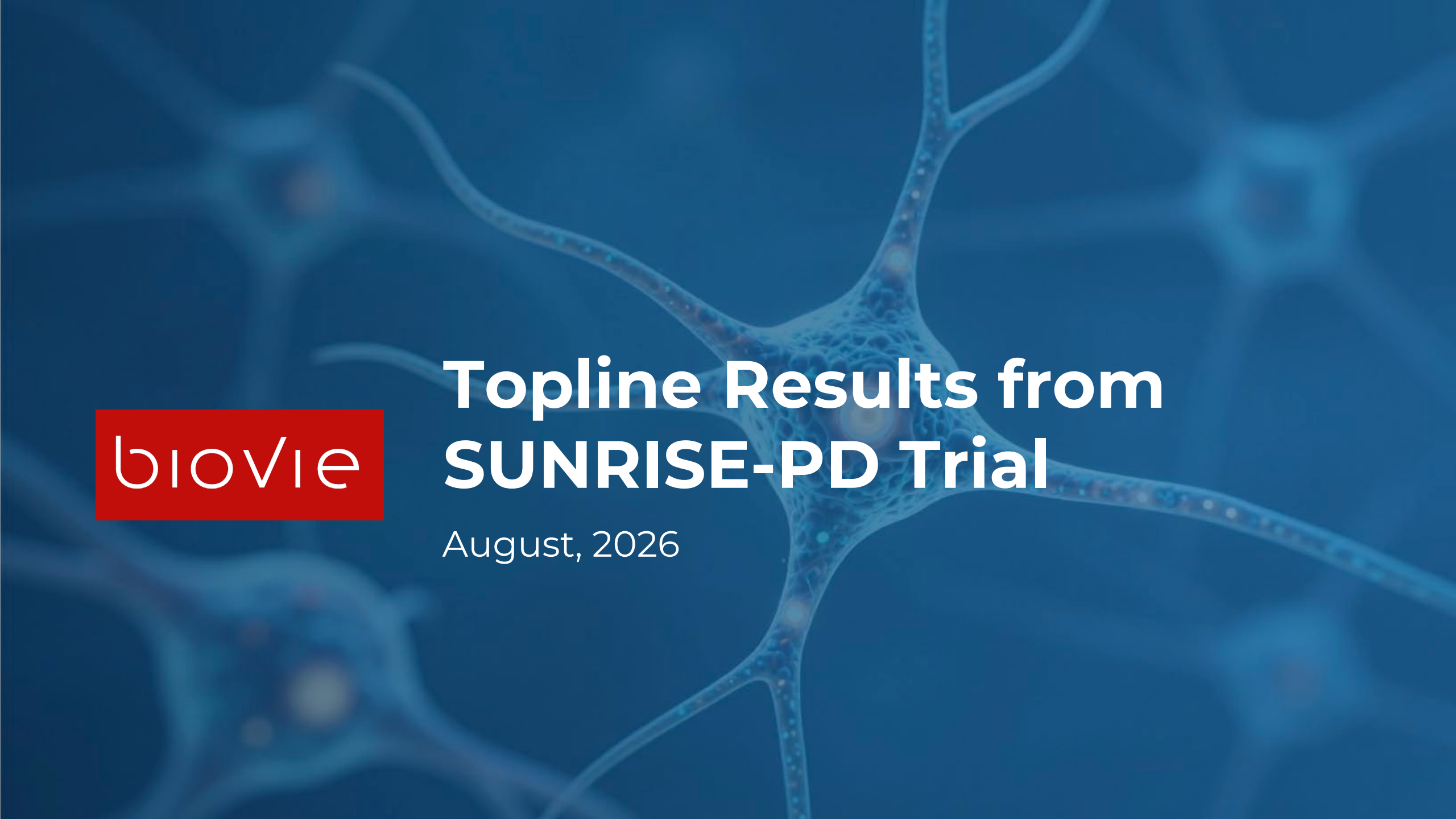


The background of the slide features a network of neurons, with a central neuron prominently displayed. The neurons are rendered in a light blue, semi-transparent style against a darker blue background, showing their cell bodies and branching processes.

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# Topline Results from **SUNRISE-PD Trial**

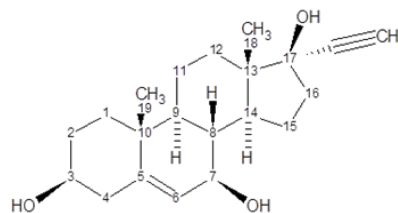
August, 2026

# FORWARD-LOOKING STATEMENTS

This presentation contains statements about BioVie's future expectations, plans, strategies and prospects which constitute forward-looking statements. These forward-looking statements are made pursuant to the safe harbor provisions of the Private Securities Litigation Reform Act of 1995. BioVie has in some cases identified forward-looking statements by using words such as "anticipates," "believes," "hopes," "estimates," "looks," "expects," "plans," "intends," "goal," "potential," "may," "suggest," and similar expressions. Forward-looking statements are subject to risks, uncertainties and assumptions that could cause BioVie's actual results and experience to differ materially from anticipated results and expectations expressed in these forward-looking statements. Among other factors that could cause actual results to differ materially from those expressed in forward-looking statements are: BioVie's ability to raise the substantial capital needed to fund its operations and research and development; risks associated with clinical development and BioVie's ability to successfully complete pre-clinical and clinical testing and be granted regulatory approval for its products to be sold and marketed in the United States or elsewhere; BioVie's reliance on third parties to conduct its clinical trials and manufacture its product candidates; BioVie's ability to establish and/or maintain intellectual property rights covering its product candidates; competition; and other risks described in greater detail in BioVie's filings with the Securities and Exchange Commission (the "SEC"). In addition to the risks described above and in BioVie's filings with the SEC, other unknown or unpredictable factors also could affect BioVie's results. No forward-looking statements can be guaranteed, and actual results may differ materially from such statements. You should not place undue reliance on any forward-looking statements. BioVie undertakes no obligation to release publicly the results of any revisions to any such forward-looking statements that may be made to reflect events or circumstances after the date that these slides are posted to BioVie's website or to reflect the occurrence of unanticipated events, except as required by applicable law or regulation.

# BEZISTERIM

Believed to block ERK/NFκB-mediated inflammatory signaling while preserving homeostatic function

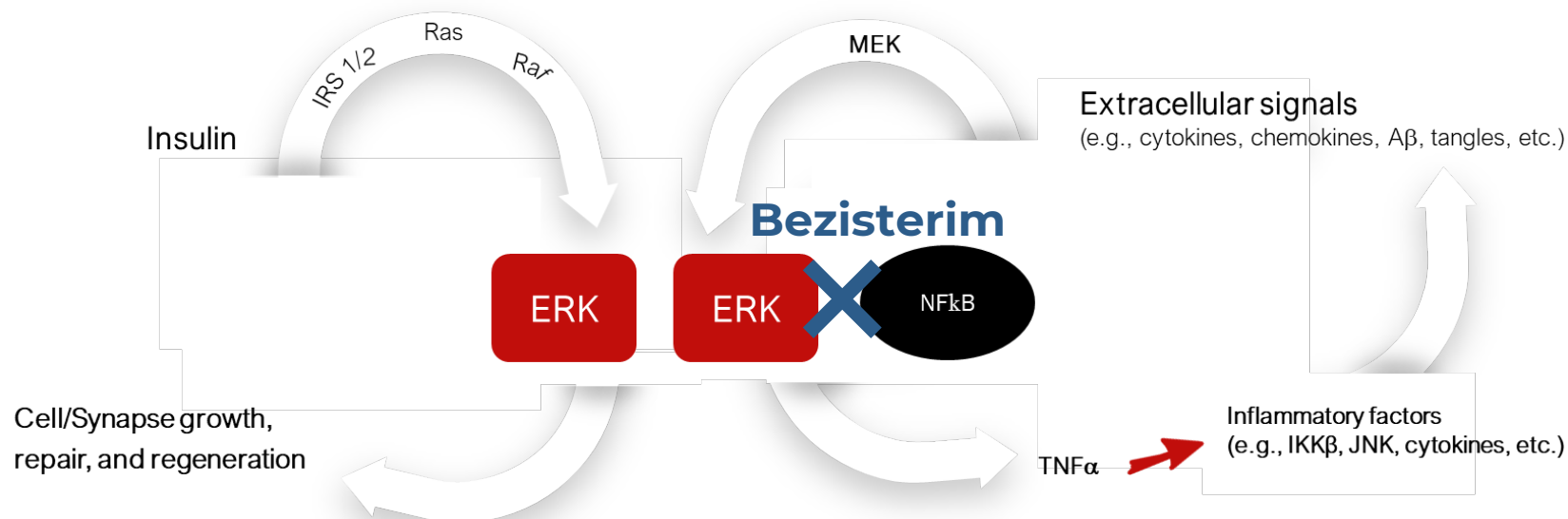


First-in-class molecule with desirable characteristics

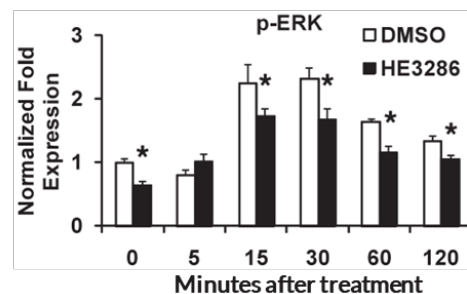
Small molecule; orally bioavailable

Crosses blood-brain barrier

No safety issues identified to date in pre-clinical and clinical trials (up to Phase 3)

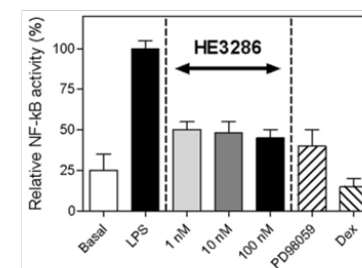


**Bezisterim Reduces ERK Activation**



Lu 2010 Am J Physiol Endocrinol Metab 298 E1036

**Bezisterim Reduces NFκB Activation**



Wang 2010 J Pharmacol Exp Ther 333 70

# SUNRISE-PD PHASE 2 TRIAL

- Designed as a **short**, signal finding trial with early-stage PD patients
  - Identify efficacy signals: endpoints with favorable treatment benefits
  - Measure the magnitude of treatment impact
  - Enable design and sizing of Phase 3 trial
- Evaluated a series of motor and non-motor endpoints
  - MDS-UPDRS Parts I (non-motor), II (activities of daily living), III (motor), and Total
  - PDSS-2 (sleep) and PDQ-39 (Quality of Life)
- Evaluated a series of biomarkers, including a proteomics battery evaluating 380 different proteins
  - Neurodegenerative (NfL, GFAP, pTau-217,  $\alpha\beta 42$ )
  - Neuroinflammation (CCL2, CHI3L1, VGF)
  - Peripheral inflammation (MLR, SIRI, IL-17A, IFN- $\gamma$ , TNF $\alpha$ , others)

# KEY FINDINGS

- Patients treated with bezisterim appeared to have experienced statistically significant treatment advantages compared to placebo
  - **Clinical outcomes:** composite of 15 motor and non-motor domains of PD
  - **Proteome-wide impact:** 283 / 380 proteins moving towards direction that has been shown to slow disease progression
  - **Neurodegeneration:** composite and individual biomarkers (NfL, GFAP, others)
  - **Neuroinflammation:** CCL2, CHI3L1, others
  - **Peripheral inflammation:** monocyte-to-lymphocyte ratio, SIRI, others
- Bezisterim appeared to demonstrate an effect across the trial population, and those with higher levels of inflammation at baseline experienced statistically significant advantages over placebo across all clinical measures evaluated, including UPDRS Parts I, II, and III, UPDRS Total, and the EPNIC-15 composite
- Biomarker results suggest that bezisterim may slow neurodegeneration over time – a hypothesis that we will test in the Phase 3 registrational trial
- Bezisterim was well-tolerated; similar to placebo

# WHAT IS PARKINSON'S DISEASE

## Progressive

A neurodegenerative disorder driven by ongoing loss of dopamine-producing neurons.

## More than movement

Causes motor symptoms (tremor, slowness, stiffness) and non-motor symptoms (sleep, mood, autonomic).

## No disease-modifying therapy

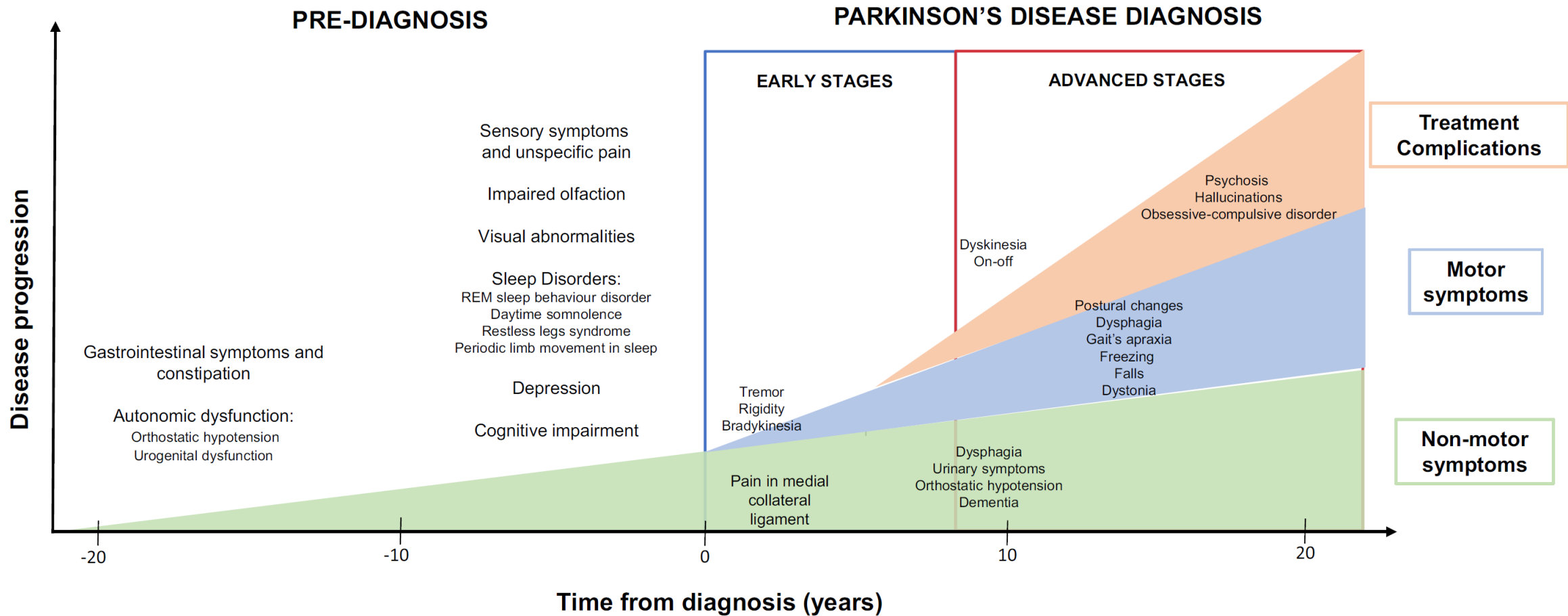
Current treatments ease symptoms; none is proven to slow the underlying disease.

## The unmet need

A therapy that addresses non-motor symptoms and slows progression – not just masks symptoms – would be first-of-its-kind.

Parkinson's disease is associated with progressive loss of dopamine neurons causing both movement and non-movement symptoms. No existing therapy slows progression.

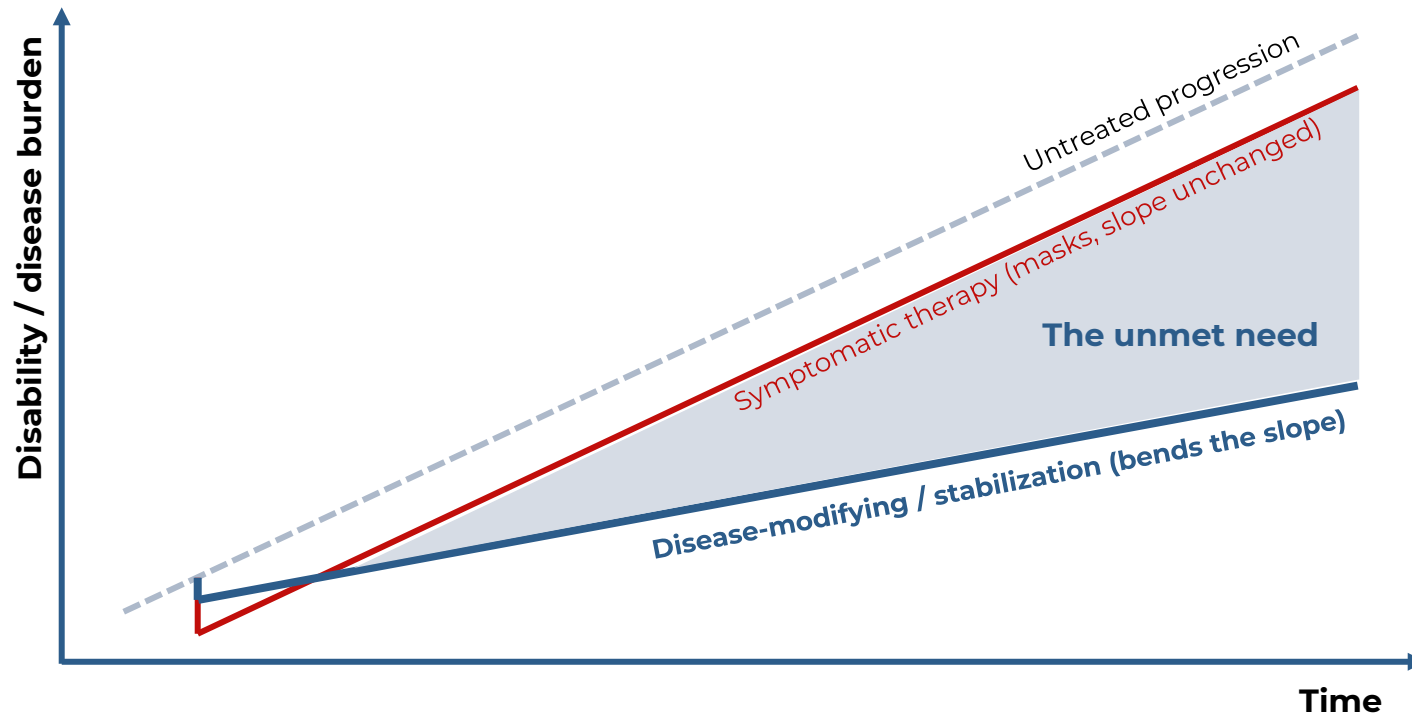
# SYMPTOMATIC PROGRESSION IN PD



# CONCEPTUAL DEPICTION OF THE UNMET NEED

Existing PD therapies relieve symptoms – not disease progression. A disease-modifier remains the unmet need

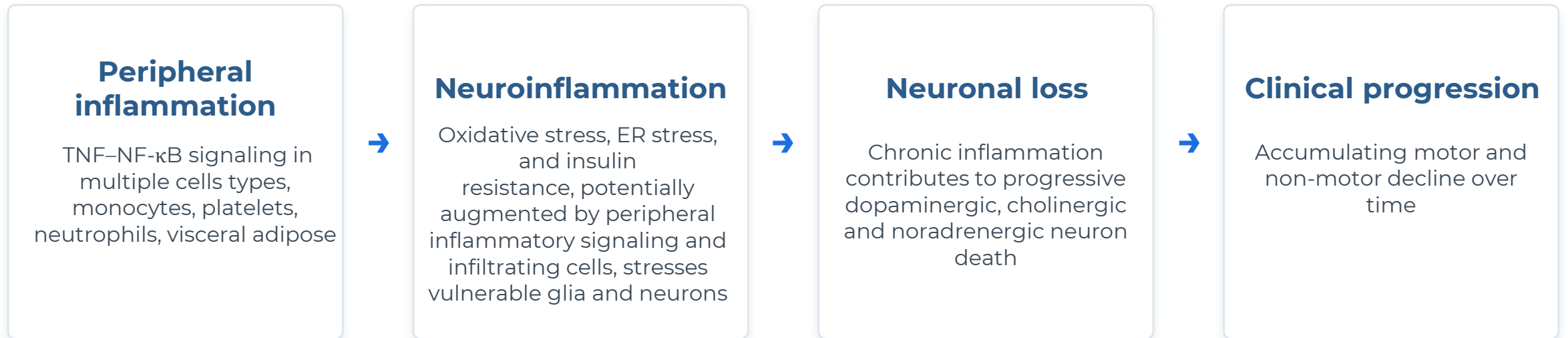
Illustration of the unmet need: Changing the slope to slow progression



- **Intercept shift** — symptomatic drugs mask while progression continues.
- **The real need** — to bend the slope
- **Benefit signal** — shows up as “less worsening”
- **Endpoint gap** — short-term motor endpoints poorly detect change

# THE NEUROINFLAMMATION HYPOTHESIS

Why an anti-inflammatory could modify Parkinson's disease.



Bezisterim is believed to dampen the TNF / NF-κB step without broad immunosuppression. Reducing inflammation could slow the disease.

# WHAT WE MEASURED IN THE SUNRISE-PD TRIAL

Four windows on the biology

1

## Inflammation

CBC-derived ratios  
(MLR, SIRI, NLR...)

*Systemic inflammatory  
tone*

2

## Proteome

380 plasma proteins  
related to inflammation  
and CNS disease

*Coordinated molecular  
shift*

3

## Nerve injury

Neurofilament light  
(NfL), Glial fibrillary  
acidic protein (GFAP),  
others

*Ongoing axonal damage*

4

## Clinical scales

MDS-UPDRS I-III,  
PDSS-2, PDQ-39

*Symptoms & function*

# HOW TO READ THE NUMBERS

Two yardsticks used throughout: effect size (how big) and the p-value (how sure)

## Cohen's d — how big is the effect?

**d ≈ 0.2**      small

**d ≈ 0.5**      medium

**d ≈ 0.8**      large

**Negative value**      Treatment improvement

*Most bezisterim effects sit in the “large” zone.*

## p-value — how sure are we?

**Smaller p = less likely to be coincidental**

p < 0.05 is the conventional bar for statistical significance

# BASELINE CHARACTERISTICS

Treatment and placebo arms appear balanced at baseline

| Characteristic                                      | Bezisterim   | Placebo      | p-value |
|-----------------------------------------------------|--------------|--------------|---------|
| n                                                   | 28           | 29           | —       |
| Age, years (mean ± SD)                              | 65.8 ± 7.8   | 66.2 ± 7.4   | 0.849   |
| Sex, Male / Female (n)                              | 20 / 8       | 16 / 13      | 0.274   |
| MDS-UPDRS Part I (mean ± SD)                        | 6.2 ± 3.9    | 5.0 ± 3.9    | 0.247   |
| MDS-UPDRS Part II (mean ± SD)                       | 6.9 ± 3.9    | 7.0 ± 4.8    | 0.903   |
| MDS-UPDRS Part III (mean ± SD)                      | 25.0 ± 7.6   | 25.5 ± 8.8   | 0.993   |
| CGI-S (mean ± SD)                                   | 2.9 ± 0.7    | 2.8 ± 0.6    | 0.716   |
| MoCA (mean ± SD)                                    | 27.8 ± 1.4   | 28.1 ± 1.8   | 0.505   |
| GDS-SF (mean ± SD)                                  | 1.9 ± 1.7    | 1.3 ± 1.0    | 0.098   |
| ESS (mean ± SD)                                     | 4.6 ± 2.2    | 5.3 ± 2.6    | 0.298   |
| Baseline Platelets, ×10 <sup>9</sup> /L (mean ± SD) | 235.1 ± 49.9 | 241.0 ± 70.7 | 0.731   |
| Baseline Platelets >230, n (%)*                     | 14 (50.0%)   | 14 (48.3%)   | 1.000   |
| Baseline Platelets ≤230, n (%)*                     | 11 (39.3%)   | 13 (44.8%)   | —       |

# WHY EPNIC-15 COMPOSITE OF CLINICAL OUTCOMES IS NEEDED

EPNIC-15 comprises 15 subdomains from MDS-UPDRS Parts I, II, and III and PDSS-2\*

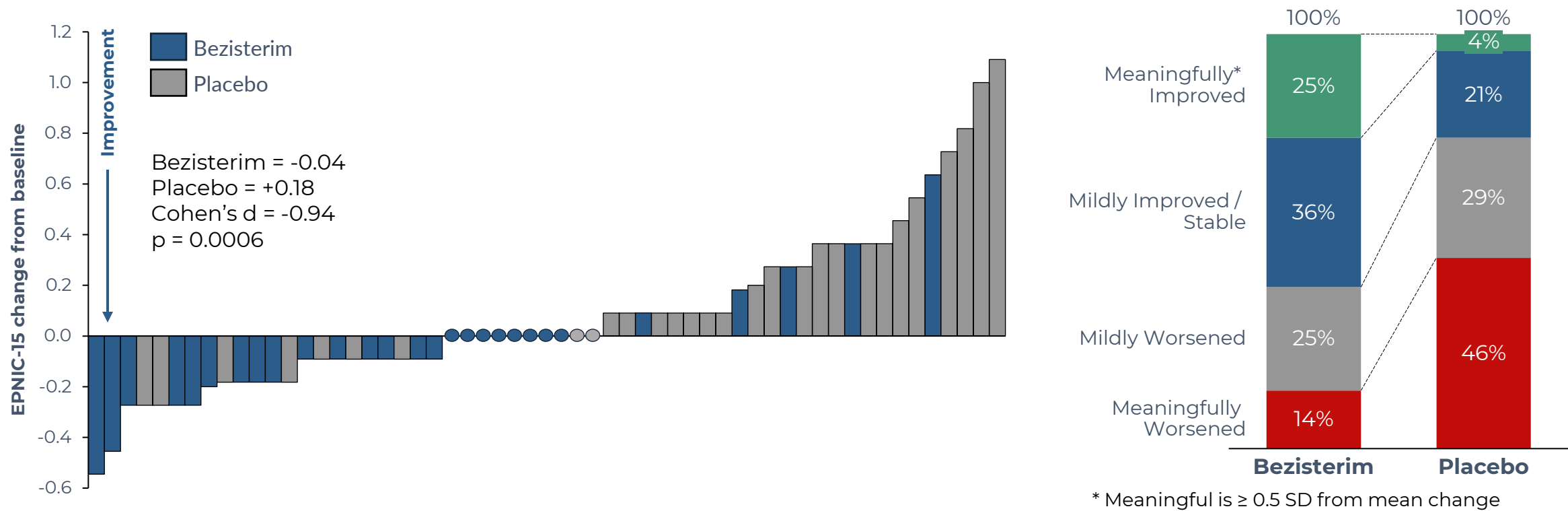
- PD therapies have historically been approved based largely on the MDS-UDPRS, which are focused primarily on motor symptoms and largely for late-stage disease
  - 6 of the last 10 approvals were based primarily on daily OFF time
  - 2 of the last 10 approvals were based on UPDRS Part III scores (motor symptoms)
- Bezisterim has shown an impact on motor symptoms. But its apparent clinical impact could be broader based on a different mechanism of action
- Therefore, using motor-focused endpoints alone may undervalue the full potential of what could be the first therapy to address both the motor and non-motor symptoms of PD
- The Early Parkinson's Neuro-Inflammatory Composite-15 (EPNIC-15) comprises 15 subdomains of UPDRS Part I-III and PDSS-2
- It was constructed based on learnings from Biohaven, Broadstreet HEOR, and Pentara's efforts to create the Parkinson's Composite Scale (PARCOMS)\*\*

| EPNIC-15 item         | Domain         |
|-----------------------|----------------|
| Sleep problems        | UPDRS Part I   |
| Daytime sleepiness    | UPDRS Part I   |
| Constipation          | UPDRS Part I   |
| Hobbies/activities    | UPDRS Part II  |
| Tremor (ADL)          | UPDRS Part II  |
| Walking & balance     | UPDRS Part II  |
| Speech                | UPDRS Part III |
| Toe tapping R         | UPDRS Part III |
| Toe tapping L         | UPDRS Part III |
| Leg agility L         | UPDRS Part III |
| Posture               | UPDRS Part III |
| Rest-tremor constancy | UPDRS Part III |
| Staying asleep        | PDSS-2         |
| Nocturia              | PDSS-2         |
| Morning tired         | PDSS-2         |

# COMPOSITE CLINICAL OUTCOMES MEASURE

Majority of treated patients improved on EPNIC-15 while placebo moved in the opposite direction

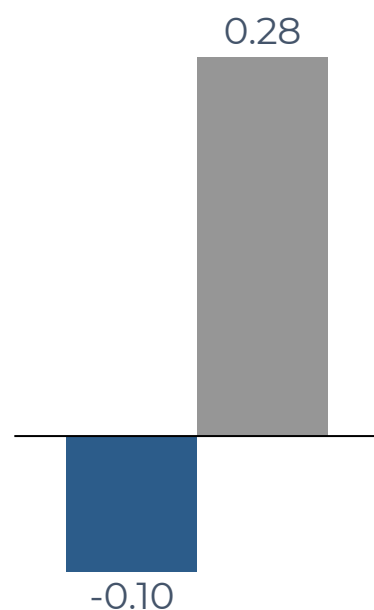
EPNIC-15 change from baseline for individual subjects



# LARGER EFFECT SIZE WITH HIGHER BASELINE INFLAMMATION

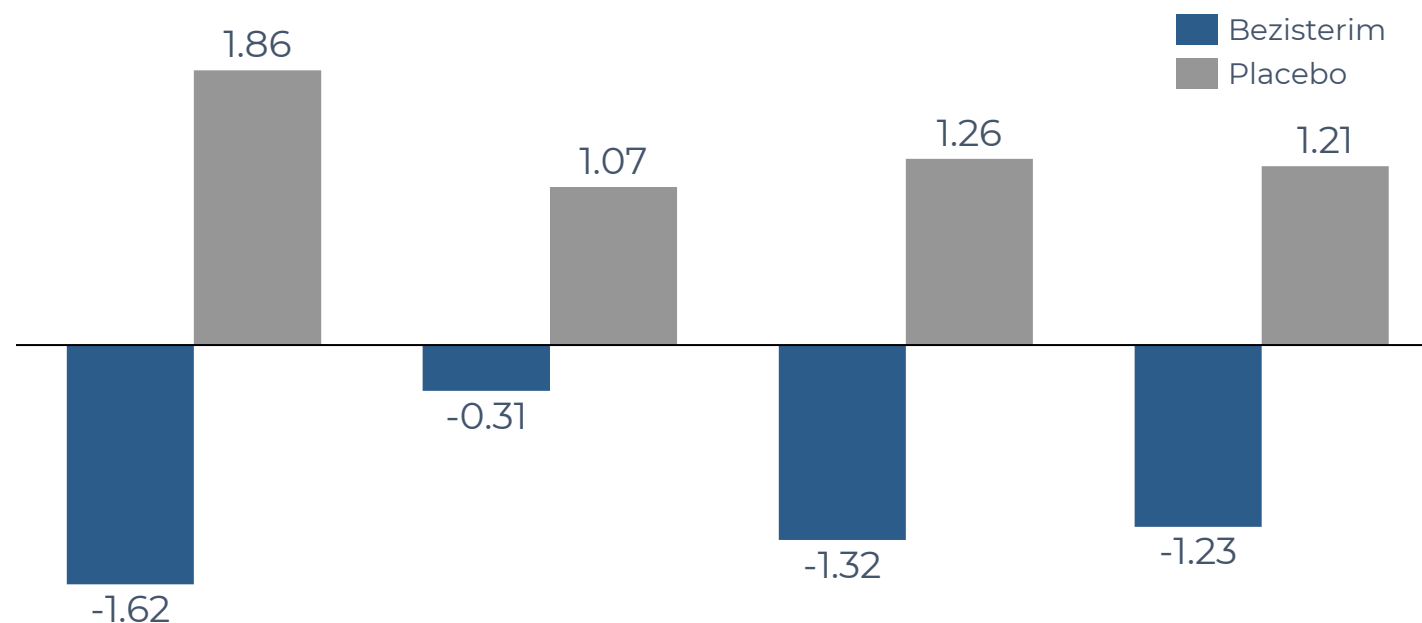
While bezisterim appeared to demonstrate an effect across the trial population, the greatest improvement was seen in those with baseline platelet counts >230

**EPNIC-15**



**EPNIC-15**  
Cohen's d = -1.61  
p=0.0002

**MDS-UPDRS Scores**



**Part I**  
Cohen's d = -1.13  
p=0.0062

**Part II**  
Cohen's d = -0.71  
p=0.0418

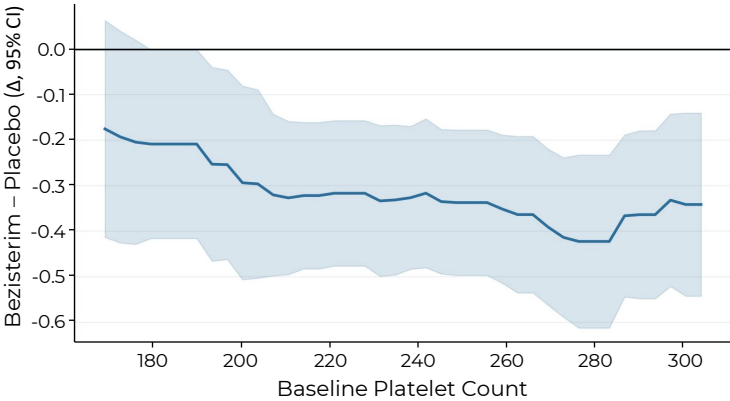
**Part III**  
Cohen's d = -0.92  
p=0.0133

**Total**  
Cohen's d = -0.85  
p=0.0198

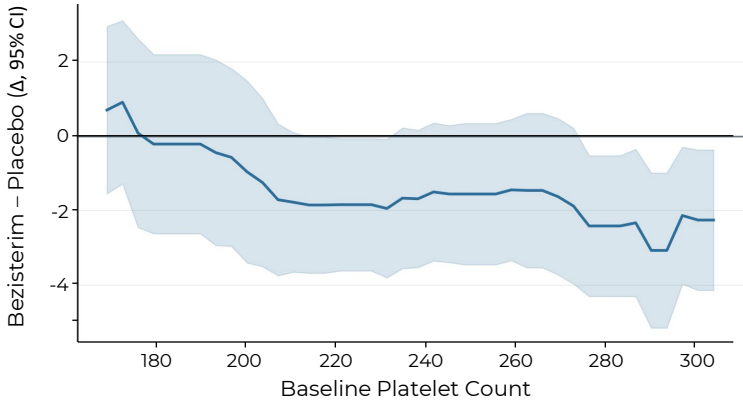
# BEZISTERIM ADVANTAGE ACROSS PLATELET RANGE

Platelet levels provide an enrichment strategy – not an exclusion of subgroups with no treatment effect

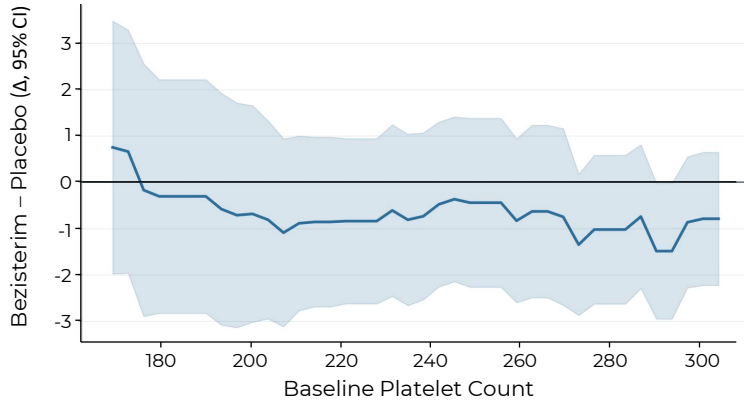
EPNIC-15



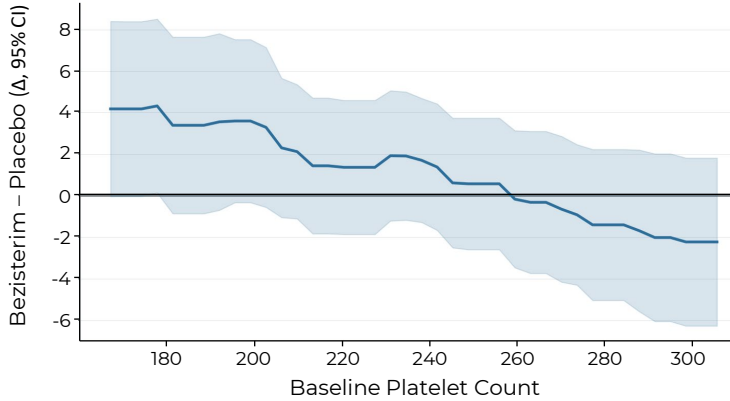
MDS-UPDRS Part I (Non-Motor)



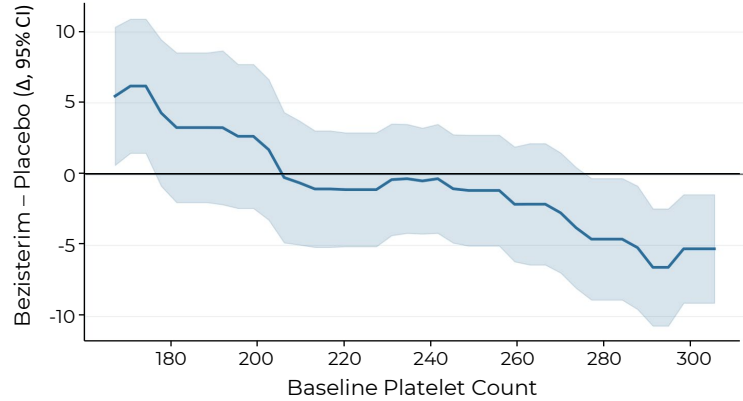
MDS-UPDRS Part II (Motor ADL)



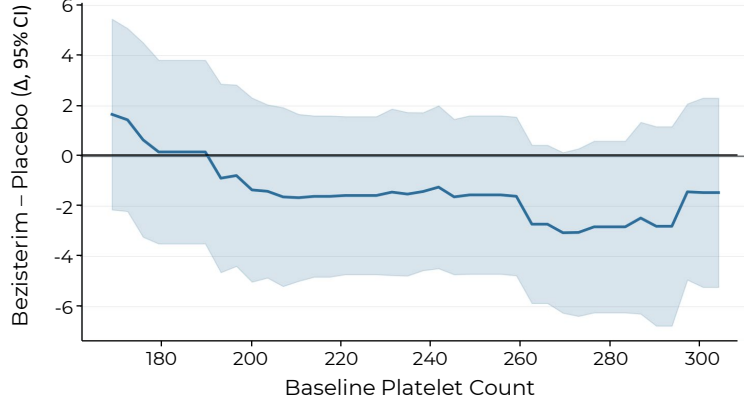
MDS-UPDRS Part III (Motor)



MDS-UPDRS Total Score



PDSS-2 (Sleep)

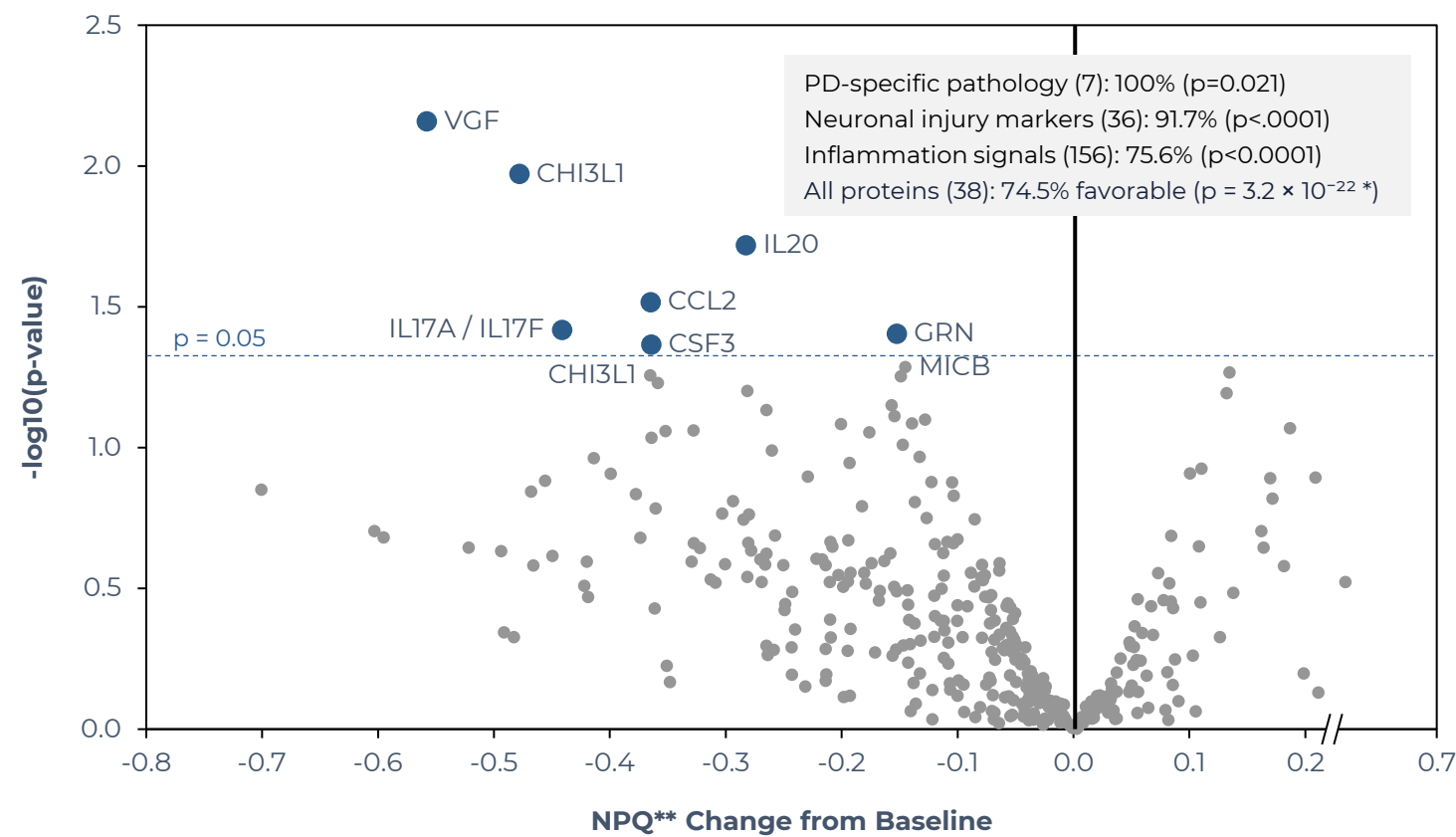


# PROTEOME-WIDE DIRECTIONALITY

283 / 380 (74.5%) of proteins moved in the direction that has been shown to slow disease progression

## Proteome volcano

Drug effect vs significance for every target



## Top biomarker effects

| Biomarker     | Panel  | $\Delta\text{NPQ}^{**}$ | Cohen's d | p      |
|---------------|--------|-------------------------|-----------|--------|
| VGF           | CNS    | -0.5580                 | -0.7730   | 0.0070 |
| CHI3L1        | CNS    | -0.4782                 | -0.7294   | 0.0107 |
| IL20          | Inflam | -0.2826                 | -0.6629   | 0.0192 |
| CCL2          | CNS    | -0.3649                 | -0.6271   | 0.0305 |
| IL17A / IL17F | Inflam | -0.4413                 | -0.5915   | 0.0382 |
| GRN           | Inflam | -0.1526                 | -0.5805   | 0.0395 |
| CSF3          | Inflam | -0.3643                 | -0.5728   | 0.0432 |
| A $\beta$ 42  | CNS    | -0.3519                 | -0.4773   | 0.0877 |
| pTau-217      | CNS    | -0.2292                 | -0.4263   | 0.1272 |

\* Binomial  $p = 3.2 \times 10^{-22}$  assumes proteins move independently. However, they are correlated. The metric provides a strong indication of the shift in the direction of a lower inflammatory burden

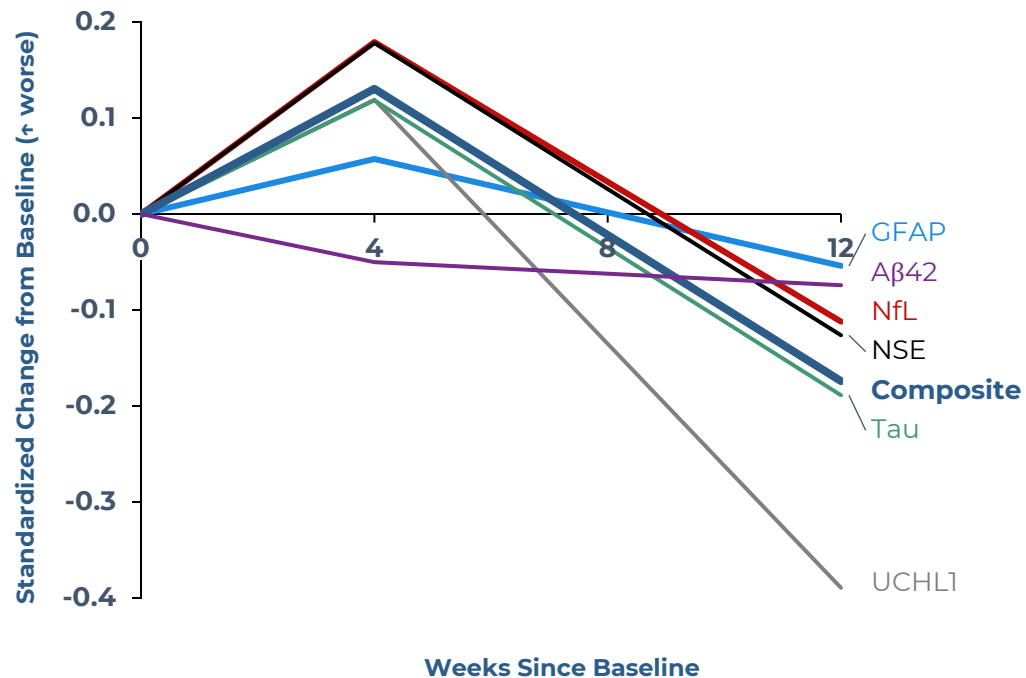
\*\* NPQ = NULISA Protein Quantification, a specialized measurement unit used in proteomics data analysis from Alamar Biosciences

VGF = VGF nerve growth factor inducible protein; CHI3L1 = chitinase-3-like protein 1 (YKL-40); IL20 = Interleukin 20; CCL2 = monocyte chemoattractant protein-1 (MCP-1); IL17A/IL17F = Interleukin 17; GRN = progranulin; CSF3 = granulocyte colony-stimulating factor (G-CSF); A $\beta$ 42 = Beta amyloid 42; pTau-217 = phospho-Tau 217

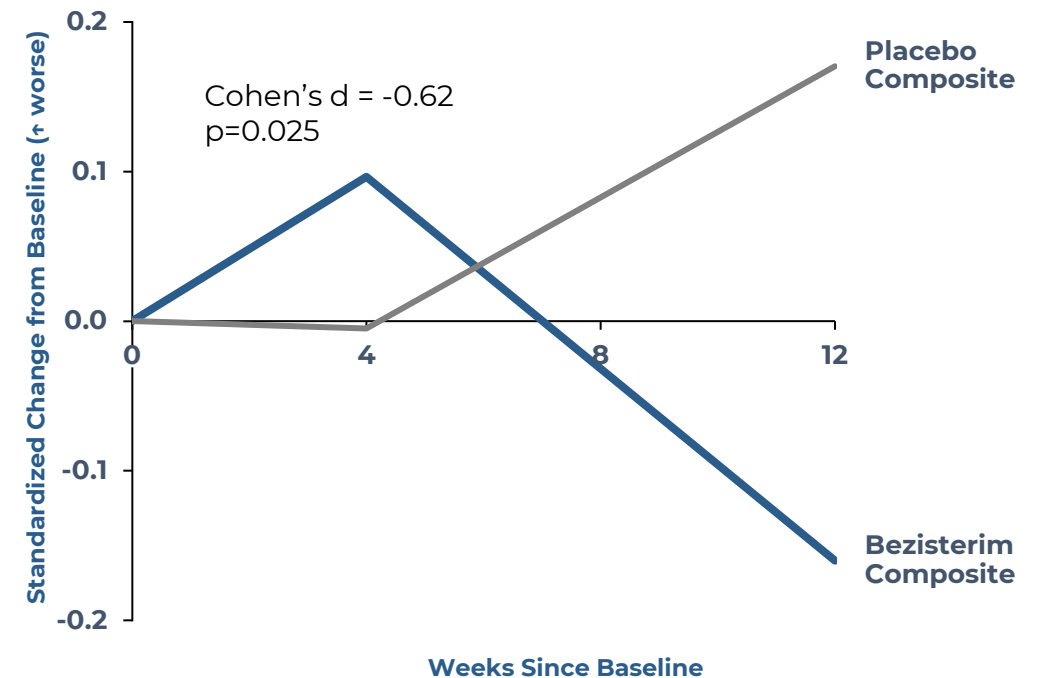
# POTENTIAL IMPACT ON NEURODEGENERATION

Neuronal injury biomarkers reduced with bezisterim

Neuronal Injury Biomarkers Change from Baseline for bezisterim treated patients



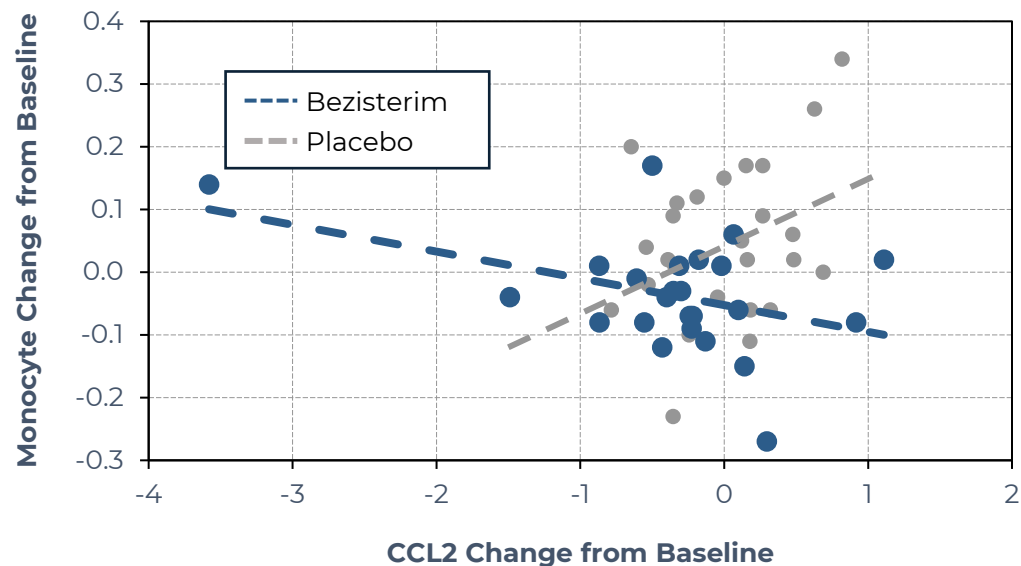
Composite Neuronal Injury Biomarkers: Bezisterim vs. Placebo



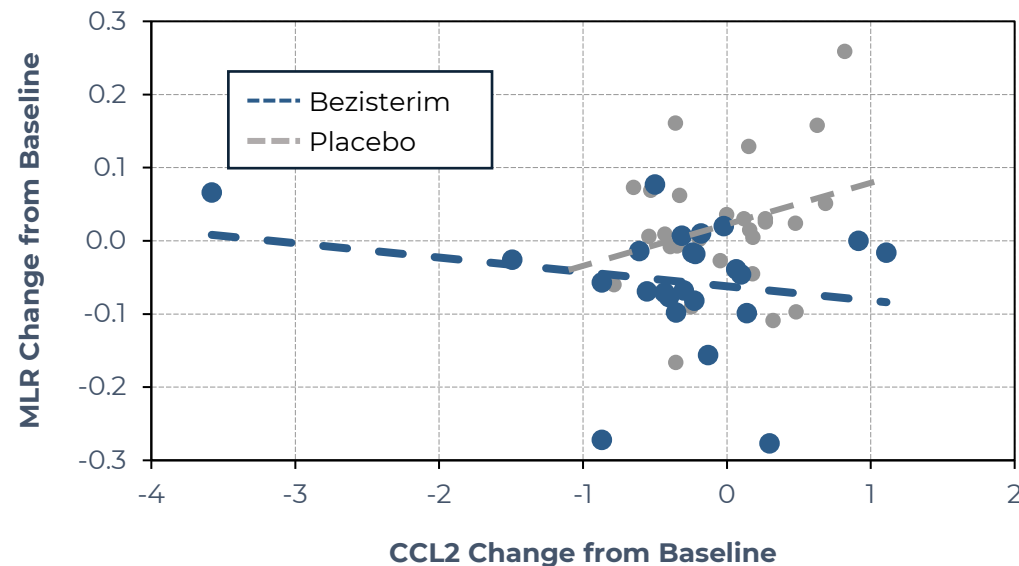
# POTENTIAL MECHANISTIC EXPLANATION

Bezisterim reduces inflammatory-driven microglia activation

Bezisterim treatment → ↓ CCL2 expression →  
↓ monocyte recruitment ...



... resulting in lower monocyte to lymphocyte  
ratio (MLR)



- MLR shows the balance between the body's general inflammation (monocytes) and immune defense (lymphocytes)
- Higher MLR is associated with chronic inflammation

# SAFETY & TOLERABILITY

Bezisterim safety and tolerability similar to placebo

|                                          | Bezisterim<br>(N=28) | Placebo<br>(N=29)    |
|------------------------------------------|----------------------|----------------------|
| <b>Subjects with any AE</b>              | <b>11 (39.3%) 19</b> | <b>15 (51.7%) 30</b> |
| Upper respiratory tract infection        | 5 (17.9%)            | 5 (17.2%)            |
| Diarrhea                                 | 1 (3.6%)             | 2 (6.9%)             |
| Headache                                 | 1 (3.6%)             | 1 (3.4%)             |
| Abdominal discomfort                     | 1 (3.6%)             | 0                    |
| Attention deficit hyperactivity disorder | 1 (3.6%)             | 0                    |
| COVID-19                                 | 1 (3.6%)             | 0                    |
| Cough                                    | 1 (3.6%)             | 0                    |
| Depression                               | 1 (3.6%)             | 0                    |
| Gout                                     | 1 (3.6%)             | 0                    |
| Musculoskeletal discomfort               | 1 (3.6%)             | 0                    |
| Uncoded                                  | 1 (3.6%)             | 0                    |
| Urinary tract infection                  | 1 (3.6%)             | 0                    |
| Vertigo                                  | 1 (3.6%)             | 0                    |
| Visual impairment                        | 1 (3.6%)             | 0                    |
| Back pain                                | 0                    | 2 (6.9%)             |
| Fall                                     | 0                    | 2 (6.9%)             |
| Abdominal pain upper                     | 0                    | 1 (3.4%)             |
| Arthralgia                               | 0                    | 1 (3.4%)             |
| Constipation                             | 0                    | 1 (3.4%)             |
| Dizziness                                | 0                    | 1 (3.4%)             |
| Gastroesophageal reflux disease          | 0                    | 1 (3.4%)             |
| Influenza                                | 0                    | 1 (3.4%)             |
| Ingrowing nail                           | 0                    | 1 (3.4%)             |
| Ligament sprain                          | 0                    | 1 (3.4%)             |
| Menopausal symptoms                      | 0                    | 1 (3.4%)             |
| Oedema peripheral                        | 0                    | 1 (3.4%)             |
| Pain in extremity                        | 0                    | 1 (3.4%)             |
| Presyncope                               | 0                    | 1 (3.4%)             |
| Sinusitis                                | 0                    | 1 (3.4%)             |
| Small airways disease                    | 0                    | 1 (3.4%)             |
| Syncope                                  | 0                    | 1 (3.4%)             |
| Viral upper respiratory tract infection  | 0                    | 1 (3.4%)             |

- **Overall AEs**

- Patients with any AEs: 57.1% bezisterim vs. 58.6% placebo
- Number of AEs: 28 bezisterim vs. 39 placebo

- **No high-severity events:** zero serious, zero severe, zero deaths.

- **Related AEs:** 1 treatment-related AE per arm; no bezisterim-related discontinuations.

# WHAT DOES IT ALL MEAN? OUR INTERPRETATION

Bezisterim has the potential to be the first of its kind treatment for Parkinson's disease

- Excellent safety profile in clinical studies to date
- Clear target engagement
  - Inflammation (CCL2, CHI3L1, MLR, SIRI, TNF $\alpha$  , others)
  - Proteomics: 380-target proteomic panels
  - Neurodegeneration markers (NfL, GFAP, others)
- Apparent clinical impact on both motor and non-motor symptoms
- Favorable biomarker signals for disease modification
- Clear rationale for Phase 3 trial design and endpoints
  - Both symptomatic relief and disease progression
  - Potential for enrichment strategy for Phase 3 trial

# LOOKING AHEAD

- Complete and report topline data for our ADDRESS-LC Long Covid trial
- Plan for End of Phase 2 meeting with the FDA
- Plan Phase 3 potentially pivotal registrational trials
- Continue analyzing the data and preparing presentations for conferences and submitting publications to scientific journals
- Reactivate potential partnering conversations in due course

# PIPELINE OVERVIEW

Multiple multi-billion-dollar US markets across the BioVie pipeline

|                                                                                         | INDICATION          | PRECLINICAL                  | PHASE 1 | PHASE 2 | PHASE 3 | Annual US Sales Potential* | Catalyst                   |
|-----------------------------------------------------------------------------------------|---------------------|------------------------------|---------|---------|---------|----------------------------|----------------------------|
| <b>Bezisterim</b><br>(formerly NE3107)<br>Oral small molecule<br>TNF $\alpha$ inhibitor | Parkinson's Disease | SUNRISE-PD · Phase 2b        |         |         |         | \$3B                       | August '26<br>Topline Data |
|                                                                                         | Long COVID          | ADDRESS-LC · \$13M DoW grant |         |         |         | \$10B                      | 3Q26<br>Topline Data       |
|                                                                                         | Alzheimer's Disease |                              |         |         |         | \$30B                      | Phase 3<br>Pending         |
| <b>BIV201</b><br>(Continuous IV<br>terlipressin)                                        | Ascites             |                              |         |         |         | \$1.6B                     | FDA Design<br>Agreed       |



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**Thank You**