# TOURMALINE

**Corporate Overview** 

**March 2025** 

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### **Our mission**

We are driven by our mission to develop transformative medicines that dramatically improve the lives of patients with life-altering immune and inflammatory diseases



### **Experienced leadership team**

### **Management Team**



Sandeep Kulkarni, MD Co-Founder and Chief Executive Officer



Ryan Robinson, CPA Chief Financial Officer



Brad Middlekauff, JD Chief Business Officer and General Counsel



Susan Dana Jones, PhD Chief Technology Officer



Kevin Johnson, PhD Chief Regulatory Officer

### **Board of Directors**

Clay Siegall, PhD Chairman

Caley Castelein, MD

**Aaron Kantoff** 

Mark McDade

Sapna Srivastava, PhD

**Parvinder Thiara** 

Sandeep Kulkarni, MD



Emil deGoma, MD Senior Vice President, Medical Research



Gerhard Hagn Senior Vice President, Head of Commercial & BD



Don Fitch Senior Vice President, Product Development



**Dora Rau** Senior Vice President, Head of Quality

## **Key highlights**



**An IL-6 renaissance is underway:** new insights emerging about a broad range of indications where IL-6 may be clinically validated



Pacibekitug (also referred to as TOUR006) has demonstrated best-in class potential: long-acting, low immunogenicity, and low-volume subcutaneous administration observed in clinical trials to date



Two paths to significant value creation: (1) cardiovascular inflammation and (2) thyroid eye disease



A late-stage clinical company: Phase 2 TRANQUILITY trial in CV and pivotal Phase 2b spiriTED trial in TED ongoing



Two potentially transformative data readouts expected in 2025: Topline data from TRANQUILITY trial expected in Q2 2025 and topline data from spiriTED trial expected in H2 2025



**Well-financed:** cash expected to fund operations into H2 2027, enabling the delivery of key anticipated milestones for both paths

# Pacibekitug: a long-acting anti-IL-6 monoclonal antibody with best-in-class potential



### Attributes observed to date

Long-acting with terminal half-life of ~7 weeks1

>90% pathway inhibition after single 10mg dose<sup>2</sup>

Fully human with ADAs in only 0.5% of patients<sup>3</sup>

**High affinity** to IL-6<sup>4</sup>

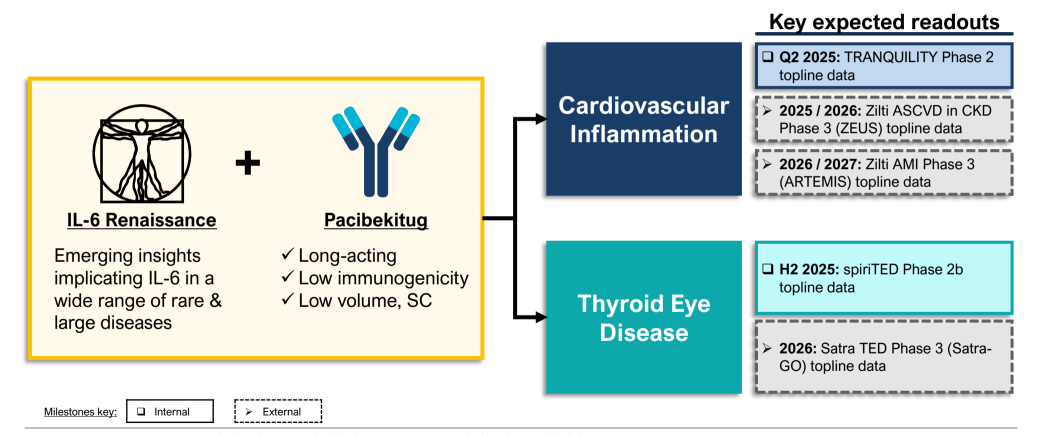
Existing data from approximately **450 study** participants<sup>1</sup>



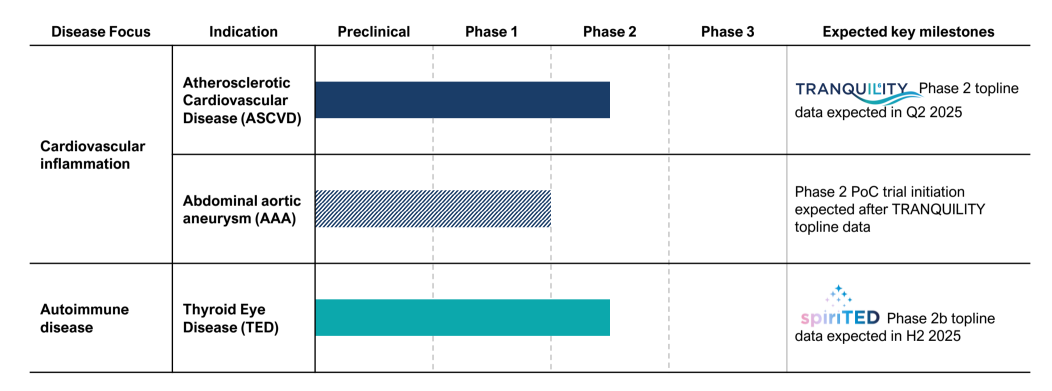
### Potential value to patients

- **Dosing quarterly**<sup>5</sup> (CV) or every 8 weeks<sup>6</sup> (TED)
- Rapid and robust impact across diseases
- Durable benefit without need to increase dose
- Volume of ≤1ml for SC injection<sup>5,6</sup>
- Generally well-tolerated safety profile observed to date

## Two paths to unlock major value creation



## Clinical development plan for pacibekitug



Note: the hatched bar represents a trial that has not yet commenced. The timing of clinical trial milestones is subject to change and additional discussion with the FDA.

## **Cardiovascular Inflammation**

### Reducing inflammation: the next frontier in CV diseases



Increasing validation for IL-6 driven inflammation as a critical and modifiable risk factor driving residual cardiovascular risk



Potential of IL-6 inhibition spans a broad range of cardiovascular indications, affecting tens of millions of patients globally



Converging lines of human evidence across multiple settings support the transformative potential of IL-6 inhibition



IL-6 inhibition is being evaluated in multiple cardiovascular outcomes trials with external readouts expected over the next 12 to 24 months



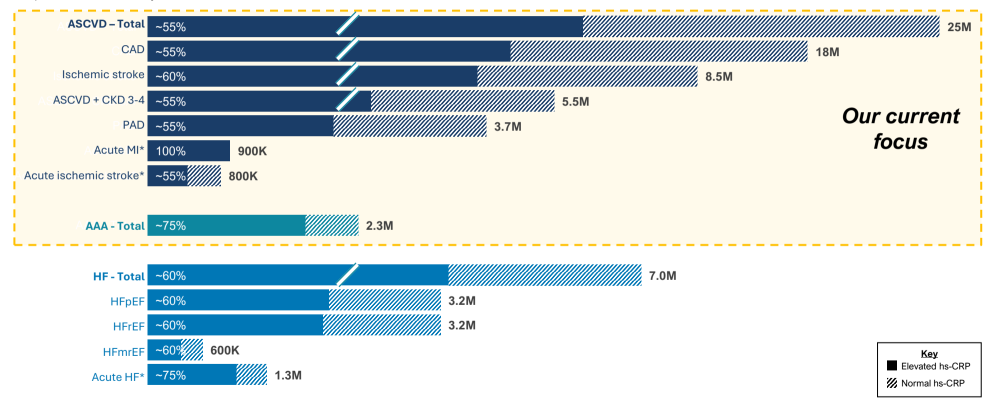
Pacibekitug's potentially best-in-class profile, including quarterly SC administration, is being evaluated in the Phase 2 TRANQUILITY trial – over-enrollment completed, topline data expected in Q2 2025

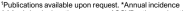
## IL-6 inhibition has the potential to address millions of patients across a wide range of inflammation mediated CV conditions

### Estimated US prevalence (2024)<sup>1</sup>

Populations are not mutually exclusive

TOURMALINE





# World-class Cardiovascular Scientific Advisory Board guiding our development strategy for pacibekitug



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Robin Choudhury, MA, DM University of Oxford



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Michael D. Shapiro, DO, MCR Wake Forest University



**Tabassome Simon, MD, PhD**Sorbonne Université
Assistance Publique-Hôpitaux de Paris



Michael Szarek, PhD University of Colorado CPC Clinical Research

## Cardiovascular inflammation largely unaddressed by existing treatments

Atherothrombotic **Pathways**  **Thrombosis** 

**Hypertension** 

**Atherogenic** lipoproteins

Diabetes, Insulin resistance. Obesity



Inflammation

**Biomarkers** 

None readily available

**Blood pressure** 

ApoB, Non-HDL-C, LDL-C, Triglycerides, Lipoprotein(a) HbA1c, Fasting glucose, Weight

C-reactive protein

**Approved Therapies** 

Aspirin P2Y12R inhibitors Factor Xa inhibitors PAR-1 antagonists

ACEI/ARB Calcium channel blockers Thiazide diuretics Renin inhibitors Beta-blockers Mineralocorticoid antagonists

Angiotensinogen inhibitors

Aldosterone synthase

inhibitors

Endothelin antagonists

Renal denervation Baroreceptor activation

Statins PCSK9 inhibitors Icosapent ethyl NPC1L1 inhibitors ACL inhibitors Bile acid sequestrants MTP inhibitors ANGPTI 3 inhibitors **Apheresis** 

**CFTP** inhibitors Lipoprotein(a) inhibitors ApoC3 inhibitors **Fibrates** CRISPR PCSK9 base editing

SGLT2 inhibitors GLP-1 agonists GIP/GLP-1 agonists

Colchicine

Therapies in Late-Stage **Development** 

Factor XI inhibitors Factor XIa inhibitors

GIP/GLP-1/glucagon agonists Amylin agonists GIP-1/amylin agonists

**IL-6** inhibitors NLRP3 inhibitors

### Increasing recognition of inflammation & IL-6 as drivers of CV risk



#### RESEARCH LETTER

Genetically Downregulated Interleukin-6 Signaling Is Associated With a Favorable Cardiometabolic Profile

A Phenome-Wide Association Study

Association of Interleukin 6 Receptor Variant With Cardiovascular Disease Effects of Interleukin 6 Receptor Blocking Therapy A Phenome-Wide Association Study

Tianot Cai, ScD, Yichi Zhang, PhD, Yuk-Lam Ho, MPH. Nicholas Link, BA, Jiehsan Sun, PhD, Jie Haang, MS, Tianrun A, Cai, MD, Scht Zhang, PhD, Zianver, MD, Yuri Ahaja, BS, Laqueiline Honerlew, MN, BSN, MPH-Jie Haang, PhD, Lianver, Costa, MPH-P terbs Cabubert, MPH-Chana Hong, PhD, David Gagnon, MD, MPH, PhD, Yan V, Sin, PhD, J. Michael Gazilian, MD, MPH-P, etek Wilson, MD, Kelly Cho, PhD, MPH, Philip Tsao, PhD, Christopher J, O'Donnell, MD, MPH-Stabriere P. Liao, MD, MPH-for the VM-Million Veteran Program

#### RESEARCH LETTER

A Missense Variant in the IL-6 Receptor and Protection From Peripheral Artery Disease

Michael G. Levin®, Derek Klarin®, Marios K. Georgakis®, Julie Lynch, Katherine P. Liao®, Benjamin F. Voight, Christopher J. O'Donnell®, Kyong-Mi Chang, Themistocles L. Assimes, ®Philip S. Tsao®, Scott M. Damrauer®, on behalf of the VA Million Veteran Program

### Interleukin-6 in Patients With Heart Failure and Preserved Ejection Fraction

Alessio Alogna, MD, PriD, <sup>a,b,c</sup> Katlyn E. Koepp, PriD, <sup>a</sup> Michael Sabbah, MD, <sup>a</sup> Jair M. Espindola Netto, PriD, <sup>d</sup> Michael D. Jensen, MD, <sup>c</sup> James L. Kirkland, MD, PriD, <sup>di</sup> Carolyn S.P. Lam, MBBS, <sup>a</sup> Masaru Obokata, MD, PriD, <sup>a</sup> Mark C. Petrie, MD, <sup>b</sup> Paul M. Ridker, MD, MPH, <sup>di</sup> Hidemi Sorimachi, MD, PriD, <sup>a</sup> Tamara Tchkonia, PriD, <sup>di</sup> Adriaan Voors, MD, PriD, <sup>b</sup> Margaret M. Redfield, MD, <sup>a</sup> Barry A. Borlaug, MD<sup>a</sup>

Research Letter

Genetically Proxied IL-6 Receptor Inhibition and Coronary Artery Disease Risk in a Japanese Population

Sizheng Steven Zhao<sup>1,\*</sup>, Dipender Gill<sup>2</sup>

<sup>1</sup> Centre for Musculoskeletal Research, Division of Musculoskeletal and Dermatological Science, School of Biological Sciences, Faculty of Biological Medicine and Health, The University of Manchester, Manchester Academic Health Science Centre, Manchester, UK

<sup>2</sup> Department of Epidemiology and Biostatistics, Imperial College London, London, UK

#### RESEARCH ARTICLE

### Circulating Interleukin-6 Levels and Incident Ischemic Stroke

A Systematic Review and Meta-analysis of Prospective Studies

Andreas Papadopoulos, MD, Konstantinos Palaiopanos, MD, Harry Björkbacka, PhD, Annette Peters, PhD, James A. de Lemos, MD, Sudha Seshadri, MD, Martin Dichgans, MD, and Marios K. Georgakis, MD, PhD Neurolesy® 2022;98:e1002-e1012. doi:10.1212/VNLI.000000000013274

Orrespondence
Dr. Georgakis
marios.georgakis@

## Quantifying inflammation using interleukin-6 for improved phenotyping and risk stratification in acute heart failure

Eleni Michou<sup>1†</sup>0, Desiree Wussler<sup>1,2†</sup>, Maria Belkin<sup>1</sup>, Cornelia Simmen<sup>1</sup>, Ivo Strebel<sup>1</sup>, Albina Nowak<sup>2,4</sup>, Nikola Kozhuharov<sup>1</sup>, Samyut Shrestha<sup>1</sup>, Pedro Lopez-Ayala<sup>1</sup>, Zaid Sabti<sup>1</sup>, Constantin Mork<sup>1</sup>, Matthias Diebold<sup>1</sup>, Tiffany Péquignot<sup>1</sup>, Katharina Rentsch<sup>5</sup>, Arnold von Eckardstein<sup>6</sup>, Danielle M. Gualandro<sup>1</sup>, Tobias Breidthardt<sup>1,2</sup>, and Christian Mueller<sup>1\*</sup>

#### ORIGINAL RESEARCH

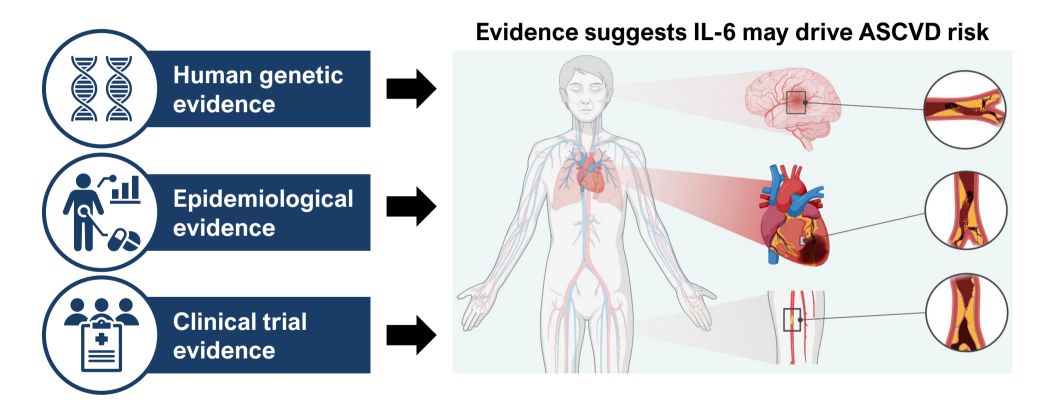
Elevated Interleukin-6 Levels Are Associated With an Increased Risk of QTc Interval Prolongation in a Large Cohort of US Veterans

Pietro Enea Lazzerini <sup>©</sup>, MD, Michael Cupelli, PhD, Alessandra Cartocci <sup>©</sup>, MSc; Iacopo Bertolazzi, MD; Vola Sakini MD, Riccardo Accioli <sup>©</sup>, MD, Falbio Sakvadori <sup>©</sup>, MD; Tormaso Marzotti, MD; Decoroso Verenega <sup>©</sup>, MD, Gabriele Cevenini <sup>©</sup>, Bioferg, Stefania Biogon, MD, Maurizo Bicchi, MD; Giovanni Donati, MD; Soiala Bernardrii <sup>©</sup>, MD, Franco Laghi-Paini <sup>©</sup>, MD, Maurizo Acampa <sup>©</sup>, MD; Per Lecooldo Capacchi <sup>©</sup>, MD, PhC Nabil E-Sherit MD. Mohamed Bouldis <sup>©</sup>, PhD

Associations of genetically predicted IL-6 signaling with cardiovascular disease risk across population subgroups

Marios K. Georgakis<sup>1,2,3</sup>\*, Rainer Malik<sup>3</sup>, Tom G. Richardson<sup>4</sup>, Joanna M. M. Howson<sup>4</sup>, Christopher D. Anderson<sup>1,2,5</sup>, Stephen Burgess<sup>6,7</sup>, G. Kees Hovingh<sup>8,0</sup>\*, Martin Dichgans<sup>3,10,11</sup> and Dipender Gill<sup>8,6,2,1,3,4</sup>\*

## Convergence of human evidence supports therapeutic potential of IL-6 inhibition for ASCVD



## Human genetic studies provide initial support for IL-6 pathway inhibition to lower ASCVD risk



Concordance between results of human genetic studies and randomized clinical trials

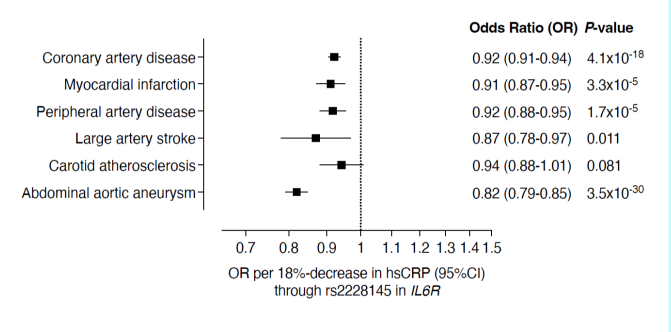
Therapeutic target	Genetic Result	RCT Result
Lowering LDL-C to lower ASCVD risk <sup>1,2</sup>	Positive	Positive
Inhibiting IL-6 to treat polymyalgia rheumatica <sup>3,4</sup>	Positive	Positive
Lowering blood pressure to lower ASCVD risk <sup>5,6</sup>	Positive	Positive
Raising HDL-C to lower ASCVD risk <sup>7,8</sup>	Negative	Negative
Inhibiting LpPLA2 to lower ASCVD risk <sup>9,10</sup>	Negative	Negative
Lowering Lp(a) to lower ASCVD risk <sup>11</sup>	Positive	Trials Ongoing
Inhibiting IL-6 to lower ASCVD risk <sup>12-16</sup>	Positive	Trials Ongoing

"Probability of success for drug mechanisms with genetic support is 2.6 times greater than those without." 17

# IL-6R gene variant mimicking low-dose IL-6 pharmacological blockade is consistently associated with a lower risk of ASCVD



### IL-6R gene variant rs2228145 associated with lower risk of ASCVD<sup>1</sup>



- rs2228145 is a common functional IL-6R gene variant (p.Asp358Ala)
- Prevalence of 30-40% (Europe)<sup>2-5</sup>
- Increased IL-6R ectodomain splicing by ADAM10/17<sup>2</sup>
- Carriers of rs2228145
   demonstrated concordant changes
   in downstream biomarkers
   compared with tocilizumab (↓CRP,
   fibrinogen;↑albumin, sIL-6R)<sup>2-5</sup>
- Modest reduction in hs-CRP ~9% per allele<sup>6</sup>

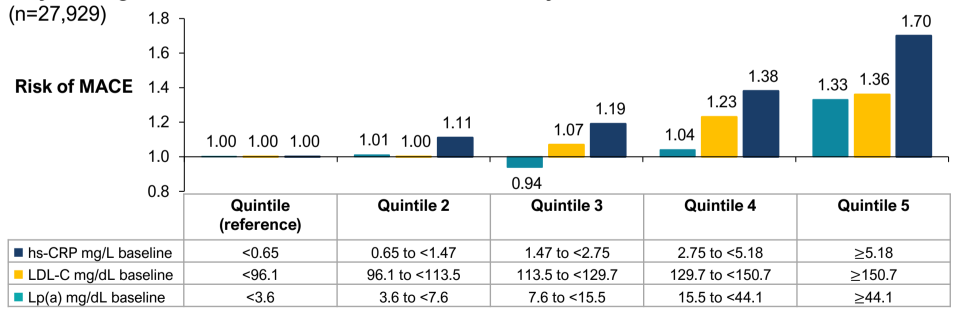


# Emerging evidence suggests that hs-CRP is more strongly associated with MACE than both LDL and Lp(a)

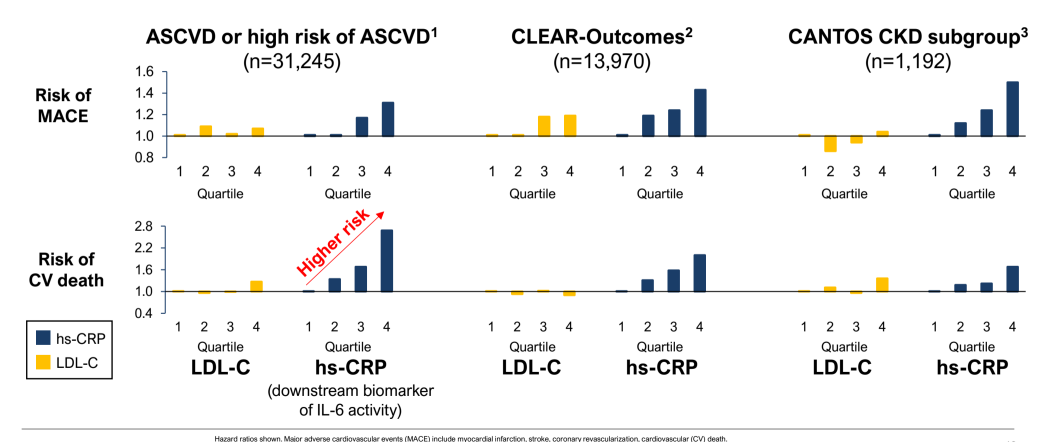


Late breaking data presented at European Society of Cardiology 2024 Congress and simultaneously published in the New England Journal of Medicine

### 30-year longitudinal data from the Women's Health Study<sup>1</sup>



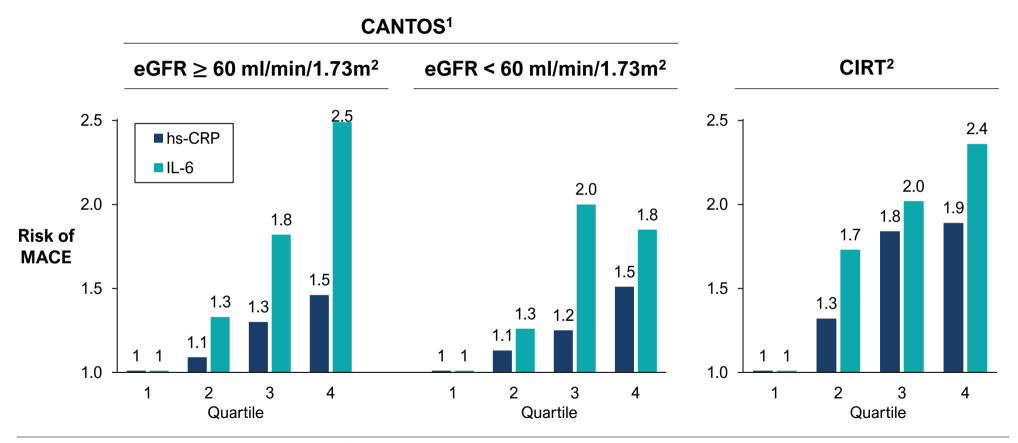
# Multiple observational studies show hs-CRP levels predict future MACE even better than cholesterol in high-risk populations





# Higher levels of IL-6, like hs-CRP, strongly and independently predicted MACE in large prospective studies

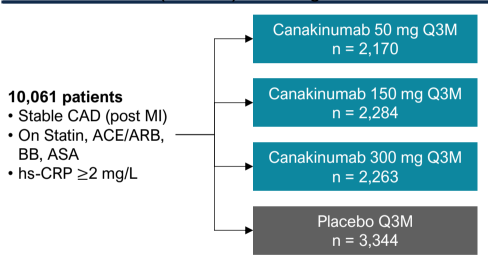




# Landmark CANTOS study validated therapeutic potential of addressing inflammation in ASCVD



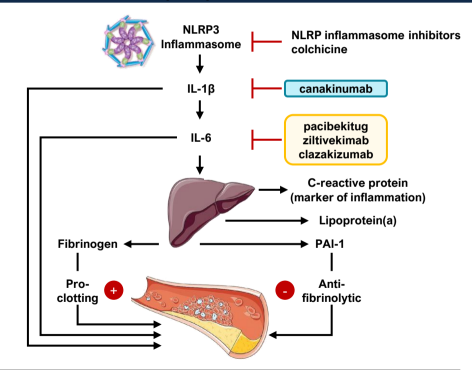
## Canakinumab Anti-inflammatory Thrombosis Outcomes Study (CANTOS) Trial Design<sup>1</sup>



### **Primary endpoint:**

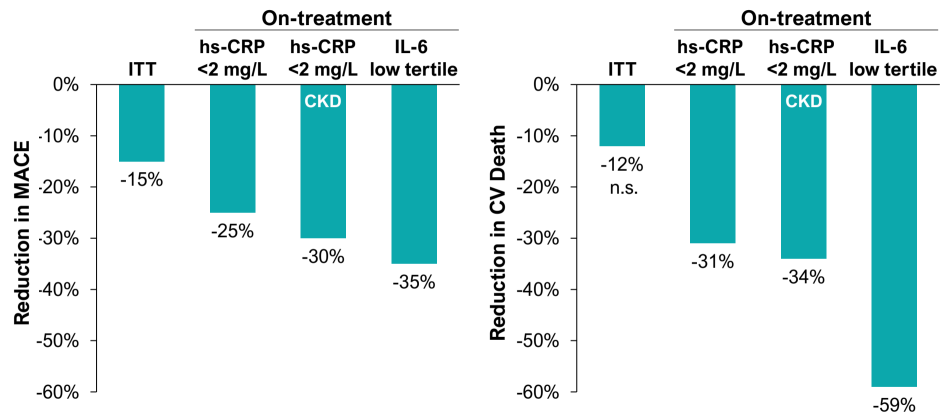
Time to the first occurrence of MACE (CV death, non-fatal MI, or non-fatal stroke)

### IL-1β is upstream of IL-62



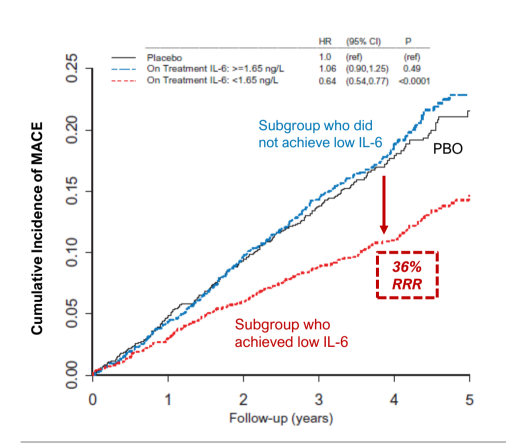
# Lessons from canakinumab (anti-IL-1β mAb): Robust IL-6 pathway inhibition associated with improved outcomes

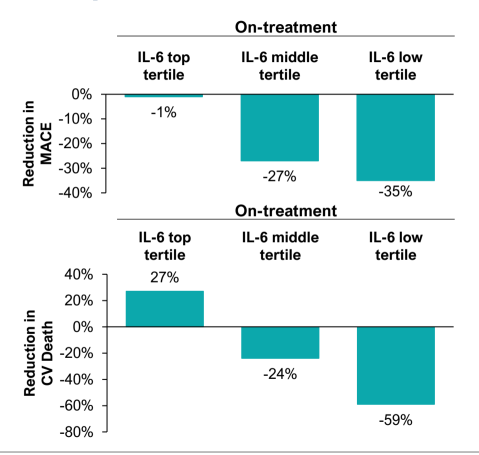




# Lessons from canakinumab (anti-IL-1β mAb): Prespecified analysis showed that reductions in IL-6 predicted CV benefit<sup>1-3</sup>

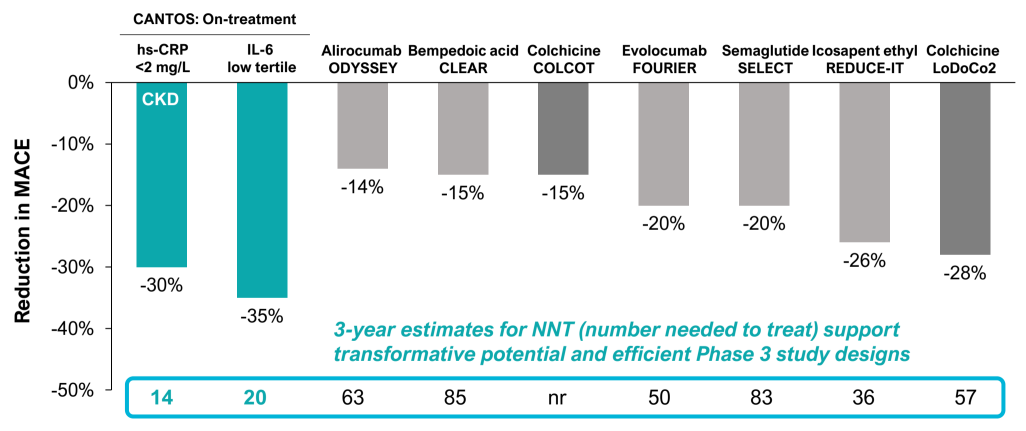






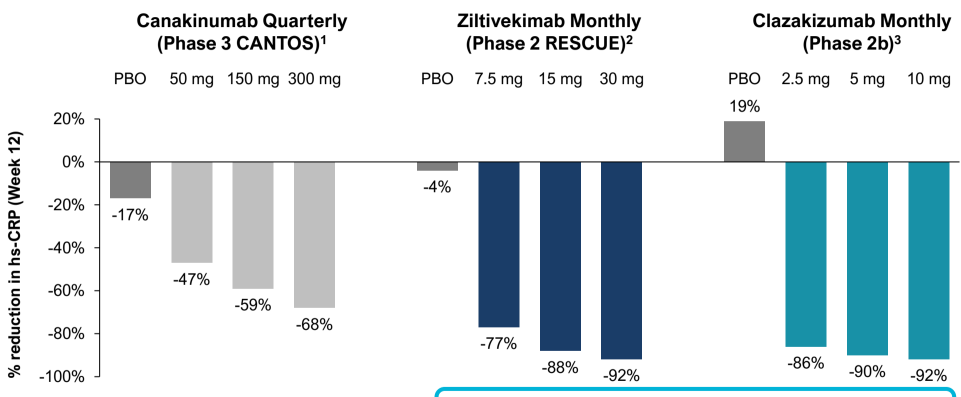
# Lessons from canakinumab (anti-IL-1β mAb): Robust inhibition of IL-6 pathway has transformative potential in ASCVD





## In independent studies, direct IL-6 inhibition lowered hs-CRP more than upstream IL-1ß blockade

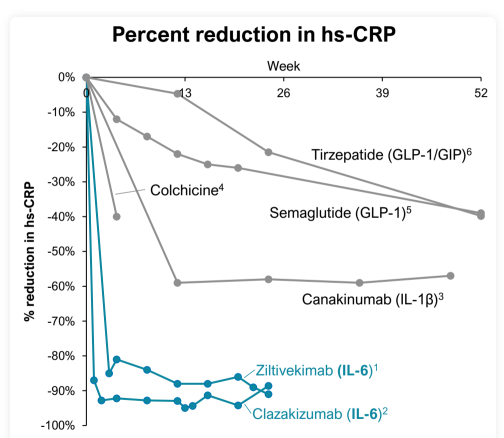


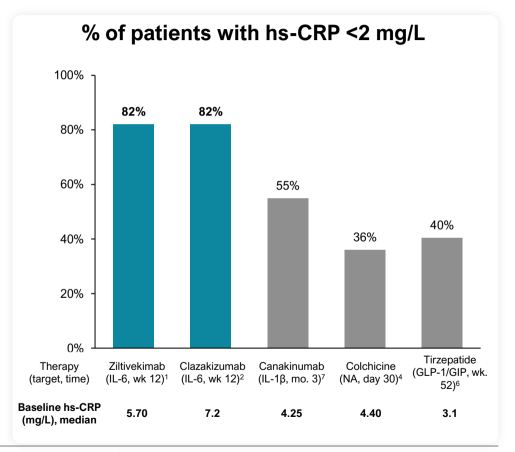


Direct IL-6 inhibition achieved ~2x placebo-adjusted reductions in hs-CRP compared to upstream IL-1β

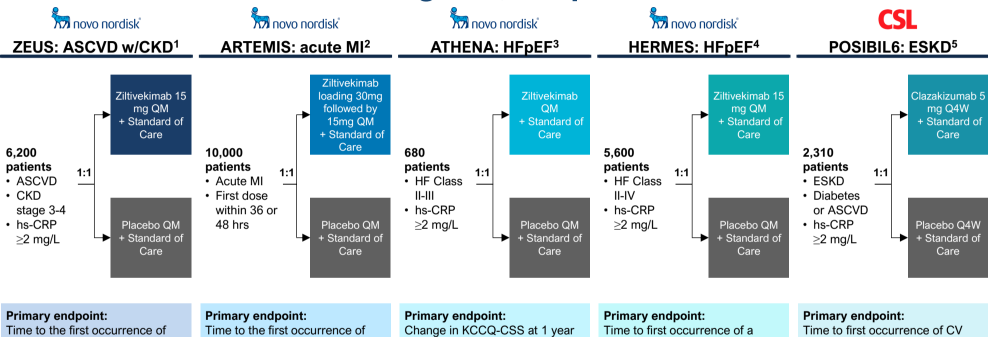
# IL-6 inhibition has produced rapid and robust hs-CRP reductions in patients with established or at high-risk of CVD







## Five Phase 3 CVOTs enrolling >24,000 patients



### Topline data readouts expected

2025 / 2026 2026 / 2027 2026 2027 2029

non-fatal stroke)

MACE (CV death, non-fatal MI, or

MACE (CV death, non-fatal MI, or

non-fatal stroke)

composite HF endpoint (CV

death, HF hospitalization, or

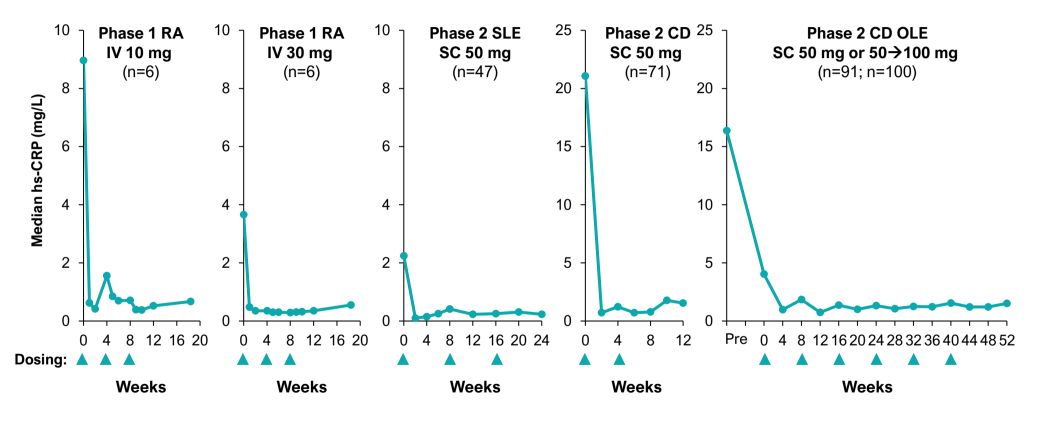
urgent HF visit)

death or MI

## Pacibekitug designed to offer best-in-class potential profile in cardiovascular diseases

	Pacibekitug	Ziltivekimab	Clazakizumab
Company	TOURMALINE	novo nordisk°	CSL
Monoclonal antibody	fully human (IgG2)	fully human (IgG1k, YTE mutation)	humanized rabbit (IgG1k)
Anti-drug antibodies <sup>1</sup>	0-1%	6-13% <sup>3,4</sup>	0-10% <sup>7-9</sup>
Route of administration <sup>2</sup>	SC 0.6 mL	SC <sup>5,6</sup> 1.0 mL	IV <sup>10</sup>
Longest dosing intervals in completed studies	Q8W (SLE, CD)	Q4W (NDD-CKD) <sup>5,6</sup>	Q4W <sup>10</sup> (HD-CKD)
Targeted dosing intervals	Quarterly	Monthly	Monthly

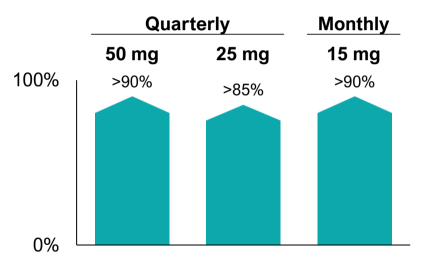
# Pacibekitug achieved robust and durable suppression of CRP in patients with high-grade inflammatory autoimmune disorders



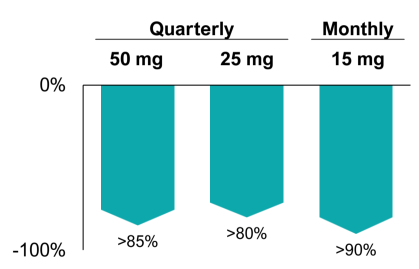
# PK/PD modeling supports potential for quarterly dosing of pacibekitug SC in ASCVD

Results of PK/PD model developed from five studies in patients with RA, SLE, CD and healthy volunteers





### Median % reduction in hs-CRP



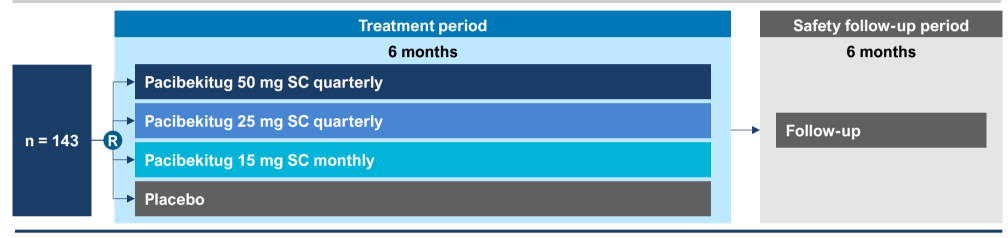
Ziltivekimab 15 mg monthly<sup>1</sup>

% achieving hs-CRP<2 mg/L: 82%

median % reduction: 88%

## TRANQUILITY Phase 2 trial supporting development in ASCVD

Double-blinded, placebo-controlled Phase 2 trial (NCT06362759) | Status: over-enrollment completed



### **Study population:**

- CKD stage 3-4 (eGFR 15-59 ml/min/1.73m<sup>2</sup>) or UPCR>200 mg/g
- hs-CRP ≥2 mg/L and <15 mg/L</li>
- Exclude patients at higher risk for safety complications (e.g., immunocompromised patients)

### Primary pharmacodynamic endpoint:

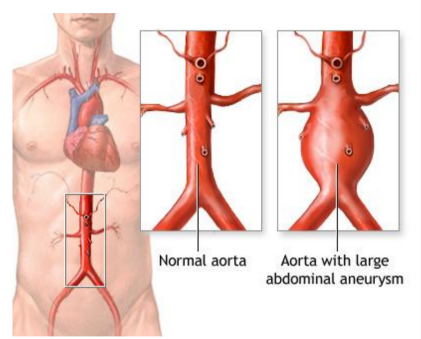
· Change from baseline in hs-CRP through Day 90

#### **Additional endpoints:**

- Percent of participants who achieve hs-CRP <2 mg/L
- Other pharmacodynamic markers, including lipoprotein (a)
- Safety and tolerability

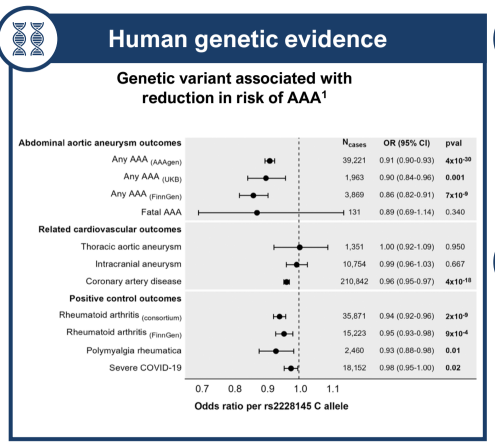
# Abdominal aortic aneurysm: a high-mortality, first-in-disease opportunity for pacibekitug

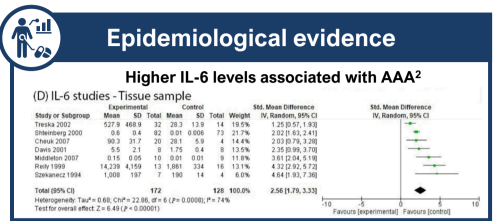
- High-risk vascular disease with significant unmet need in approximately 2M people in US<sup>1</sup>
- Strong strategic fit with ASCVD due to overlapping prescribers
- Progressive disease with increasing risk of rupture, usually a fatal event<sup>2</sup>
- In less than 5 years, majority of medium-sized AAA grow to threshold for surgical repair<sup>3,4</sup>
- Surgical repair, recommended for large AAA to prevent rupture, is **associated with complications**<sup>5-9</sup>

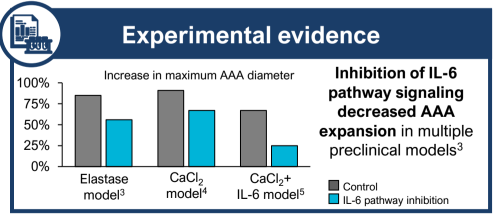


### No FDA approved treatment

## Compelling evidence supports IL-6 inhibition to slow AAA growth





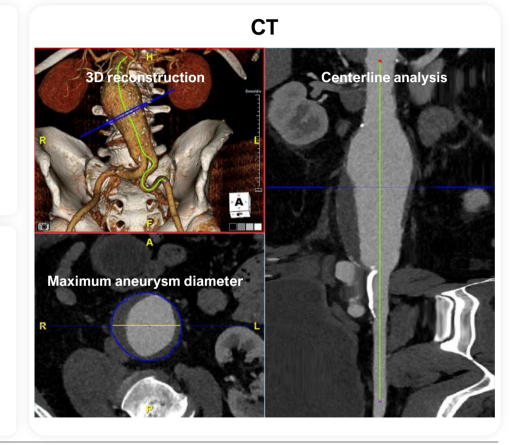


# Phase 2 PoC study expected to use imaging to evaluate the ability of pacibekitug to inhibit AAA growth

- Serial imaging is the foundation of clinical care<sup>1</sup>
- Phase 2 PoC expected to use multimodality imaging to efficiently characterize pacibekitug

### **Next steps:**

- TRANQUILITY topline data in Q2 2025
- Alignment with FDA on Phase 2 PoC design
- Details to be shared prior to study start



## **Thyroid Eye Disease**

# TED: our beachhead indication designed to validate pacibekitug's potential in autoantibody-driven diseases

- High unmet medical need with significant market opportunity
  - TED patients experience significant disease burden driven by inflammation, proptosis, double-vision, and pain
  - ~30k new patients each year in the U.S. (average age at diagnosis is ~45)<sup>1,2</sup>
  - ~80%³ of moderate-to-severe TED patients not receiving an FDA-approved treatment, which we believe may be related to significant limitations such as risk of permanent hearing impairment / loss:
    - Vast majority of US treaters report unmet need across all aspects of treatment (efficacy, safety, administration)<sup>4</sup>
- 2 Extensive third-party clinical support that IL-6 inhibition may address key unmet needs
  - 50+ publications with 400+ patients demonstrate the therapeutic potential of IL-6 pathway inhibition in TED
  - IL-6 may play a more central and upstream role in TED pathogenesis than IGF-1R or FcRn
  - Many TED treaters already routinely utilize IL-6 inhibition in their practice<sup>4</sup>
- Pacibekitug has best-in-disease potential in TED
  - Deep inhibition of IL-6 pathway observed to date offers potential for durable efficacy across many endpoints
  - Existing clinical database supports the potential for a well-tolerated profile at selected doses
  - Q8W dosing would allow for a patient-friendly, low burden treatment course

# IGF-1R class limitations present a significant opportunity for novel therapeutic approaches in TED

### TEPEZZA U.S. revenues have been stagnating since 2021...



### ...believed to be due to real-world experience

1. Safety issues: Risk of potentially permanent hearing loss<sup>2</sup>

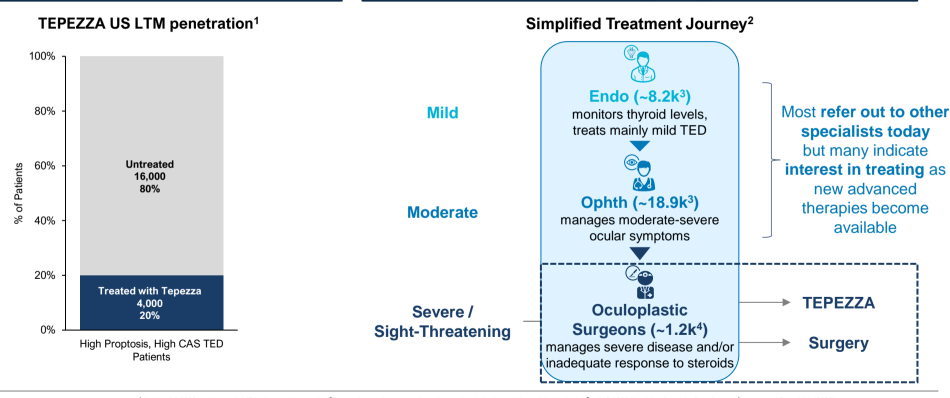
#### ------WARNINGS AND PRECAUTIONS------

- Hearing Impairment Including Hearing Loss: TEPEZZA may cause severe hearing impairment including hearing loss, which in some cases may be permanent. Assess patients' hearing before, during, and after treatment with TEPEZZA and consider the benefit-risk of treatment with patients
- Limited durability: Growing real-world effectiveness data reveals larger than anticipated number of non-responders / high relapse rate<sup>3,4</sup>
- 3. High level of inconvenience & complexity:
  - IV Q3W (n=8)<sup>2</sup> but limited access to infusion centers<sup>5</sup>
  - Numerous visits and high time commitment (HCPs and patients)<sup>5</sup>
  - Need for serial audiograms, as per label<sup>2,6</sup>
  - Burdensome reimbursement approval process<sup>7</sup>

# Despite an FDA-approved medicine, the vast majority of moderate-to-severe TED patients remain untreated

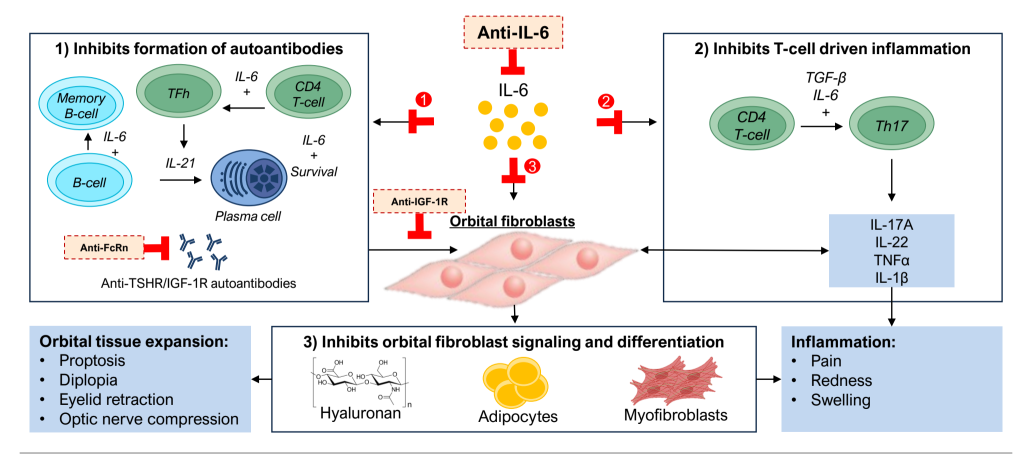
Most TED patients are not receiving TEPEZZA...

...or only get it relatively late in the treatment journey<sup>2</sup>



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## IL-6 inhibition has the potential to address a central and upstream driver of TED



# Over 50 publications support the therapeutic potential of IL-6 pathway inhibition in TED

Study	Detail	s		K	Cey Endpoin	its	Stu	dy Deta	ails		ŀ	(ey Endpoin	ts
				Proptosis	CAS	%		Í			Proptosis	CAS	%
		Study	N	response	•	autoantibody			Study	N	response	response	autoantibody
First author	Year	type	treated	rate	rate	reduction	First author	Year	type	treated	rate	rate	reduction
Pérez-Moreiras	2021	Retro	54	78	89	75	Copperman	2019	CS	2	100	C	) NR
Sánchez-Bilbao	2020	Obs	48	NR	NF	R NR	Coy	2019	CS	2	NR	50	NR NR
Atienza-Mateo	2018	Retro	29	NR	NF	R NR	Sierra Osorio	2020	CS	2	100	100	NR NR
Farde	2024	Retro	23	64	NF	75	Park	2021	CS	2	100	100	NR NR
Lee	2024	Prosp	19	11	47	56	Abeillon-du Payrat	2022	CS	2	100	50	NR NR
Pérez-Moreiras	2014	Prosp	18	72	100	76	Butnaru	2013	CR	1	NR	100	NR NR
Pérez-Moreiras	2018	RCT	15	93	60	) NS	Gómez Rodríguez	2014	CR	1	NR	100	NR NR
de la Fuente Bursón	2020	Retro	15	NR	NF	R NR	Bielefeld	2017	CR	1	CI	NF	. NR
Pereira	2023	Retro	14	NR	NF	R NR	Canas	2018	CR	1	100	NF	. NR
Habroosh	2024	Prosp	13	100	31	68	Pascual-Camps	2018	CR	1	NR	NF	. NR
Boutzios	2023	Obs	12	NR	NF	84	Garreta Fontelles	2019	CR	1	NR	NR	93
Pampín-Sánchez	2022	Retro	11	75	73	NR	Mehmet	2020	CR	1	0	NR	. NR
Moi	2022	Retro	10	CI	80	75	Kaplan	2020	CR	1	NR	C	85
Cortez	2022	Prosp	10	10	100	81	Cayon-Blanco	2020	CR	1	NR	100	NR NR
Guo	2024	Retro	10	NR	NF	R NR	Tran	2020	CS	1	NR	NF	. NR
Silkiss	2020	CS	9	CI	56	74	Ruiz	2021	CR	1	NR	NR	. NR
Smith	2021	Retro	9	78	100	54	Albrashdi	2022	CR	1	100	NF	. NR
Bielefeld	2019	Obs	8	NR	NF	R NR	Cezara	2022	CR	1	NR	C	NR NR
Ceballos-Marcias Jose	2020	CS	8	NR	75	5 41	Mohamed	2022	CS	1	0	C	) NR
Bennedjai	2020	Retro	7	NR	NF	73	Moleiro	2022	CR	1	100	NF	86
Moás	2022	Obs	7	NR	NF	92	Almazrouei	2023	CR	1	NR	NF	. NR
Toro-Tobon	2023	Retro	6	50	NF	R NR	Cuculescu	2023	CR	1	CI	C	) NR
de Pablo Gomez	2018	CS	5	NR	60	) NR	Nirmalan	2023	CS	1	NR	NR	. NR
Navarrete	2022	Retro	5	NR	NF	R NR	Pramono	2023	CR	1	NR	NF	. NR
Ribi	2017	CS	3	33	67	' NR	Rymuza	2024	CR	1	100	C	8
Maldiney	2020	CS	3	67	NF	NR NR							
Stevens	2022	Retro	3	100	67	' NR	1	Weigl	hted Mea	ın	68%	72%	72%
Russell	2017	CS	2	NR	C								
Sy	2017	CS	2	CI	50		Smith 201	٠.		,	71%	69%	N/A
•							Douglas 202	20 (tepr	o Phase	3)	83%	59%	N/A

We believe many of these reports may underestimate the true efficacy of IL-6 inhibition

- 400+ mostly steroidrefractory patients
- Late IL-6 inhibition (>9 months post symptom onset) when disease may have exited active phase
- Exposure to IL-6 inhibition may have been suboptimal (<6 months)</li>
- research with over 100 TED treaters suggests many HCPs already routinely utilize IL-6 inhibition in their practice

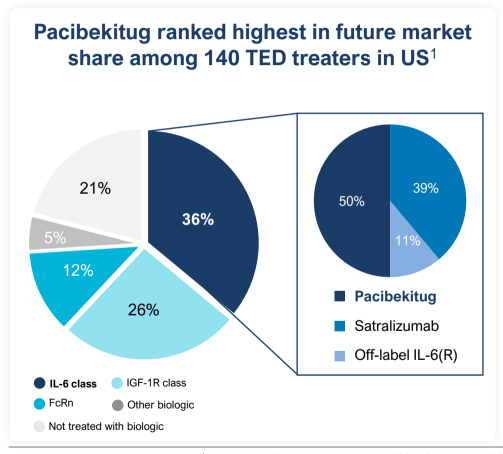


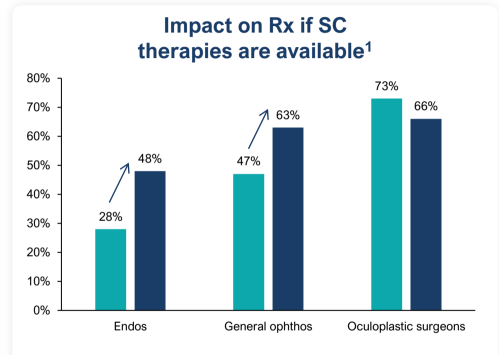
## Pacibekitug's target product profile is expected to be well-differentiated in TED...

#### **Target product profile in TED\*** Targeted points of differentiation Study population Moderate-to-severe active TED patients MOA IL-6 inhibition **Targeting inflammation** which is at core of disease Primary Proptosis endpoint **Holistic impact** on many QoL-impacting symptoms Diplopia, clinical activity score (CAS), Secondary Efficacy inflammation, and lid retraction endpoints Lower rate of relapse and retreatment Emphasis on response durability Additional Rapid time to response measures Lower rate of surgical intervention No anticipated risk of permanent hearing Warnings & **Well-tolerated** without the risk of hearing loss loss or warnings beyond typical IL-6 safety precautions considerations **Every 8-week, low volume subcutaneous** Dosina & injection through pre-filled syringe Least frequent and most patient-friendly SC dosing administration Finite dosing



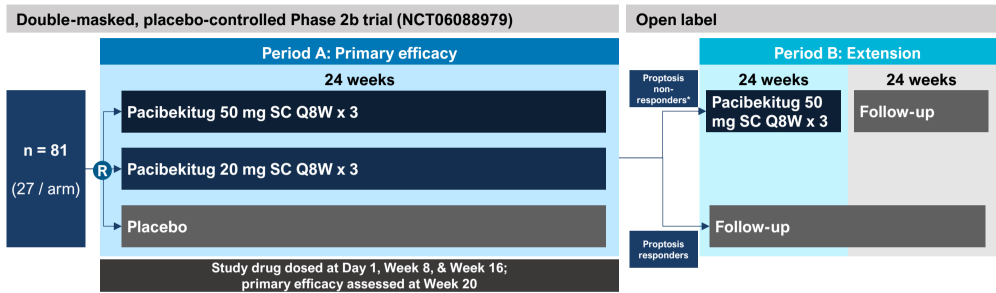
# ...resulting in leading market share, capitalizing on increasing Rx from endocrinologists and ophthalmologists





- I treat and manage moderate to severe active TED patients rather than referring out to another physician today
- As additional treatments become available for TED, including SC therapies, I will treat and manage moderate to severe active TED patients rather than referring out to another physician

# spiriTED pivotal trial in first-line TED



### **Study population:**

- Moderate-to-severe TED, with proptosis ≥ 3mm above normal (based on race and gender)
- Active phase, with symptom onset ≤ 15 months, CAS ≥ 4 and positive TSI
- First-line setting, with cap on prior corticosteroid use (< 1g methylprednisolone or equivalent)

### **Primary efficacy endpoint:**

• Proptosis response rate at week 20

### **Additional endpoints:**

- CAS
- Diplopia
- QoL, safety, PK/PD/ADA

## **Key upcoming milestones**

Disease focus	Indication	Milestone	Expected timing	
Cardiovascular	ASCVD	TRANQUILITY Phase 2 topline data	Q2 2025	
inflammation AAA	Phase 2 trial start	Post-TRANQUILITY topline data		
Autoimmune disease	TED	spiriTED Phase 2b topline data	H2 2025	

# TOURMALINE