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## Population Pharmacokinetics (PK) and Pharmacodynamics (PD) of Pegozafermin, a Novel **GlycoPEGylated FGF21**, in a Phase 2 Study in Severe Hypertriglyceridemia

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#### PURPOSE

- Pegozafermin (PGZ) is a specifically engineered, long-acting glycoPEGylated analog of fibroblast growth factor 21 (FGF21) under development for the treatment of metabolic dysfunction-associated steatohepatitis (MASH) and severe hypertriglyceridemia (SHTG).
- In a randomized Phase 2 trial (ENTRIGUE) in SHTG (triglycerides [TG]  $\geq$  500 mg/dL), PGZ-treated patients showed a significant reduction in median TG levels versus placebo (57.3% for pooled PGZ versus 11.9% for placebo, with placebo-corrected difference of -43.7%, 95% confidence interval (CI): -57.1%, -30.3%; P < 0.001).<sup>1</sup> Pharmacokinetics (PK) and pharmacodynamics (PD) data from this study were analyzed using a population modeling approach to support dose selection for Phase 3 studies. Weekly doses of 20 mg and 30 mg are currently being investigated in the Phase 3 (ENTRUST) SHTG Study (NCT05852431).

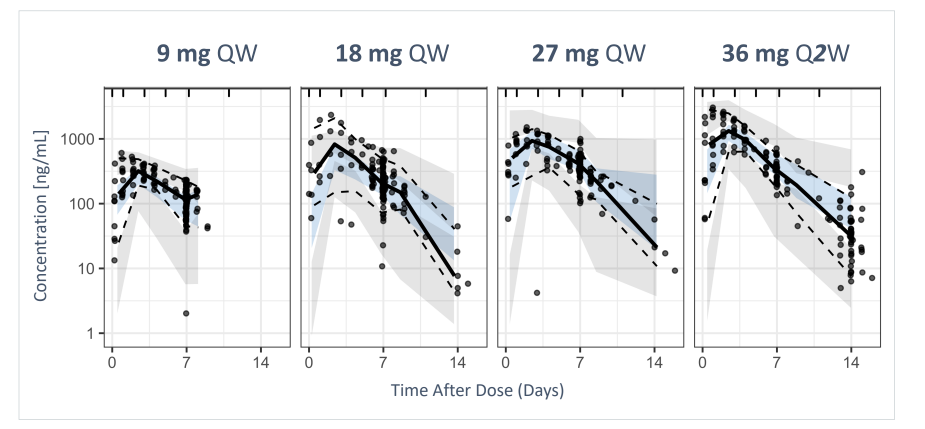
#### **METHODS**

- Subjects in this analysis included PGZ treatment groups at 9 mg (n=12), 18 mg (n=21), 27 mg (n=18) administered once-weekly (QW) and at 36 mg (n=16) administered once-every-two-weeks (Q2W).
- PK and PD (TG) data were characterized based on a population nonlinear mixed-effects modelling approach using NONMEM 7.4.2. A prior PGZ PK model was used as template and the model was then adjusted with new data.<sup>2</sup>
- POP PK/PD modeling included patients with dosing history/PK data + TG data (n= 67 IDs; 736 DVs). PK profile was first reconstructed for each patient using estimated PK individual parameters. A PKPD model for TG over time as a function of individual predicted plasma concentration (*Cp*) was then built.
- Simulation-based diagnostics, such as visual predictive checks (VPC), were used for model evaluation. Tested covariates included sex, age, weight, ethnicity, race and immunogenicity for PK and additional parameters (ALT, AST, HbA1c) added for PD. The effects of covariates were assessed using a method involving stepwise testing of linear and nonlinear relationships in a forward selection (P < 0.01) followed by a backward elimination (*P* < 0.05) procedure.

DOSE F (fixed to 1

Paramet k<sub>a</sub> (h⁻¹) MTT [1/k V (L) **CL (L/h) CL-WT** Residual Additive Proporti

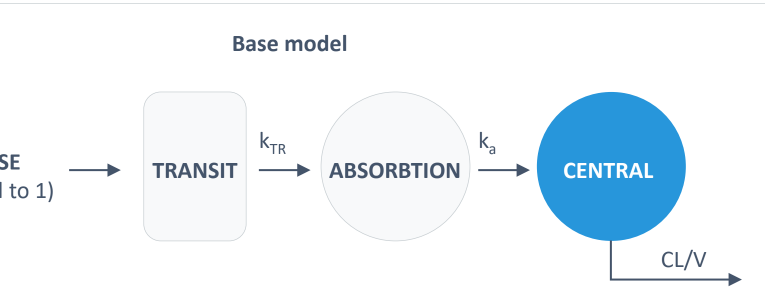
## **Pop PK Profiles**



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#### **PK POP MODEL**

#### **Pop PK Model Structure**



#### **Pop PK Model Parameters**

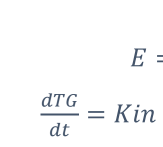
ter (units)	Estimate (RSE%)	BSV (RSE%)
	0.0381 (16)	87% (19)
<sup>[</sup> k <sub>TR</sub> ] (h)	5.68 (57)	169% (42)
	1 (fixed)	-
	18.5 (23)	29% (25)
	0.24 (11)	25% (16)
Г	0.0062 (37)	-
l variability		
e error (ng/mL)	21.3 (40)	-
ional error (CV%)	0.294 (9)	-

Upper and lower gray areas represent 2.5th and 97.5th prediction interval with uncertainty, respectively. Symbols are observed data. Dashed lines are 2.5th and 97.5th observed percentiles of the data. Blue area represents predicted median with uncertaint

#### **Pop PK Summary Results**

- **NONMEM 7.4.2.**
- 1 compartment PK model with one additional transit absorption compartment adequately describes the data. The model was adjusted for flip-flop kinetics.
- IIV was estimated on ka, CL, V and MTT parameter.
- Covariate identification was done using the stepwise covariate modelling (SCM) through the PsN software. Significant covariate included in the final model was WT on CL.
- 6% change in CL for each 10 kg change in body weight

## Longitudinal TG Model



#### **Longitudinal TG Model Parameters**

Parameters	Estimates	IIV %
Baseline	727	42%
Kout (h <sup>-1</sup> )	0.004	-
Emax (%)	7.4	86%
EC50 (ng/mL)	1180	226%
w (%)	35%	-
Kin=Base*Kout w: Proportional residual error		

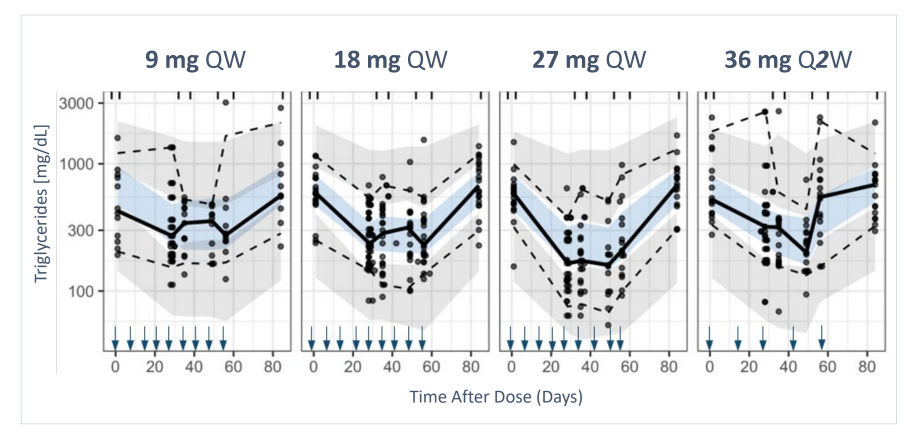
POP PD

• Pharmacokinetic data were characterized based on a population nonlinear mixed-effects modelling approach using the software

#### POP PD

### $Emax \times Cp(t)$ Cp(t) + EC50 $\frac{dTG}{dt} = Kin - Kout \times (1 + E) \times TG$

#### **Pop TG Profiles**



Jpper and lower gray areas represent 2.5th and 97.5th prediction interval with uncertainty, respectively. Symbols are observed data. Dashed lines are 2.5th and 97.5th observed percentiles of the data. Blue area represents predicted median with uncertainty Pre-dose samples for the first dose were taken on Day 0. Arrows indicate dosing day

#### **Pop PD Summary**

- An indirect response model was used to describe the longitudinal effect on TG. No significant covariates were identified.
- Pop PK/PD modeling data showed that 27 mg QW dose was the most efficacious as demonstrated by more pronounced reductions in TG as compared to the other doses.

#### CONCLUSIONS

- Population PK/PD modeling provides rationale to support dose selection for the recently initiated Phase 3 ENTRUST study in SHTG. PGZ 30 mg QW was selected to maximize efficacy and 20 mg QW was selected not only for its efficacious effect, but also to allow the sponsor to further explore the efficacy and tolerability profiles between two doses.
- Among covariates examined, only body weight can affect PGZ pharmacokinetics.

#### REFERENCES

<sup>1</sup>*Nat Med.* 2023; 29(7): 1782–1792. <sup>2</sup>Clin Pharmacol Ther 2023 Dec;114(6):1323-1331.



