

### DEVELOPING RNA-EDITING MEDICINES

for patients in need

Nasdaq: PRQR Date: January 2024

### **Forward-looking statements**

This presentation contains forward-looking statements. All statements other than statements of historical fact are forward-looking statements, which are often indicated by terms such as "anticipate," "believe," "could," "estimate," "expect," "goal," "intend," "look forward to", "may," "plan," "potential," "predict," "project," "should," "will," "would" and similar expressions. Such forward-looking statements include, but are not limited to, statements regarding our strategy and future operations, statements regarding the potential of and our plans with respect to our technologies and platforms (including Axiomer<sup>™</sup>), our preclinical model data, our pipeline targets, our other programs and business operations, our current and planned partnerships and collaborators and the intended benefits thereof, including the collaboration with Lilly and the intended benefits thereof, including the upfront payment, equity investment, and milestone and royalty payments from commercial product sales, if any, from the products covered by the collaboration, as well as the potential of our technologies and product candidates; our updated strategic plans and the intended benefits thereof, our plans to seek strategic partnerships for our ophthalmology assets, and our financial position and cash runway. Forward-looking statements are based on management's beliefs and assumptions and on information available to management only as of the date of this presentation. Our actual results could differ materially from those anticipated in these

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### **ProQR** Therapeutics

Short overview



#### Focus on Axiomer<sup>™</sup>

Exclusively focused on the development of proprietary Axiomer<sup>™</sup> RNA editing platform across multiple therapeutic areas; initial focus on liver and CNS diseases



#### Novel mechanism of action, leading patent estate

Axiomer<sup>™</sup> was discovered in ProQR labs in 2014 and uses well-proven modality of oligonucleotides to recruit a novel mechanism of action



#### Validated across multiple genes

Preclinical data demonstrate Axiomer<sup>™</sup> is broadly validated across multiple genes



#### **ADAR**

Axiomer<sup>™</sup> is ADAR-mediated RNA editing, recruiting endogenous adenosine deaminase acting on RNA (ADAR)



#### Two pillars underlie strategy

ProQR developing wholly owned pipeline with initial targets in liver-originated diseases

Selectively enter into partnerships: initial partnership with Lilly in September 2021, expansion announced December 2022





#### Cash-runway into mid-2026

Cash position of €120.6 M as of end of Q3 2023 provides runway to mid 2026, beyond multiple clinical data readouts

### **2023 accomplishments**



#### **Science and platform**

Axiomer<sup>™</sup> activity demonstrated across multiple preclinical in vitro, organoid, and in vivo models in liver and CNS - robust editing observed in mice and NHP



#### **Pipeline**

Announced initial Axiomer RNA editing pipeline programs targeting liver-originated diseases

- AX-0810 for cholestatic diseases targeting NTCP ٠
- AX-1412 for cardiovascular disease targeting • B4GALT1
- AX-2402 for Rett Syndrome a rare • neurodevelopment disorder - partnership with Rett Syndrome Research Trust focused on utilizing Axiomer to develop EONs for Rett syndrome (announced January 2024)



#### **Partnership**

A key component of ProQR's strategy

- Continued execution with Lilly partnership
- Divested sepofarsen and ultevursen to Théa who as Sepulbio will continue development of these programs for patients



IP

- Further strengthened leading global IP estate for ADARmediated RNA editing
- Multiple successful defenses against oppositions, including Europe and Japan
- Granted new patent by USPTO further underlining that the broad concept of applying endogenous ADAR by administering antisense oligonucleotides for RNA editing is proprietary to ProQR



#### Strong balance sheet

Cash position of €120.6 M as of end of Q3 2023 provides runway to mid 2026

### 2024 outlook

#### Building momentum toward development



#### **Science and platform**



- Building on robust body of preclinical data, presentation in Q1 of Axiomer data for liver NHP models
- Several additional platform data updates throughout 2024 •
- Potential for first Trident preclinical data in late 2024 •
- Continuing platform optimization and target • identification



#### **Pipeline**

- In vitro and in vivo data for AX-0810 for Cholestatic diseases targeting NTCP in mid 2024
- In vitro and in vivo data for AX-1412 for Cardiovascular diseases targeting B4GALT1 in H2 2024
- Translational data updates ahead of entry into the clinic •
- Entry into clinical trials for AX-0810 and AX-1412 on track • for late 2024/early 2025
- 2024 potential additional new pipeline target • announcement



#### **Partnership**

- Continued execution on Lilly partnership
  - Potential milestone income from existing partnership
  - potential option exercise for expansion of deal to 15 targets, with \$50 M opt-in payment to ProQR
  - potential data updates from partnership
- Potential new partnership announcements



Continued expansion of leading IP portfolio supporting that applying endogenous ADAR by administering antisense oligonucleotides for RNA editing is proprietary to ProOR

### What is ADAR editing?

**ADAR** (Adenosine Deaminase Acting on RNA)



Enzyme that performs specific form of natural RNA editing, called **A-to-I editing.** During A-to-I editing an **A nucleotide (adenosine)** is changed into an **I nucleotide (inosine)** 

#### Natural ADAR editing (A-to-I)



# Axiomer™ EONs unlock cellular machinery potential to treat diseases

By attracting ADARs and allowing highly specific editing



### **Driving the development of optimized EONs** for therapeutic use



#### **ADAR-binding region (ABR)**

Backbone modifications enable ADAR binding, and disable off-target editing

#### **Optimized sequence and chemistry define functionality**





Ensure bioavailability



ADAR: Adenosine deaminase acting on RNA, EON: Editing oligonucleotide, Nt: nucleotides

# Robust editing reported in the nervous system and liver



\*Undisclosed target. Conditions of the ACTB IPSC derived neurons experiment: gymnosis, 25µM, single dose, n=3-4, 2 weeks, dPCR and conditions of ACTB editing experiment in human primary hepatocytes experiment: gymnosis, 10µM, single dose, n=6, 48 hours, dPCR. Single doise, n=6, 48 hours, dPCR and conditions of the ACTB liver microtisus (LMTs): gymnosis, 1µM, constant dose, 3 pools of 24 LMTs here, between the mice *in vivo* experiment: intracerebroventricular (ICV), 250µg, single dose, n=6, 48 hours, dPCR and experiment: Single doise, n=6, 48 hours, dPCR and experiment: intracerebroventricular (ICV), 250µg, single dose, n=6, 48 hours, dPCR and experiment: intracerebroventricular (ICV), 250µg, single dose, n=6, 48 hours, dPCR and experiment: Single dotse, n=0 hours, n=0 hours, dPCR, mean, SD. Conditions of the actB mice experiment: Single dotse of 10 mg/kg, N=4, 1 week, ddPCR, mean, SD. Conditions of the non-human primare (NHP) in vivo experiment: intrateelar(IN), 12mg, single dose, n=3 hours, dPCR and 2 helfs not analyzable due to human error during injection procedure.

### Axiomer<sup>™</sup> creating a new class of medicines with broad therapeutic potential

Correction		Protein modulation	
Mus = Mus			
<b>Mutations correction</b> Thousands of G-to-A	Alter protein function or include protective variants	Disrupt >400 different types of PTMs	Change protein interactions
mutations, many of them described in literature	Modified proteins achieving loss- or gain-of-functions that help addressing or preventing diseases	Regulate protein activity, change localization, folding, preventing immune escape or slowing down degradation	Changes localization, folding, protein function or prevents immune escape of glycosylated tumor antigens
BROAD THERAPEUTIC POTENTIAL			





Target a wide variety of organs



Treat so-far undruggable targets

PTMs: Post-translational modifications.



### Pipeline

### **ProQR development pipeline**

	TARGET	DISCOVERY	NON-CLINICAL	CLINICAL	GUIDANCE	ESTIMATED POPULATION	
PROQR PROGRAMS							
CHOLESTATIC DISEASES	AX-0810 for NTCP				Entry into clinical trials in late 2024 / early 2025	~ 100K <sup>1</sup>	
CARDIOVASCULAR DISEASES	AX-1412 for B4GALT1				Entry into clinical trials in late 2024 / early 2025	~ 200M2	
	<b>AX-1005</b> for CVD					~ 200101-	
RARE NEURODEVELOPMENT DISORDER	<b>AX-2402</b> for Rett syndrome					~ 20K	
METABOLIC DISEASES	<b>AX-2911</b> for NASH					~ 16M	
	AX-0601 for obesity and T2D					~ 650M	
	AX-9115 for rare metabolic condition					~ 20K	
OTHERS	Multiple targets in discovery pipeline						
PARTNERED PROGRAMS	;						
<u> </u>	Initial <b>5</b> undisclosed targets	Progress undisclosed					
Lilly	Next <b>5</b> undisclosed targets	Progress undisclosed					
	Up to <b>5</b> potential additional targets						

<sup>1</sup>Approximately 100K people affected with Primary Sclerosing Cholangitis and Biliary Atresia in US and EU5. <sup>2</sup>Approximately 200 million people suffer from too high a level of cholesterol in US and EU5. *SLC10A1* is the gene that encodes for NTCP protein. CVD: Cardiovascular Diseases, NASH: Nonalcoholic steatohepatitis, T2D: Type 2 Diabetes.

References: Boonstra K, Beuers U, Ponsioen CY. J Hepatol. 2012 May;56(5):1181-1188; Karlsen TH, et al. J Hepatol. 2017 Dec;67(6):1298-1323; Dyson JK, et al. Lancet. 2018 Jun 23;391(10139):2547-2559; Sundaram SS, et al. Liver Transpl. 2017 Jan;23(1):96-109. Raghu VK, et al. Liver Transpl. 2021 May;27(5):711-718; NORD, 2019. Tsao CW, et al. Circulation. 2022;145(8):e153–e639. World Health Organization, World Gastroenterology Organization

### **AX-0810 for cholestatic diseases**

#### **RNA-editing therapy**

for Primary Sclerosing Cholangitis and Congenital Biliary Atresia



Cholestatic diseases have high unmet medical need. Patients accumulate bile acid in liver leading to fibrosis and ultimately liver failure.



Initial indications are **Primary Sclerosing Cholangitis** affecting adults and Congenital **Biliary Atresia** affecting pediatrics early in life. Both conditions have no approved therapies and require liver transplantation.



- Biliary Atresia is projected to affect ~24,000 pediatric individuals in US, EU and JP.
- **Primary Sclerosing Cholangitis** is projected to affect more than 80,000 individuals in EU, US and JP.



AX-0810 is a unique therapeutic approach leading to a potentially disease modifying therapy by targeting the NTCP channel which is responsible for majority of bile acid re-uptake in liver cells.



#### ProQR - Corporate Presentation

### AX-0810 reduces bile acids re-uptake into liver

RNA editing to a loss of function variant of NTCP can improve liver function

#### AX-0810 is a novel and "on target" approach reducing bile acid re-uptake into the hepatocytes

 Transient and controlled approach introducing a loss of function of NTCP

### AX-0810 can reduce bile acid load in the liver

- To alleviate associated pathology and symptoms in PSC and BA
- To prevent or delay the development of cirrhosis, organ failure and need for transplant

#### **AX-0810 therapy** *for cholestatic diseases*

BA: Bile acids, NTCP: Na-taurocholate cotransporting polypeptide, PSC: Primary Sclerosing Cholangitis. SLC10A1 is the gene that encodes for NTCP protein.



#### Reduced BA levels in the hepatocytes



\*LOF variant



### Well-defined development path for AX-0810



PRECLINICAL STAGE	EARLY CLINICAL	LATE CLINICAL
Preclinical models available with strong translatability into the clinic	Early insight on safety and target engagement using validated biomarkers	Clinical programs with disease specific endpoints for regulatory approval
<ul> <li>Translational models available</li> <li>Organoids models</li> <li>Animal models</li> <li>Proof of mechanism measures in animal models</li> <li>Serum levels of ALP and γ-GT</li> <li>Total bile acids in serum and liver</li> <li>Hepatic inflammation and fibrosis</li> </ul>	<ul> <li>Program with Phase 1 on healthy volunteers</li> <li>Validated biomarkers in cholestatic diseases</li> <li>Bile acids in serum, urine and feces</li> <li>Liver enzymes</li> <li>Serum cholesterol</li> <li>Disease specific biomarkers in preparation for next trials</li> <li>ALP for PSC</li> <li>Bilirubin for BA</li> </ul>	<ul> <li>Primary Sclerosing Cholangitis <ul> <li>Co-primary endpoint for regulatory approval:</li> <li>Reduction in ALP and</li> <li>Histological liver evaluation</li> </ul> </li> <li>Biliary atresia <ul> <li>Time to liver transplantation</li> <li>Mean change in total serum bilirubin levels, liver enzymes, bile acid levels, blood platelets and serum albumin</li> </ul> </li> </ul>

γ-GT: γ-glutamyl transferase; ALP, Alkaline phosphatase; BA, biliary atresia; BDL, Bile duct ligation; LMT, Liver microtissues; NTCP, Na-taurocholate cotransporting polypeptide; PSC, Primary Sclerosing Cholangitis

### AX-1412 for cardiovascular diseases

#### **RNA-editing therapy**

for cardiovascular disease (CVD)



#### Leading causes of death in the world

~18M people die from CVDs every year (32% of all global deaths) Despite therapies, the unmet medical need remains.



#### With projected increased number of patients

By 2035, >130 million adults in the US are projected to have some form of CVD with a total costs of \$1.1 trillion.



AX-1412 can become a stand-alone cardiovascular therapy that can also work synergistically with standard of care to further reduce risk of CVDs.



AX-1412 introduces a protective variant that reduces multiple independent risk factors for CVDs as was found in human genetics research.



## AX-1412 brings a novel approach to reduce residual risk for a potential cardiovascular event



RNA editing to a loss of function variant of B4GALT1 can have pleiotropic effect targeting two CVD risk factors

#### **B4GALT1** p.N352S protective allele Lipoprotein Fibrinogen **B4GALT1** mRNA Leads to hypo-galactosylation of apolipoprotein B100, fibrinogen ApoB FON AX-1412 is a novel and unique approach to address CVD ADAR Pleiotropic effects for cardiovascular protection (A-to-I-edit) **B4GALT** • Not suitable for knockdown technologies, as leads UDP-Gal 🔶 UDP to semi-lethality and severe development **B4GALT1** mRNA abnormalities in mouse studies EON AX-1412 can lower LDL-C and fibrinogen levels to reduce residual risk in cardiovascular diseases Prevent or delay the development of **B4GALT1\*** Protein

\*I OF variant

AX-1412 therapy for cardiovascular diseases

#### \*LOF variant

ADAR: adenosine deaminase acting on RNA, ApoB: Apolipoprotein B, CVDs: cardiovascular diseases, LDL-C: Low-density lipoprotein cholesterol. Reference: Montasser ME. et al., 2021 Science 374(6572):1221-1227.

#### ProQR - Corporate Presentation

cardiovascular events

### Well-defined development path for AX-1412



PRECLINICAL STAGE	EARLY CLINICAL	LATE CLINICAL
Preclinical models available with strong translatability into the clinic	Early insight on safety and target engagement using validated biomarkers	Clinical programs with disease specific endpoints for regulatory approval
<ul> <li>Organoids models for CVD</li> <li>Blood-derived myeloid cells and THP- 1 cells</li> <li>Cell-laden microtissue spheroids</li> <li>Animal models</li> <li>The Apoe-/- mouse model</li> <li>Proof of mechanism measures in animal models</li> <li>Serum lipid levels</li> <li>Atherosclerotic lesion area</li> <li>C-reactive protein (CRP) and Interleukin 6 (IL-6)</li> <li>Endothelial function</li> </ul>	<ul> <li>Programs with Phase 1 on healthy individuals</li> <li>Reduce potential signal-to-noise ratio as CVD patients have many comorbidities</li> <li>General CVD biomarkers</li> <li>non-HDL-C</li> <li>non-HDL-C</li> <li>Triglycerides</li> <li>Apoliprotein B</li> <li>Target specific biomarkers</li> <li>LDL-C</li> <li>Fibrinogen</li> </ul>	<ul> <li>Primary endpoints</li> <li>1. All-cause mortality and fatal CVD events or</li> <li>2. Composite endpoints (incl. fatal and non-fatal CVD events)</li> <li>Secondary endpoints</li> <li>Could consider using biomarkers as surrogate endpoints to reasonably predict treatment effects on outcome</li> </ul>

Apoe: Apolipoprotein E, CVD: cardiovascular diseases, HDL-C: High-density lipoprotein cholesterol; LDL-C: Low-density lipoprotein cholesterol, THP-1: human monocytic cell line

### **AX-2402 for Rett Syndrome**

#### Axiomer<sup>™</sup> technology

targeting the transcription factor MECP2 and potential to correct nonsense variants



Rett Syndrome is a **devastating and progressive neurodevelopmental disorder** caused by variants in the transcription factor Methyl CpG binding protein 2 (*MECP2*). There is a **high unmet need for a disease modifying therapy**.



Nonsense variants lead to **severe phenotypes.** They represent more than one third **of Rett Syndrome** cases and are projected to affect **20,000 individuals** in US and EU.



Rett Syndrome is **not a neurodegenerative disorders** and restoring levels of the MECP2 protein has shown to **reverse symptoms** in mice.



Axiomer has the potential to **restore the precise level of MECP2 protein regulatory function**, which is lacking in Rett Syndrome, and become a disease modifying therapy.

Krishnaraj R, Ho G, Christodoulou J. 2017. RettBASE: Rett syndrome database update. Hum Mutat 2017;00:1-10.





### Axiomer<sup>™</sup> RNA Editing Research Collaboration with Rett Syndrome Research Trust

- RSRT awarded ProQR approximately \$1M as a research grant for the initial phase of the project
  - EON design and optimization,
  - Evaluation in *in vivo* models for editing efficacy and MECP2 protein recovery
- Acting with a sense of urgency focusing on severe phenotype
- Following the initial discovery work, intent for expanded co-funding to enable continued development for the next phases
- Potential for further development for additional variants of relevance involved in Rett Syndrome



### Value creation strategy

*ProQR will develop its own pipeline and selectively enter into partnerships* 

#### **ProQR** Pipeline

- Build in-house pipeline based on Axiomer<sup>™</sup> RNA editing technology platform
- Initial focus on liver originated diseases



#### **Partnerships**

- Largely unencumbered platform, ProQR may selectively enter partnerships
- Lilly partnership with expansion announced December 2022 – total potential value of ~\$3.9B

### Well positioned

to advance Axiomer™



#### Science

- Deep understanding of basic science ADAR, oligos
- Optimization of editing oligonucleotides (EONs) for therapeutic development



#### **Axiomer<sup>™</sup> has broad applicability**

- Large number of potential therapeutic applications
- In vivo POC established in nervous system, liver



#### Advancing toward the clinic

- Extensive translational and developmental expertise with oligo modality
- AX-0810 and AX-1412 initial pipeline targets



#### Leading IP position

- Axiomer<sup>™</sup> is protected by >10 published patent families
- Continuously investing in expanding IP estate



#### Strategic partnership

- Lilly collaboration
- Rett Syndrome Research Trust
- Selectively form additional partnerships
- Optionality and multiple value creating opportunities



#### **Experienced leadership**

 Deep RNA, corporate finance, and business development expertise across Management Team, Supervisory Board, and Scientific Advisory Board

#### Strong balance sheet

- Q3 2023 cash €120.6 M
- Cash runway to mid-2026, excluding potential for additional BD-related upside



# ProQR® IT'S IN OUR RNA



### **Resource slides**



### HOW DOES ADAR WORK?

Explained in 5 minutes







# WHAT IS AXIOMER<sup>™</sup>?

Explained in 5 minutes





### **ProQR Leadership Team**



### Leading IP supporting ADAR-mediated RNA editing platform technology

- Axiomer<sup>™</sup> IP strategy commenced in 2014 with first patent application filings
- Currently 11 published patent families, comprising 29 national/regional patents
- Axiomer<sup>™</sup> IP portfolio is constantly expanding
- Oppositions/appeals and several Third-Party Observations have been filed against a variety of applications and patents in the Axiomer<sup>™</sup> IP portfolio, all by strawmen

### **Overview of Axiomer™ related patents**

Docket	Priority	Feature	Status
1 (0004)	17DEC2014	Targeted RNA Editing using endogenous ADARs	Granted BR <u>CA CN EP</u> IL IN <u>JP</u> NZ <u>US</u> ZA
2 (0013)	22JUN2016	Short EONs with wobble and/or mismatch base pairs	Granted <u>AU</u> IL <u>JP KR US</u> US
3 (0014)	01SEP2016	Chemically modified short EONs	Granted AU <u>CN EP JP KR</u> NZ <u>US US</u> ZA
4 (0016)	19JAN2017	EONs + protecting SONs (heteroduplex formation)	Granted <u>US</u>
5 (0023)	18MAY2018	PS linkages / chiral linkages ( <i>e.g.,</i> PS, PN)	Published
6 (0026)	11FEB2019	Phosphonacetate linkages / UNA modifications	Published
7 (0029)	03APR2019	MP linkages	Published
8 (0031)	24APR2019	Editing inhibition	Published
9 (0032)	13JUN2019	Benner's base (dZ)	Published Granted ZA
10 (0039)	23JUL2020	Split EONs	Published
11 (0045)	14FEB2022	PCSK9 editing	Published

In addition to the above, numerous patent applications are pending but have not yet been published.

ProQR expands its Axiomer<sup>™</sup> IP portfolio continuously.

### ProQR Axiomer<sup>TM</sup> IP

Broad coverage

- Axiomer<sup>™</sup> patent claims are broad and cover:
  - Any type of chemically modified oligonucleotide aimed at RNA editing of any possible target and any possible disease using endogenous ADAR
  - Specific targets
  - Oligonucleotides with chirally-controlled linkages
  - Oligonucleotides with all sorts of chemistries (also in the 'Central Triplet'), including **DNA**
- To note: claims directed to chemically modified oligonucleotides do not cover viral delivery of the oligonucleotide

### **Overview of key claims – 1**

Granted claims in the 1st Axiomer<sup>™</sup> patent family relate to (chemically modified) oligonucleotides that comprise:

- **A targeting portion** for binding to a target RNA incl. target adenosine
- A recruitment portion (hairpin structure) for recruiting endogenous ADAR to edit the target adenosine



EP 3 234 134 B1	<u>l</u> - Granted; appeal pending
<u>US 10,676,737</u>	- Granted
<u>US 11,781,134</u>	- Granted

Claim 17. A method for making a change in a target RNA sequence in a human cell, comprising the steps of:

- introducing into the cell an oligonucleotide construct that is sufficiently complementary to bind by nucleobase pairing to the target RNA sequence, wherein the target RNA sequence comprises a target adenosine;
- allowing the formation of a double-stranded structure of the oligonucleotide construct with the target RNA sequence upon base pairing;
- allowing the double-stranded structure of the oligonucleotide and the target RNA sequence to recruit an hADARI or hADAR2 enzyme naturally present in the cell;
- allowing the hADARI or hADAR2 enzyme to perform deamination of the target adenosine to an inosine in the target RNA sequence.

### **Overview of key claims – 2**

Granted claims in the 2nd Axiomer<sup>™</sup> patent family relate to oligonucleotides that do **not** have a hairpin structure, but instead have one or more wobbles and/or mismatches, and chemical modifications in the base, ribose sugar and/or linkage to increase stability and are still able to recruit **endogenous** ADAR to edit the target adenosine.



<u>US 10,988,763</u> - Granted <u>US 11,649,454</u> - Granted US 18/296,912 - Allowed

Target-specific claims:

- An AON capable of forming a double stranded complex with a target RNA in a cell, wherein: the target RNA encodes CFTR, CEP290, alpha1- antitrypsin (A1AT), LRRK2, or BDNF, or the target RNA is encoded by the IDUA gene
- The AON is complementary to a target RNA region comprising a target adenosine
- The AON comprises one or more nucleotides with **one or more sugar modifications**
- The AON does **not** comprise a portion that is capable of forming an intramolecular stem-loop structure that is capable of binding an ADAR enzyme
- The AON is shorter than 100 nucleotides
- The AON optionally comprises 1, 2, 3, 4, 5, 6, 7, 8, 9 or 10
  mismatches, wobbles and/or bulges with the complementary target
  RNA region, and, wherein formation of the double stranded complex
  between the AON and the target RNA results in the deamination of
  the target adenosine by an ADAR enzyme present in the cell

### **Overview of key claims – 3**

Granted claims in the 3rd Axiomer<sup>™</sup> patent family relate to oligonucleotides that do **not** have a hairpin structure, but have **chemical modifications** in the base, ribose sugar and/or linkage to increase stability and are still able to recruit **endogenous** ADAR to edit the target adenosine.



<u>US 10,941,402</u>	- Granted
<u>US 11,851,656</u>	- Granted
EP 3 507 366 B1	- Granted; opposition pending

An antisense oligonucleotide (AON) capable of forming a double stranded complex with a target RNA sequence in a cell, preferably a human cell, for the deamination of a target adenosine in the target RNA sequence by an ADAR enzyme present in the cell, said AON comprising **a Central Triplet** of 3 sequential nucleotides, wherein the nucleotide directly opposite the target adenosine is the middle nucleotide of the Central Triplet, wherein 1, 2 or 3 nucleotides in said Central Triplet comprise **a sugar modification and/or a base modification** to render the AON more stable and/or more effective in inducing deamination of the target adenosine; with the proviso that the middle nucleotide does not have a 2'-O-methyl modification.

### ProQR Axiomer<sup>TM</sup> IP

Summary

- ProQR's Axiomer<sup>™</sup> IP contains 3 early RNA editing platform patent families covering single-stranded oligonucleotides that recruit **endogenous** ADAR
- Oppositions/appeals and Third-Party Observations have been filed throughout these three patent families
- First (2014): oligonucleotides with a complementary (**targeting**) and a stem-loop (**recruiting**) portion
- Second (2016): oligonucleotides without a stem-loop structure but with one or more mismatches and chemical modifications
- Third (2016): oligonucleotides **without a stem-loop structure** but with specific chemical modifications in the '**Central Triplet**'

# ProQR® IT'S IN OUR RNA