

We are

An expert preclinical CRO delivering services for metabolic disorders and dedicated R&D studies to evaluate your drugs targeting, founded in 2003. We offer a wide range of tests to demonstrate the effectiveness of our clients' drug candidates.

ABOUT 20 YEARS OF EXPERTISE

specializing in obesity, type 2 diabetes, NASH / Fibrosis, diabetic nephropathy, inflammation, dyslipidemia and cardiovascular diseases for preclinical drug development, post launched studies and consulting.

We built our expertise over the years by developing innovative and predictive animal nutritional models to accelerate your drug discovery.





More than 10 of top 15 international biotech and pharma companies choose Physiogenex and gave the best chance of success to their lead compounds





Physiogenex and Cardiomedex

Innovative twin sister-CRO companies to evaluate your drugs on cardiometabolic disorders





- Physio-pathological predictive models
- Customized preclinical pharmacology studies
- In vitro, ex vivo, in vivo exploration







Benefits of our synergies

INVENTIVA AND LANIFIBRANOR: an example of unique synergy between Physiogenex and Cardiomedex

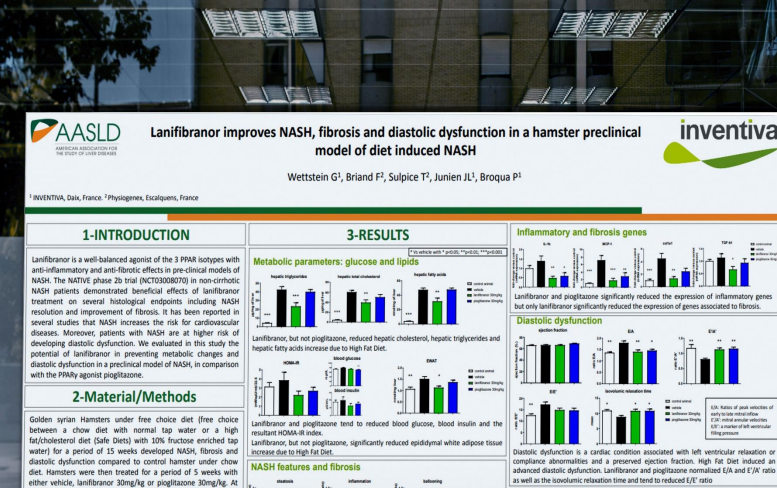
HIGH-VALUE RESULTS

Lanifibranor improves NASH, fibrosis and diastolic dysfunction in a Golden Syrian hamster model of diet induced obesity and NASH.

This new study realized at Physiogenex generated data that further support the development of Lanifibranor as a treatment for patients with NASH who are at cardiometabolic risk.

This leading anti-NASH drug is now evaluated in phase III (NATiV3).





compliance abnormalities and a preserved ejection fraction. High Fat Diet induced an

4-CONCLUSION

inventiva

Lanifibranor but not pioglitazone led to improvement in NASH and fibrosis suggesting that a panPPAR activation leads to a greater efficacy than a PPARy activation alone for the treatment of NASH-related liver features. Lanifibranor also markedly and significantly improved diastolic dysfunction similarly to pioglitazone. Activation of PPARy is therefore sufficient to correct the diastolic dysfunction in this model. However Lanifibranor as a pan-PPAR activator has the potential to limit the development of diastolic dysfunction via its beneficial effect on NASH but also by a direct effect on diastolic dysfunction through its PPARy component. This new data further support the development of lanifibranor as a treatment for patients with NASH who are at risk for cardiovascular diseases.

Contact information

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the end of the treatment, liver histology, genes expressions and biochemical analysis were performed. Diastolic dysfunction was evaluated by echocardiography and defined as an absence of change in left ventricular ejection fraction, an increase in E/A and E/E' ratio and a decrease in E'/A' ratio as well as in isovolumic



Group 2: Free choice diet + vehicle p.o. QD; n=10 Group 3: Free choice diet + lanifibranor p.o. QD; n=10 Group 4: Free choice diet + pioglitazone p.o. QD; n=10

High Fat diet produced a significant increase in steatosis, liver inflammation and fibrosis but not ballooning.

Lanifibranor, but not pioglitazone, significantly decreased steatosis, inflammation, fibrosis and consequently the total score taking into account the 3 NASH features and

A management team of experience & expertise



Thierry Sulpice, Ph.D

CEO, CSO, Founder

- 9 years at Sanofi –Aventis
- In charge of drug discovery optimization and preclinical development for projects targeting the cardiometabolic syndrome.
- PhD in Physiology and Pharmacology University of Grenoble, France.
- Founder and CEO/CSO of Cardiomedex





Director, Research and Business Development

- Expert in metabolic diseases since 2007.
- Worked with Novo Nordisk during his PhD in Nutrition and Metabolism, and with Glaxo Smith Kline and Merck as a postdoctoral fellow with Dr. Dan Rader at Upenn, Philadelphia.





Highly skilled metabolic disease experts working towards innovation, client satisfaction and quality compliance



Estelle Grasset, Ph.D – Project Manager



Natalia Breyner, Ph.D - Project Manager



Emmanuel Brousseau – team manager



Marjolaine Quinsat – team manager



Sura Setau – quality department manager



Julie Arasse – administrative department manager



Pr. Rémy Burcelin – Scientific expert consultant





Drug efficacy testing pipeline

In vitro/ex vivo In vivo In vitro/Ex vivo *In vivo* efficacy: *In vivo* drug Your small platform: Ex vivo platform: screening: obese/type 2 classical cell lines isolated tissues molecules/peptides normal or disease diabetic/MASH/nephropathy/ (adipose, muscle) /nutraceutics and human animal models HFpEF/fibrosis animal models organoids Cell Blood parameters Glucose Blood/tissue biochemistry

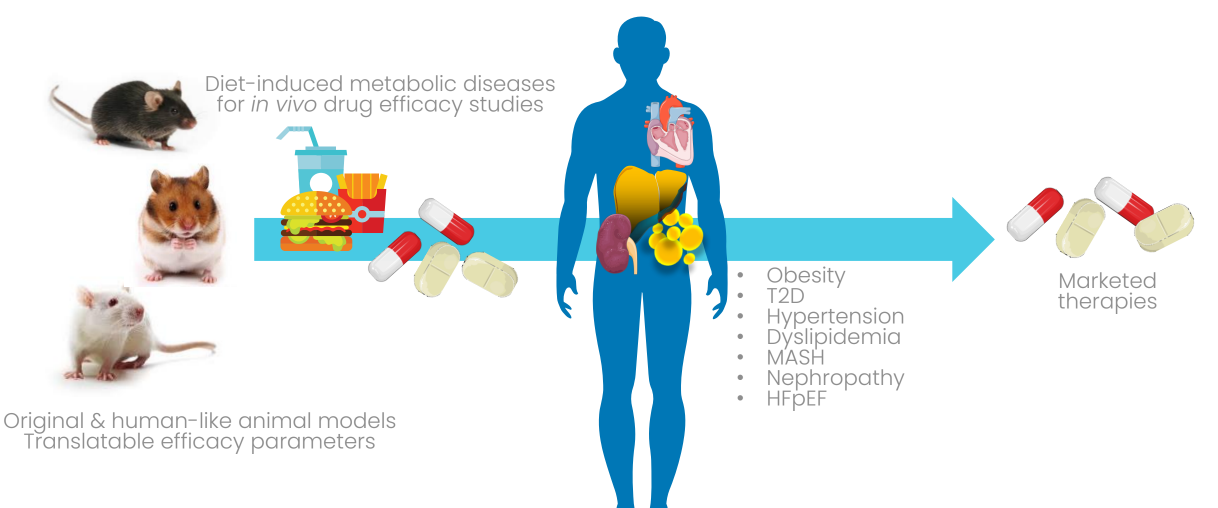
- viability/cytotoxicity
- Gene expression
- Protein/lipids levels
- uptake
- Fatty acids oxidation
- Glucose/insulin/lipid tolerance tests
- Kidney function
- NAFLD/NASH

- Glucose/insulin/lipid tolerance tests
- *In vivo* lipogenesis
- Hyperinsulinemic euglycemic clamp
- Kidney glomerular filtration rate
- Echocardiography
- Liver/lungs/kidney/heart histology





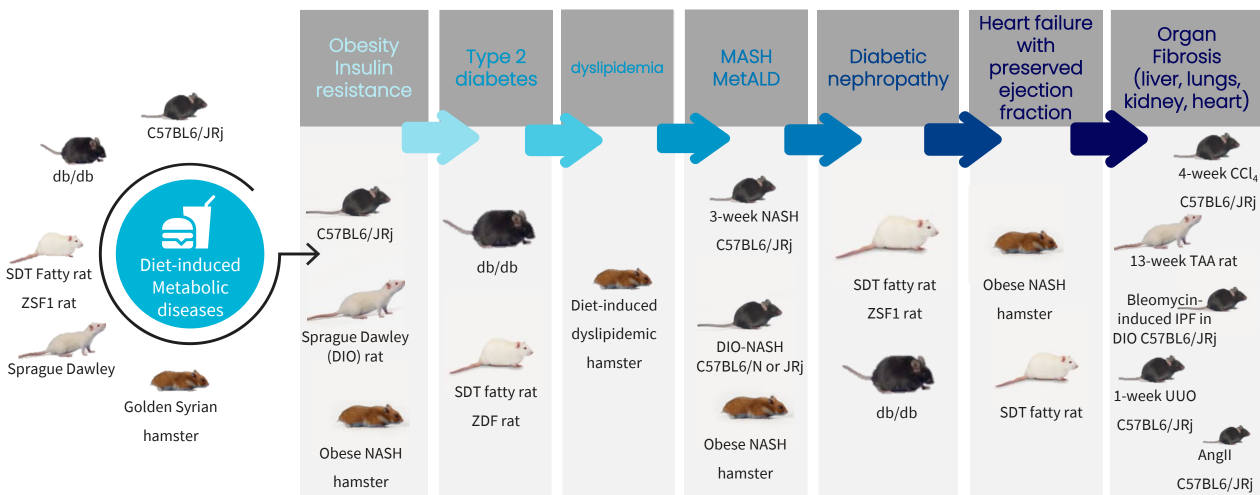
Translational data: Derisk your drug efficiently and early.







Innovative animal models developed to optimize and expedite your drug efficacy studies







Platform To fully characterize your drug candidates mechanism of action and efficacy



Colorimetric assays

- Plasma and hepatic Lipids (cholesterol, triglycerides fatty acids)
- Lipoproteins (direct LDL-C and HDL-C assays, lipoprotein FPLC profile, apo A-1, apoB)
- Plasma ALT /AST
- Total ketone bodies
- Glycerol
- Phospholipids
- Albumin
- Creatinine
- Faecal total cholesterol and bile acids



ELISA and multiplex assays:

- Hormones (insulin, glucagon, adiponectin, Leptin, etc.)
- Cytokines panel (IL-1b, IL-6, MCP-1, TNF-alpha, etc.)



Western Blot analysis

(WES technology) for any protein. on any sample/tissue.



Microbiome, lipids, bile acids profiling Gene expression by qPCR on any tissue:

- Lipogenesis
- Glucose
 metabolism
- ER and oxidative stress
- Inflammation
- Fibrosis



Histology analysis:

- H&E
- Sirius Red
- Masson Trichrome
- ORO
- PAS staining
- Immunohistochemistry (F4/80, CD68 ED1, collagen III alpha-SMA, etc.)
- NAS scoring (steatosis, inflammation, hepatocyte ballooning tibrosis) Nephropathy histopathology scoring (glomerulosclerosis interstitial fibrosis, etc.)

+ 25 publications demonstrating Physiogenex expertise in high impact scientific journals

Arteriosclerosis, Thrombosis, and Vascular Biology

JOURNAL OF THE AMERICAN HEART ASSOCIATION

Upregulating Reverse Cholesterol Transport With Cholesteryl Es Inhibition Requires Combination With the LDL-Lowering Di Dyslipidemic Hamsters

François Briand, Quentin Thieblemont, Elodie Muzotte and T

Diabetes Volume 65, July 2016

Arterioscler Thromb Vasc Biol. 2013;33:13-23; originally published on doi: 10.1161/ATVBAHA.112.252932

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Peptides 114 (2019) 44–49

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Peptides

journal homepage: www.elsevier.com/locate/peptides

BZ043, a novel long-acting amylin analog, reduces gastric emptying, food intake, glycemia and insulin requirement in streptozotocin-induced diabetic



Bisman Biopharmacanical SA, Rau Viccomè de Projú 6,523, 9th flore, Rio de Janviero, R.J. 224/10.003, Parail Federal University of Bio de Janviero - PREJL, CSS, Back J. and he Francisky, 224/15-500, Rio de Janviero, R.J. Brazil National Institute of Science and Technology for Structural Biology and Bisineaging (RNBEN BICT), Federal University for the de Janviero, Rio de Janviero, Rivald Laborators fro Marcomoleculos, (AMAC DIMAY). Destallors de Marcologo, des de Herroleco, Oscilla, Vanderonicalos, (AMAC DIMAY). Destallor Astronicalos de Marcologo, del Service de Marcologo, del Service de Technologo de Parail Technologo

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European Journal of Pharmacology

2032

journal homepage: www.elsevier.com/locate/eipha

Full length article

Nephropathy in diabetic db/db mice is accelerated by improved by the SGLT2 inhibitor dapagliflozin

Sisse Andersen Nørgaard^{a,c,1,*}, François Briand^{b,1}, Fredrik Wolfhager Elisabeth Douglas Galsgaard^a, Henrik Søndergaard^a, Dorte Bratbo Sø

- ^a Pharmacology, Novo Nordisk A/S, Novo Nordisk Park, Måløv, Denmark
 ^b Physiogenex S.A.S, Prologue Biotech, 516 rue Pierre et Marie Curie, 31670, Labège, France
- ^c Department of Veterinary Disease Biology, University of Copenhagen, Grønnegårdsvej 15, 1870, Frederiksberg C, Denu

Elafibranor improves diet-induced nonalcoholic steatohepatitis associated with heart failure with preserved ejection fraction in Golden Syrian hamsters

François Briand ^{a.*}, Julie Maupoint ^b, Emmanuel Brousseau ^a, Natalia Breyner ^a, Mélanie Bouchet ^a, Clément Costard ^b, Thierry Leste-Lasserre ^c, Mathieu Petitjean ^d, Li Chen ^d, Audrey Chabrat ^e, Virgile Richard ^e,

François Briand,¹ Eric Mayoux,² Emmanuel Brousseau,¹ Noémie Burr,¹ Isabelle Urbain,¹ Clément Costard,¹ Michael Mark,² and Thierry Sulpice¹

Empagliflozin, via Switching Metabolism Toward Lipid Utilization, Moderately Increases LDL Cholesterol Levels Through Reduced LDL Catabolism

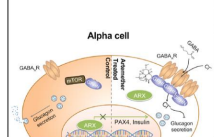
Diabetes 2016;65:2032-2038 | DOI: 10.2337/db16-0049

Cell

Artemisinins Target GABA_A Receptor Signaling and Impair α Cell Identity

Graphical Abstract

American Heart



Authors

Jin Li, Tamara Casteels,
Thomas Frogne, ..., Patrick Collombat,
Jacob Hecksher-Sørensen,
Stefan Kubicek

Article

Correspondence

skubicek@cemm.oeaw.ac.at

In Br

The anti-malarial drug Artemisinin can drive the in vivo conversion of pancreatic α cells into functional β -like cells through enhanced GABA signaling and may have potential as a therapeutic for diabetes.







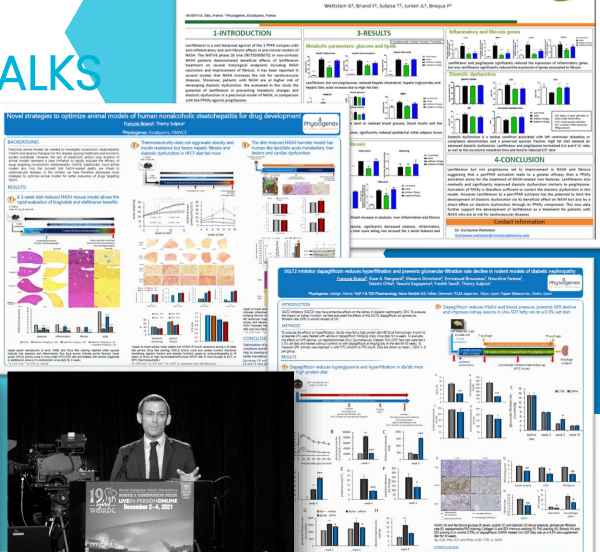
+120 POSTERS and TALKS

in international conferences

To make our innovative new services and models known for your drug evaluation success, early.







Lanifibranor improves NASH, fibrosis and diastolic dysfunction in a hamster preclinical model of diet induced NASH





They trust us

















































































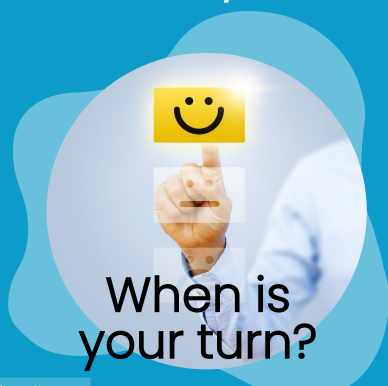
High success rate customer satisfaction over the last 3 years: 97.5%

« I can't thank you enough for all your great work and support! It was really nice collaborating with you all and I am very much looking forward to the next collaboration with you! »

- US Pharma customer

« Great working with Physiogenex: great competence in the area, straight answer and very good team available to support all our needs »

- European biotech customer



"Excellent scientific advice provided for the study design. Studies tailored for small biotechs with limited budget. Timelines met, rigor in execution. Indeed a great partner for drug development in the metabolic disease area." – US biotech customer

"High level of expertise and timely discussions duringthe study plan and execution

Very co-operative" - India Pharma customer

"Once again, I would like to appreciate for all your assistance during the study execution and all technical issues that Physiogenex experts helped us to understand. Physiogenex is a very strategic partner and in the future, we certainly will consider you to perform our preclinical studies." – South American Pharma customer

"We really appreciate what you did for our study. A new assay development was successful. The study report was well-documented."

- Korean Pharma customer

What if we work together? Major steps ahead



FIRST QUOTATION

- Scientific consulting
- Strategic approach
- Price
- Timeline (study start)
- Study design

CONFIRMATION

- Price
- Timeline (study start)
- Study plan

STUDY RUN

- Data production
- Project management

STUDY REPORT

- Data
- Interpretation
- Scientific & strategic recommandations



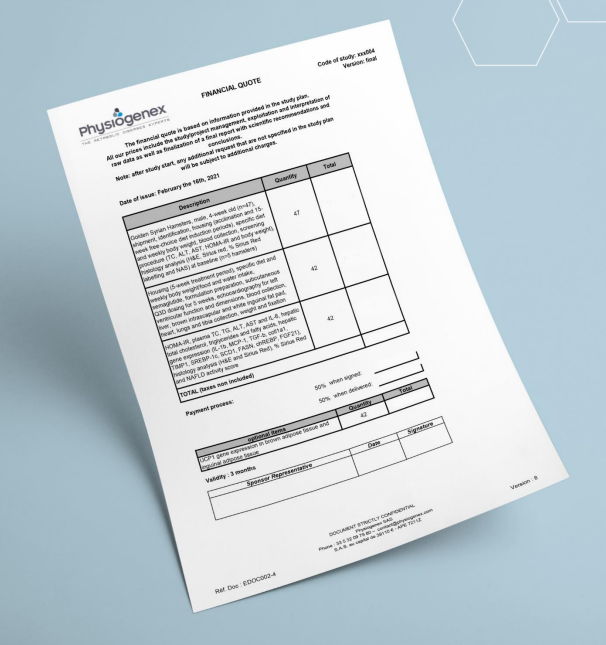






Quotation and study confirmation

- Discussion with our experts to setup the best cost-effective study design for your drug efficacy project
- An experimental design and quotation are then proposed with study timelines
- 3. Upon you agreement, a study plan is issued and once signed your study starts (2-3 weeks average)





Study run

- ✓ Studies led by a dedicated project manager
- ✓ Fast and efficient communication: Weekly follow-up
- Raw data delivered as soon as available







Study report

- A clear description of your study results provided on Word format for your review
- ✓ Includes statistical analysis, data expertise and recommendations to go further with your drug development
- Reporting includes qualitycontrolled data on Excel format, statistics (GraphPad)





Partening with us to launch Your new therapies successfully

Obesity - Type 2 diabetes - Dyslipidemia Inflammation - Diabetic nephropathy - NASH -Fibrosis and cardiovascular complications

- Over 20 years of experience/expertise in drug development with major pharmas & biotechs
- 97.5% customer satisfaction rate
- Unique, translational preclinical models published in major scientific journals
- Time and cost-effective preclinical studies to reach the clinical stage development
- High added value partnering with Cardiomedex
- As R&D studies providers we propose and set-up dedicated tolls/animal models to target your needs







Physiogenex and you



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