

# Bio-Modeling Systems The R&D booster for life Sciences discoveries

# Bio-Modeling Systems

The Mechanisms-Based Medicine Company



We changed the discovery paradigm to create novel medical meanings from unreliable heterogeneous sources of data

Short corporate Presentation
This is not a pitch presentation
This document is for download only
We added the necessary details and explanations in the slides to help the reader

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### Why do we need to change the discovery paradigm?



1-The industry is under pressure by too high failure rates and payers no more willing to pay



SANOFI













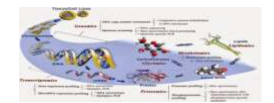








2-New players are entering the global domain, but no one has an integrative picture, because they are still "limited" by their own regulations, experiences, histories.



3-Pharma industry already invested for decades in Omics technologies and Systems Biology programs for no relevant results. The key reason: Life mechanisms are complex not complicated!



4-The future should be digital and biology but, Artificial Intelligence (AI) MUST follow rules while humans do not! Currently the "Garbage in garbage out" reality is not correctly treated.

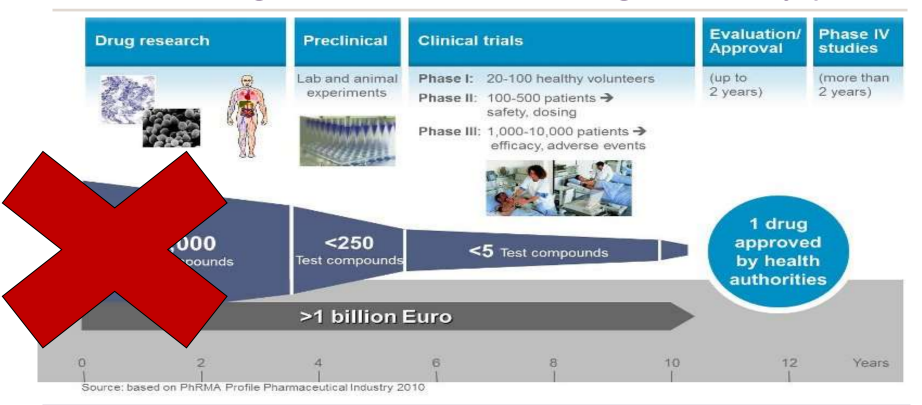


5-The unreliability of scientific and clinical publications is increasing. "Many published research findings are false or exaggerated, an estimated 85% of research resources are wasted." (Stanford university), and the valuable negative results are not published.

So why despite massive investments in technology and IT, the success rate of the industry is still declining? The challenge is not a question of technologies only!



### The misleading classical Pharma drug discovery process

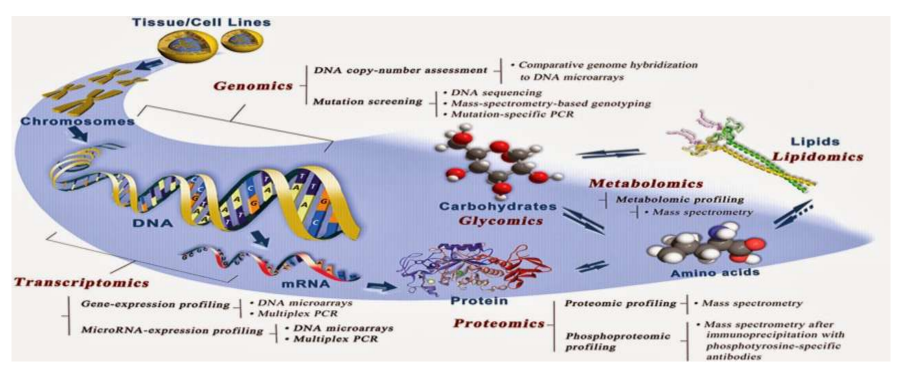


This Big Pharma R&D model focused on testing new patentable compounds for novel targets based on KOL concepts is clearly not performant!

- 1. Is 1 billion € per drug approved a fatality or a Discovery paradigm failure?
- 2. How are KOL concepts generated and evaluated?
- 3. Has Evidence based Medicine reached its limits with chronic complex human diseases?
- 4. Are the data produced and the scientific publications reliable and robust enough to feed algorithms that MUST follow rules?



# The discovery of molecular biology, and the endless Omics story that began in the 70's generated 3 major side effects



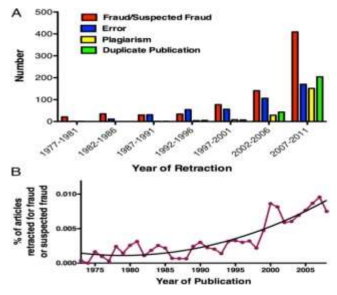
- 1. Medical research focused on patient's diseases became life sciences research driven by data, technologies and IT outputs.
- 2. The leadership switched from MDs & biologists to molecular & IT scientists.
- 3. The discovery issue: Tools, algorithms & concepts from Digital and Technologies giants, valid for complicated systems, cannot address complex systems such as life



# The unreliability of scientific and clinical publications is unacceptable and increasing

- 85% of research resources are wasted. Currently, many published research findings are false or exaggerated (John P. A. Ioannidis METRICS Institute Stanford University. <u>Published</u> in Plos medicine 2014)
- 90% of 53 studies were not reproducible.
   Amgen's scientists couldn't reproduce the findings of 53 "landmark" articles in cancer research (C. Glenn Begley ex Amgen. Published in Nature, 2012)
- 79% of 67 projects were not reproduced by Bayer's scientists trying to reproduce the findings of 67 target-validation projects in oncology, women's health, and cardiovascular medicine. (Florian Prinz, Thomas Schlange and Khusru Asadullah Reu Bayer. Published in Nature discovery 2011)

Number of retracted articles for specific causes by year of retraction

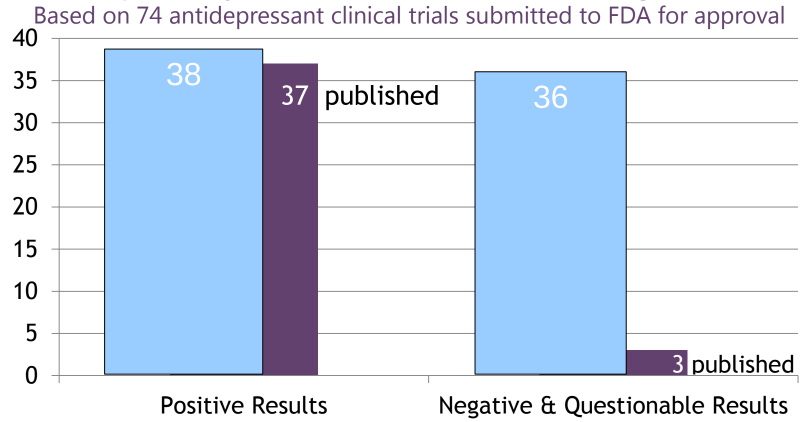


Ferric C. Fang et al. PNAS 2012;109:17028-17033

The "garbage in, garbage out" reality demonstrates that a wrong hypothesis, even if generated or treated by the best Digital and IT technologies, remains a wrong hypothesis



# Publications do not represent the real knowledge especially when the results are negative



clinical trials submitted to FDA compared to those published. An enormous bias. A critically misleading issue if not contextualized



# The Life-modeling issue illustrated

1-If you dream of creating the first operational model of a bird...



2-... a "basic" living Complex System that not only flies...

3-Be sure to use the appropriate modeling concepts & tools. If you don't ...



4-...you'll get a Complicated "Cartesian" system. It flies... But the major issue is that, for modelers, **this is a bird!**\*

The challenge is clearly not a question of technologies only! Even with expensive efforts, this model will never become a "bird"!

A valid solution must address both the complexity of life's mechanisms and the unreliability of scientific and clinical publications to create novel & pertinent medical meanings

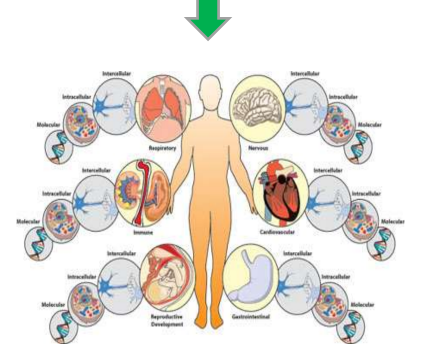
<sup>\*</sup> Based on this model,1) when birds lay eggs, they explode; 2) the rear end of a bird is extremely hot when it flies; 3) a bird has three legs, etc.... You may think this stupid, but it is what is being done with systems biology.

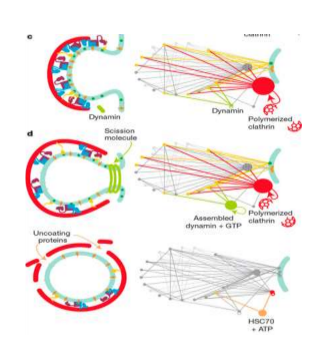


# What leads to Therapeutic Success?

The success of a therapeutic approach largely arises from the coherent manipulation of a physiological system as a whole

and not from that of a target in a molecular context.





Therefore, any given medical problem should be approached from a "systems medicine" standpoint In this context, novel therapies can be combinations of drugs, nutriments, devices, e-health, etc... (while targeted therapies belong to the "target in a molecular context" concept)



### The mechanisms-Based Medicine Principles

The Global Discovery stepwise approach places diagnostic / therapies / prevention solutions & validation processes *in the right order*:

#### 1-DISEASE\*

- •Redefine the definitions and descriptions of the physiopathology of the disease/disorder/syndrome with physiologists, clinicians and patients feedbacks.
- \* Do not forget but integrate that for a disease/disorder/syndrome, similar symptoms can have very different functional origins, while similar dysfunctions can produce different symptoms. Download\*\* the dedicated presentation with the psychiatry case study

#### 2-MECHANISMS

•Discover the causal versus symptomatic mechanisms of the disease/disorder.

#### **3-BIOMARKERS**

•Indirectly based on causal mechanisms, identify relevant biomarkers or specific biomarkers combination/signatures (biological, imagery, physical signals, etc....) that could measure defined mechanistic deregulations at different stages of disease/disorder progression.

#### **4-TARGETS**

•based on the causal mechanisms, identify what could be the best targets (not only one) to specifically address the causative deregulations.

#### **5-SOLUTIONS**

•We harness the mechanisms to propose the most practical solutions addressing the relevant mechanistic deregulations.

•It is important to notice that the proposed solutions, integrating diagnostics, therapies & patients follow-up, can be new drugs, combinations of existing drugs, nutriments, devices, e-health, disease prevention tools and services, etc ...

#### **6-VALIDATION**

•Global validation loop at each steps of the process: Integrate the results from e-R&D or e-Health experimentations into the validation process to improve global patient and disease/disorder follow-up.

Understanding and validating the mechanisms of a disease/disorder becomes the first objective. Finding the most adapted solutions is a necessary consequence of the first objective



# CADI™ Discovery Principles

| "Mechanis | sms-Based  |
|-----------|------------|
| Medicine  | Principle" |

- ☐ Answers the failures of the pharma Research Process & of the "KOL dominant thinking" by fostering the discovery & selection of novel concepts.
- ☐ Need to separate causal mechanisms understanding from solutions discovery.
- ☐ Discovery of lower risk & cost effective multi-technologies and integrated solutions.

# "Architectural Principle"

- ☐ Mechanisms of life are complex, non-linear and integrative.
- ☐ Heuristic Modeling (the Architects) searches for satisfactory solutions to describe the mechanism of a poorly defined system.
- ☐ Mathematical Modeling (the Engineers) simulates, when correctly described, the dynamics of the system .

#### "Negative Selection Principle"

- ☐ "An estimated 85% of current published research findings are false or exaggerated" J.P.A Joannidis, 2014 Stanford University [PLoS Med]).
- ☐ "It is always possible to demonstrate a statement to be false" Karl Popper.1963.
- Only working hypotheses that resist destruction are worth retaining.

# "4 Steps Validation Principle"

- ☐ Only mechanisms that resisted the "Negative Selection Process" are worth testing.
- ☐ Iterative validation process with the necessary scientists, clinicians, MDs, and patients.
- ☐ Construction of dedicated experimentations to evaluate the predictions of the model.
- □ Necessary bridge between R&D, clinic and real life.

# "Integrated Solutions Principle"

- ☐ Can be combinations of drugs, diagnostics, medical devices, nutriments, e-health, cosmetics, for treatments, and prevention programs, etc. ...
- ☐ Access to end user is strategic, and digital technologies are essentials to connect all the components of the solutions.

CADI™ Discovery is the world's first and, to date, only operational platform that addresses life's mechanisms complexity and the unreliability of scientific and clinical publications by combining the strengths of human and artificial intelligences in the right order.



# Mathematical & Heuristic approaches can be complementary, provided they are harnessed in the proper order.

Mathematical approaches are of limited usefulness when applied to <u>poorly</u> <u>defined</u> <u>multicellular</u> <u>physiological</u> <u>systems</u> because they cannot efficiently <u>reveal</u> & <u>define</u> the functional states within such a system (cross-talks alterations, etc...).



But heuristic approaches are very efficient at doing precisely this.

Heuristic models are of limited usefulness when addressing the <u>dynamics</u> of <u>defined</u> complex <u>physiological</u> pathways structures and cross-talks because they are not open to mathematical manipulations.



But Mathematical models are very efficient at doing precisely this.

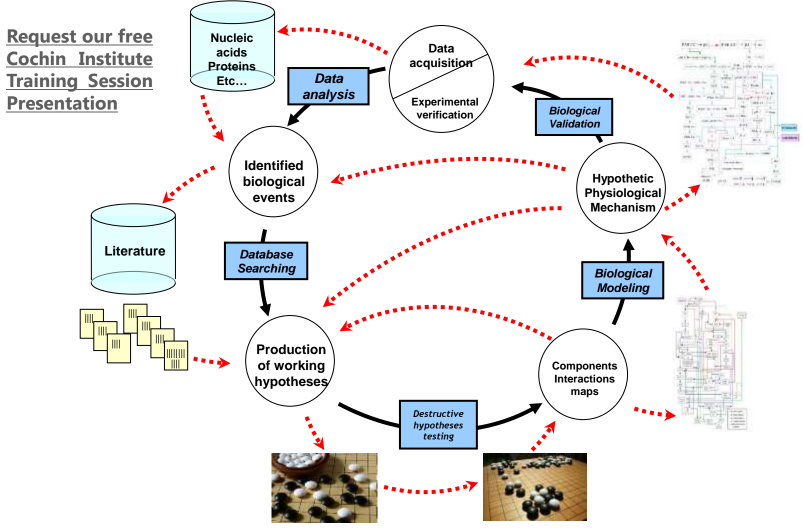
To efficiently address the translation of systems biology to clinical & medical interventions (dominated by patient's data heterogeneity and largely unstructured documents), ways to achieve synergy between Heuristic and Mathematical approaches can be effectively designed.

We apply first Heuristic modeling and then propose the outputs for Mathematical modeling when the system is correctly described



# The CADI™ Integration workflow

More details in the Full Presentation with CADI full Description, publications and the 10 CADI™ programs & POCS

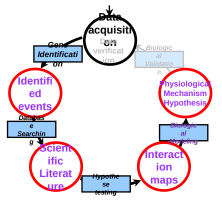


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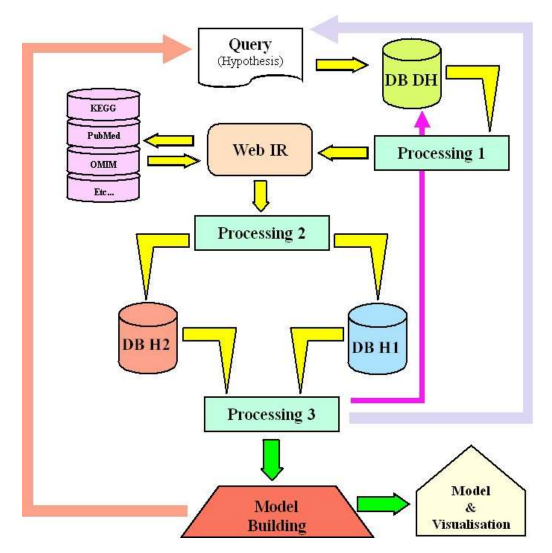
### The CADI™ Integration & Modeling Process

More details in the Full Presentation with CADI full Description, publications and the 10 CADI™ programs & POCS



This iterative process does three things:

- It largely resolves the coherence issues attached to the classical approach;
- It reveals hitherto unknown mechanisms/processes, and
- It allows the translation of systems biology to clinical & medical interventions.

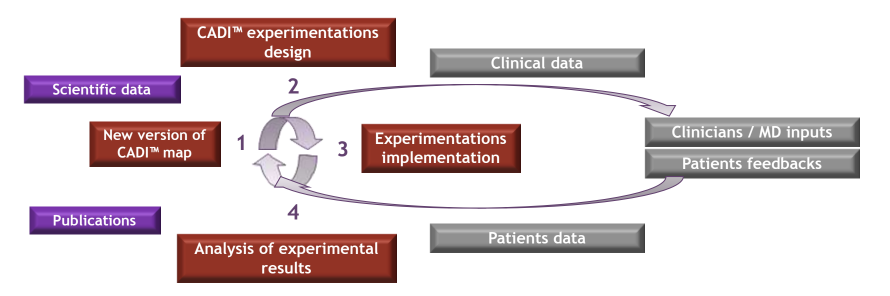




## CADI™ Discovery Global validation Principle

exploiting Smart Data (contextualized, with patients based lines, related to mechanisms data)

CADI™ Discovery from bench to bed to real patient health processes



E-R&D Continuum/Synergies E Health

#### Information technologies

Data acquisition, Simulation, collaborative, data Storage, Big Data, Smart Data, Mobility

**CADI™** Smart Data (contextualized, with patients based lines, related to mechanisms)



### An experienced multidisciplinary founders' team



Dr. François Iris (PhD), Chairman, CSO-CTO - Heuristic modeling specialist

**French-New-Zealander.** Geneticist, physiologist & molecular biologist. **40 years of experience in life sciences in academia and industry**: Dept. of Medicine University of Otago, The Christchurch School of Medicine (NZ) Millennium Pharmaceuticals' (USA) collaborator of Nobel Laureate Prof. Jean Dausset. Inventor of CADI<sup>TM</sup> and of new technologies in molecular biology. MRC Overseas fellow, Member of H.U.G.O., Wellcome Trust; etc..



Manuel Gea, C.E.O & VP R&D Information Systems – Operational Research & business development specialist

**30 years of experience in IT and life sciences**. Scientific Engineering Degree from Ecole Centrale Paris. Various experiences R&D and business from consumer goods Industry to cosmetics, biotechnology & pharmaceutical companies: Colgate-Palmolive McKinsey, Boehringer Ingelheim, HemispherX Biopharma, Pherecydes-Pharma, BMSystems; etc..



Gérard Dine (MD, PhD), Chief Medical Officer - Physician, biologist

**35** years of experience in clinical and medical research. Head of hospital's Hematology Dept. Former President of the Institute for Sports Medicine; etc..



Paul-Henri Lampe, CIO & Systems Integration Director - Systems Integration specialist

**French-American. 20 years of experience in Systems integration in healthcare**. Scientific Engineering Degree Ecole Centrale Paris. Former IBM Systems Integration Manager. Former Information Systems Manager, Hospital in Paris.



Pablo Santamaria, IT & Internet Systems Director - Internet technologies specialist

**30 years of experience in Internet technologies and life sciences**. Scientific Engineering Degree from Ecole Centrale Paris, Founder and President of the computing firm Formitel, Glaxo Pharma (Evreux, France)

### A selection of Prestigious collaborative R&D partners.

7 publications, 2 reference books, 4 patents



























#### BMSystems' cost-effective, lower risk, patentable novel integrated solutions

| Program Domain               | Partners | CADI™<br>compliance | CADI™<br>vers. 0 | Ind. Valid. | Patents<br>/ Publi. | First Proof of<br>Concept<br>(POC) | Mid scale or preclinic. P.O.C. | Business<br>launched |
|------------------------------|----------|---------------------|------------------|-------------|---------------------|------------------------------------|--------------------------------|----------------------|
| Infection-Immunology         |          |                     |                  |             |                     |                                    |                                | Phase I/II           |
| CNS-PNS                      |          |                     |                  |             |                     |                                    |                                | Phase Ib             |
| Oncology                     |          |                     |                  |             |                     |                                    |                                |                      |
| Metabolism                   |          |                     |                  |             |                     |                                    |                                |                      |
| <b>Dermatology/Cosmetics</b> |          |                     |                  |             |                     |                                    |                                |                      |

**BioProcesses / Synthetic Biology** 

#### BMSystems' first outstanding POCs completed (World's first in vivo validations, 2 spin-offs)







**World's first** *in vivo* **validation** of the mechanisms of Creutzfeldt-Jakob disease pathogenesis & progression. Two Awards (2009 and 2010)



<u>Pherecydes-Pharma</u>: **BMSystems' spin-off, 2006**, novel M.R. antibacterial nano-agents biotherapies), two indications: <u>Multi-resistant Skin infections</u> in Phase I/II. and osteoarticular infections.

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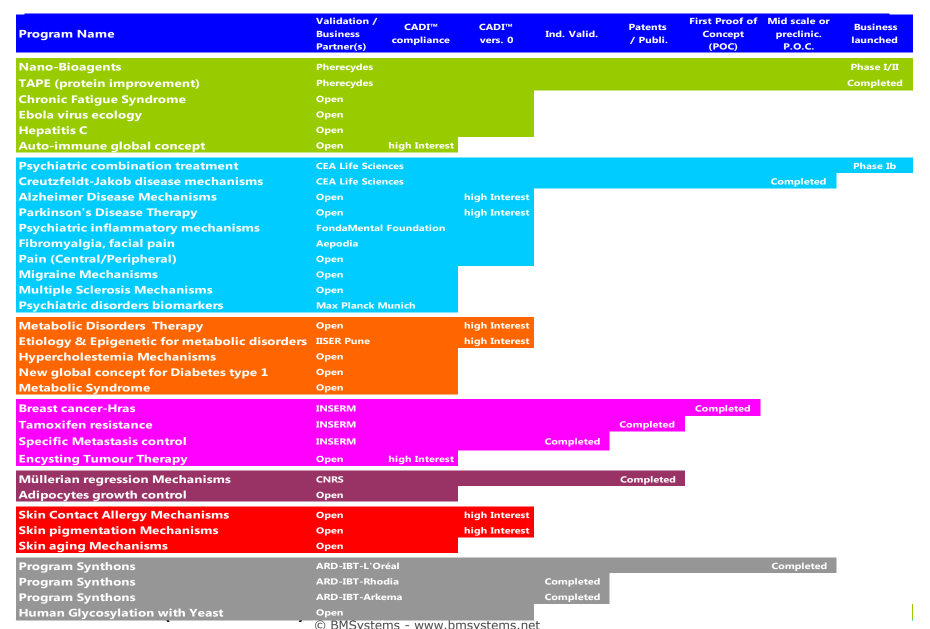


Theranexus: CEA's spin-off, 2013, (Therapeutics CLASS co-patent, WO/2010/029131, innovative combined therapies for Psychiatric disorders). Phase I success & Phase Ib launched for narcolepsy, a rare disease

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#### BMSystems' cost-effective, lower risk, patentable novel integrated solutions





# What we already achieved with our partners This list excludes our contractual research programs with our clients



CEA: Project "Creutzfeld-Jacob Disease CJD" World's first in vivo validation of the mechanisms of Creutzfeldt-Jakob disease pathogenesis & progression. US, EU & French Awards; Awards (2009 and 2010).

Successfully completed; 1 therapeutic CLASS co-patent CEA/BMSystems; 1 publication.



<u>Theranexus</u> **CEA's spin-off, created in 2013**, exploiting the <u>Therapeutics CLASS co-patent</u> <u>WO/2010/029131</u> **CEA/BMSystems**, <u>innovative combined therapies</u> for Psychiatric disorders. <u>Phase I success</u> & Phase Ib in 2015 in narcolepsy, a rare disease.



<u>Pherecydes-Pharma</u> BMSystems' spin-off created in 2006, novel M.R. anti-bacterial nano-agents biotherapies 3 patents. Two indications: <u>Multi-resistant Skin infections</u> in Phase I/II in 2015 and osteo-articular infections.



Max Planck Institute (Munich): Project "Chronic Anxiety".

Successfully completed; 3 publications & a Reference Book "Biomarkers for Psychiatric disorders" chapter 19.



INSERM: 3 Projects "Tumoral Progression"; "Therapeutic Resistance"; "RGD 15 & Metastasis".

All 3 successfully completed, **3 publications**.



CNRS: Project "Müllerian Regression" Tissue differentiation Successfully completed, **1 publication**.



Foundation FondaMental: Project "Bipolar Disorders & Schizophrenia". Immuno-inflammatory hypothesis. On going, **1 publication pending** 



L'OREAL Arkema, Rhodia/Solvay ARD: "Synthons" IAR cluster Industrial Biotech program

Feasibility study Completed 16 molecules evaluated, 2 strains built, 1 program with 1patent pending

Skin Homeostasis: Reference book "Computational Biophysics of Skin" chapter 15 with Dr. Querleux (L'Oréal)



Centre of excellence in Epigenetics IISER Pune India: Project "Etiology & Epigenetic for metabolic disorders" Etiology & Epigenetic for metabolic disorders, on going **1 publication pending** 



### BMSystems' solutions to critical & high impact issues

#### GO-NO GO decision before product acquisition or for portfolio risk analysis.

- **Why**: With a success rate around 10%-15% in the Pharma industry, be smarter by not investing in the wrong asset, increase your ROI.
- Objective: Identification and evaluation of the potential hidden issues in acquisitions. Investment savings. Refine acquisition value.
- Who is interested: VCs, Angels, TTO, Corporate funds, Consulting companies and life science industry managers.

#### **GO-NO GO decision before next development phase.**

- Why: When pros and cons are really mitigated and no more robust facts available from existing expertise.
- **Objective**: Address the possible safety and efficacy issues before launching the next phase. **Costs and time/resources savings**.
- Who is interested: Pharma, Diagnostics experts, Biotech, e-Health and cosmetics, preclinical and clinical development managers.

#### R&D program Rescue for a program facing critical issues during its lifetime.

- **Why**: There are multiple reasons for specific problems. Some can be addressed only when functionally understood.
- Objective: Identify the roots of problems and try to propose a pertinent solution. Investments & costs savings.
- **Who is interested**: Pharma, Diagnostics, biotech, e Health and cosmetics, preclinical, clinical and post-marketing development managers.

#### External R&D "B plan" program when the "A plan" cannot be rescued.

- Why: The reasons for failure are systemic, the concepts or the solutions could be wrong.
- **Objective**: Propose an alternative solution to secure company's business development. Business opportunity, new products launch.
- Who is interested: Pharma, Diagnostics, biotech e Health and cosmetics R&D managers, CEOs.

#### **Exploratory Discovery program to generate novel causal mechanisms concepts.**

- Why: Complex human diseases/disorders need to be revisited to build novel hypotheses.
- **Objective**: Propose novel causal mechanisms concepts for cost-effective novel solutions. **Business opportunity, new products launch**.
- Who is interested: Pharma, Diagnostics, biotech e Health and cosmetics R&D managers, CEOs.



## BMSystems' CADI™ publications to date

#### CADI™ Models published in prestigious peer-reviewed journals: (click on the grey links to get the pdf)

- <u>2014: CNS Psychiatry publication</u>: American Journal of Psychiatry and Neuroscience. Second publications with the Max Planck Institute of Psychiatry in Munich: Differential proteomics analyses reveal anxiety-associated molecular and cellular mechanisms in cingulate cortex synapses. The first output of the DECIUS CNS research program.
- <u>2012, CNS NEURODEGENERATIVE & PSYCHIATRY</u>: PharmacoPsychiatry publishes the first review describing a productive vision of Systems Medicine that will change R&D organization and interactions between clinicians & researchers & reveals how the world's first explanation of the mechanisms of the Creutzfeldt-Jakob disease led to the discovery of a truly innovative psychiatric treatment.
- <u>2011, CNS PSYCHIATRY</u>: Pharmaco Psychiatry publication: Proteome-Based Pathway Modelling of Psychiatric Disorders. Publication with The max Planck Institute of Psychiatry in Munich
- <u>2010, INFECTIOUS DISEASES</u>: Biosecurity and Bioterrorism: Biodefense Strategy, Practice, and Science :Genetically Engineered Virulent Phage Banks in the Detection and Control of Emergent Pathogenic Bacteria. Publication with Pherecydes-Pharma.
- <u>2009, TISSUE DIFFERENTIATION</u>: Médecine & Sciences: Müllerian duct regression explanation. Integrative systems biology & experimental Biology. Publication with CNRS experimental data.
- <u>2005, CANCER</u>: Journal of molecular Endocrinology: Integrative analysis of gene expression patterns predicts specific modulations of defined cell functions by estrogen and Tamoxifen in MCF7 breast cancer cells. Publication in collaboration with INSERM unit 553.
- <u>2003, CANCER</u>: Nucleic Acids Research: Integrated transcriptome analysis of the cellular mechanisms associated with H-ras-dependent malignant transformation of the human breast epithelial MCF7 cell line. Publication in collaboration with INSERM unit 553. World first. First in-silico model of a complex human disease validated in-vitro and published.

#### **Collaboration to scientific reference books:**

- <u>2014: Dermatology Cosmetics.</u> The first reference book on "Computational Biophysics of the Skin" edited by Prof. Bernard Querleux, scientific chairperson of the International Society for Biophysics and Imaging of the Skin
- <u>2011: Phage Nano Technology</u> book published by <u>Valery Petrenko</u>. Chapter 8: Genetically Engineered Virulent Phage Banks for the Detection and Control of Bacterial Biosecurity Threats.
- 2008: CNS: Biomarkers for Psychiatric Disorders. (Ref. ISBN: 978-0-387-79250-7, November 2008). Dr. François Iris, is the author of the Integrative Biology chapter of the book. The editor, Prof. Christoph W. Turck, is head of the Proteomics and Biomarkers branch at the Max Planck Institute for Psychiatry
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## BMSystems' 10 CADI™ programs & POCs

#### Selected POCs and their outputs of CADI™ Programs (all details in <u>full presentation</u>):

- 1. Case study A; Domain: CNS neurology and Psychiatry. Collaborative CADI™ program with CEA life sciences (*1 patent*, *1 publication*, *1 spin-off*).
- 2. Case study B; Domain: Metabolism: First disease application: Parkinson's disease. Collaborative CADI™ program (novel combined therapy proposed for POC in humans).
- 3. Case study C; Domain: Infectious diseases. Collaborative CADI™ program with Pherecydes-Pharma (our first spin-off) (*3 patents*, *1 publication*, *1 spin-off*).
- 4. Case study D; Domain: Industrial biotech. Collaborative CADI™ program with ARD, IBT, CVG, L'Oréal, Rhodia, Arkema (*1 patent filed by an industrial partner*).
- 5. Case study E: Domain: Synthetic biology: Yeast-Based Human-Glycoylation Project CADI v0 produced
- 6. Case study F; Domain: Oncology. Collaborative CADI™ program with Inserm unit 553 (*2 publications, Novel strategy proposed for R&D collaboration*)
- 7. Case study G; Domain: Dermatology. Contractual program CADI™ for a client (8 new targets, cosmetic company confidential).
- 8. Case study H; Domain: Cosmetics. Collaborative CADI™ program) (synergistic low allergy mechanisms identified for safety issues).
- 9. Case study I; Domain: Type 2 diabetes. Contractual CADI™ program for a client (*NO GO decision for safety issue, pharma company, confidential*).
- 10. Case study J; Domain: Tissue differentiation/embryogenesis. Collaborative CADI™ program with CNRS (*1 publication*).

### A new paradigm qualified for industrial use



# BMSystems' Group at a glance

Independent Private Company incorporated in 2004. 100% owned by its founders. Profitable since 2006, thanks to our recurrent clients. We only sell the results of the R&D programs, not our proprietary technologies. 100% biology driven company focused on discovery. Proprietary CADI™ Knowledge Database of mechanisms & interactions. Not domain-dependent, but information-dependent. Markets: Pharma, Cosmetics, Nutrition, Health Technologies, Connected health, Highly productive 24 FTE\* of which 9 FTE on CADI™ Discovery programs only. Strong & long term strategic R&D collaborations (>100 people collaborating). Dual business model: Contractual and/or Collaborative R&D programs. Outstanding internal pipeline of programs ready for collaborations. 2 therapeutic spin-offs\*\* at clinical stage, 4 issued patents, 10 publications. Competition: Key Opinion Leaders, dominant thinking companies or pharma Systems Biology or bioinformatics teams argue they can do the same. We are always open for discussions & comparisons on success rates and outputs for patients.

The World's first Mechanisms-Based Medicine Company You have a R&D issue or a decision to make, we may have a solution for you.



# BMSystems' Management Summary

- □ BMSystems, the world's first Mechanisms-Based Medicine Company, invented CADI™\* Discovery in 2004. **The results:** 
  - 1. Thanks to our recurrent clients, BMSystems, created in 2004, is profitable since 2006;
  - 2. 4 issued patents, including a therapeutics class patent;
  - 3. 2 therapeutic spin-offs\*\* with products now in phase Ib and I/II;
  - 4. a world's first in neurodegenerative diseases;
  - 5. complex biological processes explained (some had remained obscure for over 60 years);
  - 6. 10 peer-reviewed publications and books, ranging from oncology to neuropsychiatry; etc.
- With CADI™ Discovery, BMSystems possesses a truly unique competitive advantage: the ability to address and exploit the two major issues in life sciences
  - 1. the enormous complexity of life's mechanisms, and
  - 2. the significant and growing unreliability of scientific and clinical publications.
- ☐ Through its dual contractual & collaborative business model, BMSystems
  - proposes robust R&D and Business decision making support, and
  - discovers cost-effective, lower risk, patentable novel integrated solutions, for medicine, health technologies, nutrition and cosmetics
- > You have a critical R&D issue or high impact decision to make, we may have a solution for you.



# BMSystems' science at a glance

Correct understanding of disease/disorder/syndrome mechanisms is the first objective. Finding the most pertinent biomarkers and therapeutic solutions is the necessary consequence of the first objective. We check the "CADI™ compliance" of requests in novel domains for GO NO GO decision before launching the full CADI™ Discovery program. We are information-dependent not domain-dependent. CADI™ Discovery was invented in 2002 by Dr. François IRIS, geneticist, physiologist & molecular biologist. Our IT people, from the digital world, developed the platform to "help" our biologists work. CADI™ Discovery addresses the recurrent causes of failures in the "dominant thinking" systems biology programs: the issues of "life mechanisms complexity" and publications unreliability. CADI™ Discovery is operated by biologists to generate and destroy the maximum of working hypotheses before starting the experimental validation phase. Our biologists build heuristic non-mathematical holistic models to generate novel disruptive physiological/ medical meanings from scientific, medical & health smart data. CADI™ Discovery, our proprietary CADI™ Knowledge Database of mechanisms & interactions and our CADI™ domains cross-fertilization process cannot be compared to classical systems biology or bioinformatics. Only discovery processes already delivering novel therapies should be the benchmarks. Highly productive, we successfuly conducted R&D programs in the fields of neurology, inflammation, metabolism, immunology, addressing neurodegenerative diseases (Creutzfeldt-Jakob's, Parkinson's & Alzheimer's diseases), psychiatry, autism, cancer, diabetes, longevity/aging, infections, dermatology, skincare and cosmetics.



# Why do we need to change the dominant discovery paradigm? (supporting documents: click on the links for details)

| The <u>industry is under critical pressure</u> due to a <u>too high failure rate</u> and <u>payers no longer willing</u> to pay premium prices.                                                                                                      |
|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| The Pharma industry has for decades invested in Omics data production, IT technologies and Systems Biology programs for <u>remarkably few relevant results</u> .                                                                                     |
| The consequences of <u>life's mechanisms being complex</u> , as <u>opposed to complicated</u> , are dramatically underestimated by data-treatment scientists and their algorithms.                                                                   |
| "Currently, many published research findings are false or exaggerated, an estimated 85% of research resources are wasted". (John P.A. Ioannidis, MD, DSc PLOS medicine METRICS, Stanford University).                                                |
| The <u>unreliability of scientific</u> and <u>clinical publications</u> used by these algorithms is strongly increasing.                                                                                                                             |
| Negative experimental results are seldom published, generating an enormous bias.                                                                                                                                                                     |
| The "garbage in, garbage out" reality demonstrates that a wrong hypothesis, even if generated or treated by the best Digital and IT technologies, remains a wrong hypothesis                                                                         |
| Mathematical models are remarkable validation/fine-tuning tools when applied to well defined processes. They are inadequate discovery tools when applied to multicellular processes poorly understood and/or created form unreliability information. |

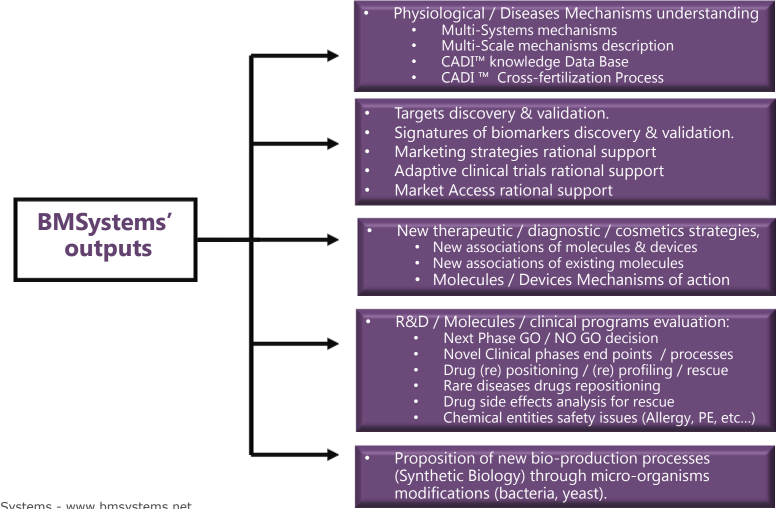
R&D managers aware of these critical & underestimated issues should ask their suppliers to prove that their operational solutions are really able to address these issues.



### BMSystems' detailed answers to clients/partners issues

Reduce time to result, improve success rate and reduce development costs to address specific markets:

biomedical, diagnostic, Pharma, cosmetics, nutrition, food, chemistry, environment, energy.





### **Useful Downloads**

#### Download the Full Presentation with CADI Description, publications and the 10 CADI POCS

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- ☐ Request our Cochin Institute Paris "Integrative Analyses" Training Session Presentation
- The discovery of Innovative Therapeutic Approaches: Under the street light is not the right place to search BIT's 10th Annual Congress International Drug Discovery Science and Technology 2012 November 8-10, 2012, Nanjing, China
- The Differences & Complementarities Between « Heuristic » and « Mathematical» approaches. The scientific presentation given by Dr. François IRIS (CSO BMSystems) during the EPA (European Psychiatric Association) conference in 2011.

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