# Resource

# Protein Function Easily Investigated by Genomics Data Mining Using the ProteINSIDE Online Tool

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### ABSTRACT

Nowadays, genomic and proteomic studies produce vast amounts of data. To get the biological meaning of these data and to generate testable new hypothesis, scientists must use several tools often not designed for ruminant studies. Here we present ProteINSIDE: an online tool to analyse lists of protein or gene identifiers from well-annotated species (human, rat, and mouse) and ruminants (cow, sheep, and goat). The aims of ProteINSIDE modules are to gather biological information stores in well-updated public databases, to proceed to annotations according to the Gene Ontology consortium, to predict potentially secreted proteins, and to search for proteins interactions. ProteINSIDE provides results from several software and databases in a single query. From a list of identifiers, ProteINSIDE uses orthologs or homologs to extend analyses and biological information retrieval. As a tutorial, we presented how to launch, to recover, to view, and to interpret the results provided by the two types of analysis available with ProteINSIDE (basic and custom analyses). ProteINSIDE is freely available using an internet browser at www.proteinside.org. The results of this article are provided on the home page of ProteINSIDE website as the example of an analysis result.

### **AVAILABILITY AND REQUIREMENTS**

- Project name: ProteINSIDE
- Project home page: <u>http://www.proteinside.org</u>
- Operating systems: Linux, MacOS, Windows
- License: No

### **KEYWORDS**

Online tool, workflow, protein function, protein interaction, protein secretion, gene ontology, networks, ruminant, genomics.

### INTRODUCTION

Given the increasing amount of genomic and proteomic data produced even in ruminants [1, 2, 3], there is a challenge for the bioinformatic data processing, which has not yet been completely solved [4]. Such bioinformatic data processing has to proceed to data gathering and database searching in order to produce a functional interpretation of large datasets. For this purpose, workflows integrating several bioinformatics analyses are now available [5-8] and were developed to mine dataset from specific species (BioMyn [9] for human, DroPNet [7] for Drosophila, TAIR [10] for Arabidopsis thaliana, EcoCyc [11] for Escherichia coli ...) or to identify candidate genes related to diseases as ToppGene [12] or NetPath [13]. The few workflows currently used for the bioinformatics data processing of ruminant datasets are multispecies. Consequently, the data source of the results proposed is not available because of the privacy of databases (as the licensed software Pathway Studio [14] or Ingenuity Pathway Analysis (<u>www.ingenuity.com</u>; Redwood City, CA, USA). An alternative for scientists working with ruminant datasets is to use dedicated and complementary bioinformatics tools implemented as web services. These tools are dedicated to one type of analysis, as for example the annotation according to the Gene Ontology (GO) [15], the prediction of signal peptide to identify putative secreted proteins [16], or the molecular interactions identification [17] and visualization as networks [18, 19]. Whatever the analysis carried out, the first step is to connect a protein name to a unique identifier (ID). Conversely to gene names that have been standardized, protein names or IDs can differ between databases or tools, especially for ruminant data that remains to be largely curated in most of databases. Thus, the use of several bioinformatics tools to mine ruminant datasets leads to a substantial loss of information and time.

A strategy to perform a systematic and integrative analysis of biological protein information from ruminant datasets is to develop an online workflow that integrates several analysis steps in one package and from a unique ID. Thus, we propose ProteINSIDE [20], a web service dedicated to a systematic and integrative analysis of protein's biological information from ruminant datasets. Unlike human, mouse or rat, ruminant species are less annotated and protein sequences or information are not always curated. Often, scientists working with ruminant use orthologs or homologs with the aim to increase the meaningful

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biological contexts for proteins thanks to knowledge available in well-annotated species. Thus, ProteINSIDE was designed to run using lists of protein or gene IDs from 6 species (cow, sheep, goat, human, rat, and mouse) to annotate biological and molecular functions and cellular location, predict secreted proteins, search for interactions between proteins within and/or outside a dataset. The objective of this article is to propose a tutorial to use ProteINSIDE and interpret generated results.

# **METHODS**

This section lists necessary equipment, ProteINSIDE resources and describes the dataset used to assess the functionalities of the tool.

### **ProteINSIDE's features**

ProteINSIDE is an online workflow with an interface devoted to accessible and fully customisable analyses from lists of protein or gene IDs. Registered users have access to an analyses manager to run and save analysis, and visualise the results. Unregistered users can use ProteINSIDE, but there is no analyses manager and analyses are deleted each month. ProteINSIDE is divided into three parts: the workflow, the database, and the web interface (Figure 1).

The web interface, designed to easily use ProteINSIDE, helps the user to create the analyses, to have access to the results thanks to a balance between technical functionalities and visual elements, and to inform about updates (Figure 1). ProteINSIDE proposed two types of analysis to be launched: the basic analysis (automatic settings) and the custom analysis (user's settings). There is also a pre-set analysis for registered users only who want to make a new analysis with settings of a previous analysis.

The basic analysis performs a:

- Functional annotation using GO terms by querying QuickGO database [21] without electronic annotation.
- Prediction of secreted proteins using SignalP [16] and TargetP [22] software. We improve the prediction by giving GO terms related to the

cellular location of the protein and the processes of secretion.

• Search of proteins interactions curated and listed in IntAct [23], UniProt [24], and BioGrid [25] databases.

The custom analysis performs programs and their options that have to be selected by the user in order to:

- Perform a functional annotation using GO terms from QuickGO, with the options to select also electronic annotations (predicted and scripted annotations), and to generate a GOTree view of linked GO terms (pathways of functional annotation).
- Predict secreted proteins with the option to increase the software's sensitivity of prediction and by this way to increase the number of predicted proteins, however with a higher number of false positive results.
- Search for protein interaction within (core network) and outside (extended network) the uploaded dataset. Options propose to select interactions stored within 1 to 31 databases gathered by the PSICQUIC website [17]. User can select the databases depending on the type of interactions (PPi, Nucleic acid-Protein interaction (NPi), and Smallmolecule-Protein interaction (SPi)) and the data (curated, predicted, curated according to the IMEx project [26] or the MIMIx curation [27]). PSICQUIC service or some databases could be offline, that's why the status of each website is indicated in the table.

To submit an analysis, users either directly paste a list of IDs or upload a file of IDs. Inputs can be protein (e.g., ADIPO\_HUMAN) or gene (e.g., ADIPO or gi[62022275) ID, or protein accession numbers (e.g., Q15848) from six species: cow, human, rat, mouse, sheep, and goat (Figure 2). A new analysis is run directly or is placed on a waiting list if the workflow is overloaded. Uploaded data and results remain confidential.



Figure 1. Flow chart of ProteINSIDE structure. The four workflow's modules are either all launched in the basic analysis or individually selected in the custom analysis. These modules aims to query the available biological information, annotate according to the gene ontology, predict signal peptide and visualized protein-protein interactions.

In addition to the web interface, ProteINSIDE is composed of a database and a workflow.

The database (invisible to any user) collects and stores the information required for the proper functioning of ProteINSIDE. It stores analysis settings and results to reduce server load (Figure 1). The database stores also a gathering of biological information from the NCBI [28] (Gene, Protein, and HomoloGene for known orthologous proteins between the 6 species) and UniProt [24] databases (for the ID Mapping module), and QuickGO [21] and AmiGO [29] (for the GO annotation module). A script updates automatically and monthly the database by extracting IDs, homologs, biological function, FASTA sequence, and other information from the latest releases of these databases.

The workflow uses uploaded data. It is a combination of Perl and R scripts to query databases, recover protein data, perform calculations and run algorithms for signal peptide predictions and network visualisation (Figure 1). The workflow is invisible to any user. The workflow is composed of 4 parts: the "ID Mapping", the search of annotations according to GO, the prediction of secreted proteins, and the search of protein-protein interactions (PPi). The workflow always starts by the ID Mapping program which searches the biological information available for each protein or gene of the input within the ProteINSIDE database. Gathered biological information is required to run the 3 other modules of the workflow: "Gene Ontology", "Secreted Proteins", and "Protein Interaction" (described in the "Results" section of this article). The GO program gueries QuickGO and ProteINSIDE's databases to perform the functional annotation. The GO program analyses over- and under-represented terms to highlight the most relevant GO terms related to the input. These statistical calculations are made with an R script performing a Fisher's exact test (functional enrichment first proposed by FatiGO [30]) and the resulting *p*-value is corrected or not by the Benjamini & Hochberg (BH) test [31]. The prediction of secreted proteins is made using a local version of SignalP (version 4.1) that looks for a signal peptide on amino acid sequence of each protein [16] (cutoff of 0.45 and 0.34 for SignalP prediction, in the basic and custom analysis with the sensitive option selected, respectively; for more information see the tutorials of SignalP<sup>a</sup>. To ascertain that proteins are secreted, ProteINSIDE uses TargetP [22] (version 1.1) to predict the cellular location of each protein. ProteINSIDE uses a pre-set cutoff option to get a significant prediction (higher than 95%) according to TargetP instructions<sup>b</sup>. Protein interactions are searched using PSICQUIC service [17] and statistical calculations are made with an R script and the package "tnet" [32]. ProteINSIDE performs sequence alignment thanks to a local version of NCBI BlastP [33] against UniProt/Swissprot databases [24, 34]. Lastly and as an additional valuable tool, ProteINSIDE lists in one table all known IDs for an input of proteins or genes thanks to the ProteCONVERT module. This list is the result of a search and of a gathering of IDs thank to the ProteINSIDE biological database. Only registered users have access to the ProteCONVERT module.

elect your species:	of your IDs
) Input IDs; se a sample file: SAMPLE xample of IDs that can be used; Q15848 INPUT C ile format (250 ko max / 4000 IDs max). C · Microsoft Excel file (.xls or .xlsx) · Tablulation file (.tab or .txt)	or ADIPO_HUMAN or ADIPOO or gil62022275 (or login and use ProteCONVERT to get right IDs) PASTE PASTE your IDs (8000 IDs max):
put file tutorial : How to create my input f	file? Which file can I use?
ettings for this basic analysis : Gene On owered with <u>Psicquic and SignalP 4.1</u> )	tology / Signal Peptide prediction / Interactions research within the input dataset (IntAct / UniProtKB / BioGrid

Figure 2. Setting up a basic analysis. First, enter a name for the analysis and select the species of study. There are two ways to submit a protein or gene list; you can use an input file or directly paste your IDs. The input file must be less than 250 kb and the file format must be specified. There is also a "Sample" button that loads parameters for an example analysis. Once everything is filled, click on the button "Run the job" to submit.

The results of an analysis with ProteINSIDE are available online or downloadable. Unregistered user gets results through a unique code and a link provided after the submission of a list. Registered users have access to the results through their analysis manager after their connection with their login and their password. Four separated pages provide results from the four analyses ("ID Mapping", "Gene Ontology", "Secreted Proteins", and "Protein Interaction"). The results are dynamic tables and charts that can be sorted and filtered by specific criterion such as biological function, protein or gene ID, or p-value. Tables and charts are made using Google Charts package (https://developers.google.com/chart/) and are downloadable, diagrams and histograms are printable. Networks are downloadable as image (.pdf or .png) or as network viewer input files (Cytoscape .cys or graphml and xgmml). The results interpretations are detailed in the results section.

# Implementation

ProteINSIDE is freely available online at <u>www.proteinside.org</u> and doesn't require an installation on a computer. ProteINSIDE is completely adapted for any internet browser and tablet. We recommend multiprocessors computer with at least 2 GB of ram to get better performances for huge network visualization and filtering.

The web interface is programmed in PHP, HTML, and JavaScript. The workflow has been completely programmed in Perl (version 5.10.1; CPAN modules (Comprehensive Perl Archive Network) and BioPerl [35]) and R scripts (version 3.0.1). The database was made in MySQL (version 5.5) (Figure 1).

# Sample dataset

We have created a dataset to assess ProteINSIDE performances. This dataset is composed of the UniProt accession numbers of 133 proteins (Table 1): 34 proteins related to the glycolysis cycle, 11 proteins from the respiratory chain, 5 proteins from the tricarboxylic acid cycle (TCA), 79 hormones or secreted proteins, and proteins with very specific functions unrelated to the others. We also included a duplicated ID among proteins of the glycolysis to verify its recognition by ProteINSIDE. ProteINSIDE is able to detect duplicate protein even if the IDs are different: a Gene Name, a UniProt accession number, and a Gene Identifier related to a same protein will be taken into account as a single protein. We have created this dataset on bovine species, but the number of annotations and PPi weren't sufficient for a complete overview of the functionalities of ProteINSIDE. Then, we used the same proteins using human IDs to test ProteINSIDE with the basic and the custom analyses (Table 1).

# RESULTS

Here we present how to run a basic or custom analysis and how to view the results. We explain how to interpret the results and we discuss the relevance of biological information extracted by ProteINSIDE for our sample dataset of 133 proteins.

# Setting up a Basic Analysis: a standard overview of a dataset

ProteINSIDE performs a basic analysis (in which settings are locked and the workflow provides GO terms, list of putative secreted proteins, and PPi data from IntAct [36], UniProt [24], and BioGrid [37] databases). A basic analysis gives a complete overview of a dataset. To set up a basic analysis, user has to follow these steps (Figure 2):

- Click on "Basic Analysis" menu on the homepage
   of ProteINSIDE
- Fill in "the job name" box
- Select the species of study (the same species as the uploaded IDs)
- Upload an input file or directly paste IDs
- Click on the "Run the job" button to submit a new analysis

The analysis status is indicated by the colour of a button: red for "analysis on the waiting list", yellow for "the analysis is running" and green "analysis done". The blue globe is the link to access to the online results:

- Click on the blue globe button to view the results (or use the trash to delete them)
- Visualise the results summary produced by the four modules of analysis on the first default page (entitled "Results Summary", Figure 3)
- Navigate between the four module's results pages by clicking on the module's name on the toolbar menu.

Analyses and data	Glycolysis	Hormones	TCA	Analysis duration (min)
Dataset	33+1 duplicated	79	5	-
Basic analysis	27	78	3	2
Custom analysis	33	79	5	10

Table 1. Results summary of ProteINSIDE analysis performances. The numbers are the proteins that belong to main pathways in the sample dataset, that are properly annotated by GO terms relevant to glycolysis and tricarboxylic acid (TCA) pathways, and that have been predicted as secreted by SignalP (and confirmed by GO terms, TargetP, and subcellular location) for hormones.

For our sample dataset, the "Results Summary" page reported that all 133 proteins were recognized by ProteINSIDE and the protein in duplicate was identified and excluded from the analysis (Figure 3). Thus, 132 proteins were submitted to the four modules of analysis.

The "ID Mapping's" module aimed to retrieve and gather basic biological knowledge, results are directly viewed on the "ID Resume" web page of ProteINSIDE. This module compares each submitted IDs to the database of ProteINSIDE to ascertain a match with genes or proteins from human, rat, mouse, cow, sheep or goat species. The local biological database of ProteINSIDE is a combination of NCBI Gene/Protein, NCBI HomoloGene [28], and UniProt [24] databases. These databases were chosen because data are easily extractable, curated and daily updated. For each uploaded ID, ProteINSIDE obtains and summarises as a downloadable table (Figure 4): gene or protein ID, gene or protein name, a summary of protein function, gene chromosomal location, and information on tissue expression and cellular location. The module also recovers the protein sequence of each input ID. Each protein and gene ID listed on this web page are linked to corresponding UniProt and NCBI web pages. FASTA amino acid sequences of each input are also downloadable.

The module dedicated to the functional annotation according to the GO consortium, produces results that are viewed on the "GO" web page of ProteINSIDE. ProteINSIDE imports GO terms by querying the QuickGO database [38]. QuickGO was chosen because of its daily update, accessibility, and performances. ProteINSIDE only imports GO terms that have been selected by evidence codes (GO Inferred from Electronic Annotation codes (IEA) are excluded by the basic analysis) and confirmed by curators. The GO script of ProteINSIDE analyses over- and under-represented terms to identify the most relevant and the most specific terms associated with the uploaded list. Within a GO, ProteINSIDE compares the number of genes or proteins from the dataset to the total number of gene products (for a species) declared in the AmiGO database [29] to provide a coverage frequency, and thus, to identify the most representative pathways associated to a dataset. The result is viewed on the "GO" web page of ProteINSIDE as tables and diagrams. Three tables (Figure 5) report the GO terms that annotated two or more proteins (Figure 5-B), the GO terms that annotate one protein (Figure 6-C), as well as all GO terms for a protein (Figure 5-D). Each annotation is informed with an evidence code (that reflects the type of experimental evidence or analysis to describe an annotation between a GO term and a gene product) and the database source. Tables are automatically sorted by the best enrichment p-value to help the user to view the most significant GO terms related to a dataset. Tables can also be sorted by ontology group, p-value range for enrichment, GO term description, gene name or any input IDs (Figure 5B). From the sample dataset of 132 proteins, ProteINSIDE annotated 128 proteins with 624 GO terms. The most significant enriched GO terms is "hormone activity" (that annotated 31 proteins over the 79 expected; not shown) and "glycolytic process" (that annotated 27 proteins over the 33 expected; Table 1). The low number of annotated proteins may be related to our choice to use only GO terms that have been confirmed by curator in the basic analysis. This means that the basic analysis doesn't use the annotation with IEA (Inferred by Electronic Annotation) evidence code. However, the option to use IEA is provided in the custom analysis to extend the annotations.

	General information	Analyze general results				
Name: 133prot_GCB Type: Basic Species: HUMAN Date: 2015-01-14 Code access: ydcn07a7kxjxh0 Download all results:	Jsqc9p6e0tbs8enta	Query find in ID Mapping DB: 132     All GO detected: 624 for 128 annoted protein(s)     Signal peptides detected: 85     Interactions detected on the dataset: 29				
(lot of results = more treatmen Analysis parameters: ID Mapping / Gene Ontology /	t time) SignalP / Interaction research (IntAct / UniProtKB / BioGrid)	Number of proteins involved for each module				
Incomplete information: Incomplete query: 12 + show/hide + Duplicate query: 1 + show/hide +	Incomplete information: - Query misinformed by the ID M - Duplicate query on the dataset - ID Not-found on the dataset	lapping module				
Not imported query: 0 + show/hide +	(click on "show/hide" to view the I	Ds)				

Figure 3. Main page of results produced by a basic analysis. This is the first page of the results. It shows the number of proteins or genes successfully analysed by each module.

The module that aims to predict potentially secreted proteins provides results on the "Secreted Proteins" web page of ProteINSIDE (Figure 6). To identify proteins that are putatively secreted, ProteINSIDE first predicts the presence of a signal peptide on a protein sequence (imported by the "ID Mapping" module) through a local version of the SignalP tool [16]. SignalP was chosen because of its high prediction score in comparison with other available tools [39, 40]. To ascertain the prediction, a local version of TargetP software [22] predicts the subcellular location of the proteins. ProteINSIDE also checks the subcellular location of the protein using UniProt source to confirm TargetP results. As a final verification step, ProteINSIDE selects the GO terms related to secretory pathways for each SignalP prediction. For this purpose, we have selected about 1,000 GO terms related to secretion (monthly updated) as for example: secretion, vesicle, or extracellular region. This four-step analysis improves the reliability of proteins proposed to be secreted thanks to a signal peptide and to our knowledge is unique to ProteINSIDE [40]. However, proteins are also secreted by pathways that do not involve signal peptide such as: endosomal recycling, plasma membrane transporter, membrane flip-flop, and membrane blebbing including the formation of vesicles or exosomes [41]. Thus, ProteINSIDE was designed to predict the proteins secreted by other pathways, by gathering the data of subcellular location provided by UniProt, GO terms, and TargetP results (Figure 6-B). From our sample dataset of 132 proteins, ProteINSIDE has predicted 85 proteins as potentially secreted outside the cell by a signal peptide, among them 78 over the 79 proteins that were expected (Table 1). This lack of perfect prediction can be explained by the false positive and false negative prediction rates of SignalP, as already evaluated by Petersen et al. (Supplementary materials and methods of [16]). Over the 85 predicted secreted proteins, 65

were also annotated by GO terms related to the secretion. The subcellular locations of 81 proteins were both confirmed by TargetP and UniProt source. Additionally, 30 proteins were predicted to be secreted without signal peptide.

The fourth module is dedicated to PPi analysis and results are viewed on "Protein Interaction" web page of ProteINSIDE. PPi identification and visualisation within a network conveyed how various genes or proteins contribute to cellular or metabolic processes. ProteINSIDE uses the PSICQUIC service [17] to identify PPi and imports PPi identified by their "interaction detection methods" with experimental proofs and confirmed by curator. The basic analysis identifies PPi within the uploaded dataset (core network) using the preselected databases IntAct, UniProt, and BioGrid. These PPi databases were chosen as a default option because there are daily updated and reviewed by curators as well as by the curation processes of the IMEx project (that ensures reliable interactions data using experts and curation rules shared between many interaction databases [26]) or MIMIx (a guideline of the minimum information required for reporting a molecular interaction experiment, thus advising the user on how to use the interaction data [27]). Moreover, BioGrid is the biggest PPi database that has its own curation workflow (more than 740000 curated PPi) and is not a partner of IMex curation program. IntAct is another big PPi database with more than 380000 PPi currated according to IMex and MIMIx curation rules and that are often listed in several databases. UniProt is a major database dedicated to the study of proteins. Thus, it possesses its own curated PPi but in lesser amounts compared to the two other specialized databases (less than 13 000 PPI; UniProt is a partner of IMEx project). By using 3 databases as a default option, the aim of ProteINSIDE is to favour the use of multiple PPi databases in order improve the PPi to data gathering [42].

R	esea	rch are	а		Dov			
Gene Na	me	e Proteins Functi		Func	tion Tissue Subcellular location Number of rows townload t	table 🗶 / Sequences (FASTA) 🌿		
Query	Proteins	Protein ID	Gene Name	Gene Entrez	Function	Chromosite	Tissue	Subcellular location
1	<u>Q15848</u>	ADIPO_HUMAN	ADIPOQ	<u>9370</u>	Important adjockine involved in the control of fat metabolism and insulin sensitivity, with direct and-fability, and-informationy carities. Stimulate: AMPK phosphoryhoutian and caritalion in the liver and the solvential musi- glucose utilization and fatty-acid combustion. Antaporizer: NFF-alpha by negatively regulating its expression in such as liver and macrophages, and allow by counteracting its effects. Inhibits endoffeital NF-alphage Signaling dependent pathway. May play a role in cell growth, angiogenesis and tissue remodeling by binding and sequest growth factors with district binding antilinities, depending on the type of complex. LUW, MLMV er MW.	nti-atherogenic and cle, enhancing n various tissues g through a cAMP- tering various	Synthesized exclusively by adipocytes and secreted into plasma.	Secreted.
2	<u>Q7Z4H4</u>	ADM2_HUMAN	ADM2	<u>79924</u>	IMDL and IMDS may play a role as physiological regulators of gastrointestinal, cardiovascular bioactivities mee CALCRU/RAMPs receptor complexes. Activates the cAMP-dependent pathway.	diated by the 22q13.33 NC 000022.11 (5048155650486437)	Expressed in the esophagus, stomach, jejunum, ileum, ileocacum, ascending colon, transverse colon, descending colon and rectum. Expressed in myocardial cells of the heart, renal tubular cells, hypothalamus, and pitulary.	Secreted.
3	<u>P35318</u>	ADML_HUMAN	ADM	<u>133</u>	AM and PAMP are potent hypotensive and vasodilatar agents. Numerous actions have been reported most re physiologic control of fluid and electrolyte homeostasis. In the kilowy, am is diuretic and natriuretic, and both an aldosterone secretion by direct adrenal actions. In physikary gland, both peptides at physiologically relevant door ACTH secretion. Both peptides appear to act in brain and pituitary gland to the patient by the loss of plasma volume complement their hypotensive effects in blood vessels.	elated to the am and pamp inhibit es inhibit basal e, actions which	Highest levels found in pheochromocytoma and adrenal medulla. Also found in lung, ventricle and kidney tissues.	Secreted.

Dynamic table: results are sorted and filtered by clicking on columns; a specific search is made possible by a search depending on gene name, protein accession number or protein function.

Figure 4. Biological knowledge retrieval. The ID Mapping module results are listed in a table. This table provides protein IDs, gene names, summaries the protein function, chromosomal locations, data on tissue expression, and subcellular location. User can sort the table by using the dynamic table research area.

These 3 PPi databases ensure the good recovery of known interactions for an overview of interactions within or/and outside of a new dataset. Then, ProteINSIDE lists pairs of proteins known to interact between each other in a downloadable table (Figure 7) and constructs a network (Figure 8) using the PPi identified within the uploaded list. The dynamic network is available by using the "Cytoscape" button on the "Protein Interaction" web page ("Dynamic Cytoscape view of PPi", Figure 7-A). Within the network, edges are experimental detection methods used to identify the PPi. Consequently, several edges may link two proteins. Network can be sorted by the number of interactions by node, the proximity of a node to other nodes (closeness centrality; CC) and the shortest paths between nodes (betweenness centrality; BC) (Figure 8-A). These centralities criteria were already proven to be efficient to select key nodes/proteins within a pathway [43]. From our sample dataset of 132 proteins, ProteINSIDE has identified 29 PPi that involved 28 different proteins (Figure 7-B). As expected from our small dataset, ProteINSIDE linked, within sub networks, proteins involved in glycolysis, TCA or respiratory chain as protein complexes (partially on Figure 8-B).

# Setting up a Custom Analysis: an addedvalue provided by the extension of the analysis

We made a custom analysis using the same major settings as for the basic analysis with additional options (GO network, GO electronic annotations, and extension of PPi to proteins outside of the dataset in the same species, extended network). To set up the custom analysis, user has to follow these steps (also explained by Figure 2):

- Click on the "Custom Analysis" menu on the homepage of ProteINSIDE
- Fill in "the job name" box
- Select the species of study (the same species as the uploaded IDs)
- Upload an input file or directly paste IDs

Then, user has to select the settings of either all or only one module of analysis on the section "4" of the page, by following these steps (Figure 9):

All GO det     Dynamic p     General Anno     Cellular Comp     Molecular Fur     Biological Pro	ected: 624 for 128 an ie chart: tation conent notion ccess	oted protein(s) in HUMAN logg groups 0 0 0 0 0 0 0 0 0 0 0 0 0	Download complet GO table: Download GO by protein table P-value range 1 > 10e-3 f0e-3 > 10e-5 f0e-5 >	106.5 106.9 > 0	Download results								
P-value rar	notations GOs	Proteins	Gene Name		Ontology Groups	# GO per p	age "Comple	ete term annotation" fro	om				
Choose a	value •				Choose a value	10 🔻		HUMAN		<b>B</b> )			
GO	Function	Prote	ins		Gene Name		Ontology Groups	GO frequency within the dataset (%)	GO frequency within the genome (%)	P- P-v value 🔺 (Bł	value H)		
<u>GO:0005179</u>	hormone activity	A6NKQ9 P01233 P01222 P81 O6NT52 P61278 Q9Y581 O43 P01242 P09681 P16860 P0133 P01308 O95399 P09529 P012 P06860 Q15848 P01189 P048	277         P01148         P06307         P10082           555         P01275         P09683         P01215           50         P01270         P01298         P01282           25         Q14406         P08476         P12272           08         P05111         P0         P0	CGB1 CGB8 T CGB2 SST INS GH2 GIP NPPE UTS2 INHBB F CRH ADIPOQ	CGB1 CGB8 TSHB PRLH GNRH1 CCK PYY CGB2 SST INSL6 GNRH2 GCG SCT CGA GH2 GIN PRPS GAST PTH PPY VIP INS UTS2 INHBB FSHB CSHL1 INHBA PTHLH CRH ADIPOQ POMC RLN1 INHA					9.847e-74 3.02	!3e-71		
<u>GO:0006096</u>	glycolytic process	P00338         P08237         P52789         P193           P06733         P09972         P14618         P0919           P05062         P00558         P15259         P0444           P60174         O14556         O60825         P139	67         P36871         P30613         Q01813           04         P17858         Q1683         6744           06         P07205         P0407         Evide           29         Q16877         P16118         Evide	LDHA PFKM H ENO1 ALDOC	K2 HK1 PGM1 PKLF PKM ENO2 PFKL PF A / Database sour	60.000 27 /45	8.638e-62 1.76	8e-59					
<u>GO:0005576</u>	extracellular region	P01588 P20396 Q15848 Q9U8 P01229 P09681 P01308 P203	<u>3U3 P06307 Q9UEF7 P01225</u> 82 P0DML2 P06850 P01215	EPO TRH ADII	POQ GHRL CCK KL PMCH CSH1 CRH C	FSHB GA VIP	CC	49.219 63 /128	63 /3705	2.138e-50 3.28	82e-48		
60	oluceso 6 phosphate	Function	Proteins	Gene Name	Ontology Groups		GO frequenc	y within the genome (%)	P-value +	P-value (BH)	2020.2		
<u>GO.0004347</u>	giucose-o-priospriate	somerase activity	<u>P06/44</u>	GPI	MP		1 /1		1.0056-5	0.	2028-5		
GO:0004382	guanosine-diphospha	ase activity	<u>075356</u>	ENTPD5	MF		1 /1		1.869e-3	6.	202e-3		
<u>GO:0004742</u>	dihy drolipoy lly sine-re	idue acetyltransferase activity	P10515	DLAT	MF	_	1 /1		1.869e-3	6.3	202e-3		
<u>GO:0004807</u>	triose-phosphate isor	erase activity	P60174	TPI1	MF		100.000 1 /1		1.869e-3	6.	202e-3		
Proteins         O           P08476         IN           P01270         PT	Sene Ime HBA GO:0000082 G GO:0045578 G H GO:0045578 G GO:004558 G	C.0001541 GC.0001942 GC.0002244 GC C.0017046 GC.003154 GC.0030218 GC 0.0045548 GC.004556 GC.0045786 GC 0.0045125 GC.003705 GC.0045786 GC 0.0045225 GC.0051428	2 0005125 GC 0005160 GC 0005179 G 0 0033308 GC 0032270 GC 0032924 G 0 0045893 GC 0045944 GC 0046880 G 0 0005576 GC 0005615 GC 0006366 G	C 0005515 GO 00055 C 0035987 GO 00423 C 0046881 GO 00468 C 0007186 GO 00072	GO 76 GC 0006357 GC 0006 26 GC 0042476 GC 0042 26 GC 0048468 GC 0050 167 GC 0008628 GC 0010	852 GO:00070 493 GO:00425 021 GO:00602 468 GO:00305	50 GC-0007166 41 GC-0042701 79 GC-0060395 01 GC-0030819	GC 0007267 GC 0007399 G GC 0042802 GC 0043408 Gr GC 0061029 GC 0070699 G GC 0031855 GC 0034645 Gr	0.0008083 GO.0008285 G 0.0043509 GO.0043512 G 0.0097191 GO.2001241 0.0045453 GO.0045725 G	0.0008584 0.0045077	53 21		
P00133	GO:0000122 G	0:0045892 GO:0070062	0000010 00.0000615 00.0005634 0	GO0005757 GO00058	ka GO.0005375 GO.0006	00 GO:00060	94 00.0006096	GO.0009402 GO.0009615 G	0.0010020 GO:0030308 G		21		

**Figure 5. Functional annotation according to the Gene Ontology.** GO results are first extracted and classified by the number of GO terms related to Molecular Functions, Biological Processes, and Cellular Components, then visualised as diagrams or downloadable tables. (A) Main menu of GO results page, to download the results as Excel files, to view the significance of *p*-value range colours, or a proportion of major annotation categories as diagram. (B and C) GO terms are also sorted as two dynamic tables (a table for GO terms that annotate more than one protein on the dataset - B, and a second table for the GO terms with a single protein annotated - C). Tables can be sorted by GO term, function, protein name or ID, gene name, number of annotations, annotation frequency or annotation enrichment. (D) A third table lists all GO codes for a given protein. Users can move the cursor over a protein to be informed about the evidence code and the database source of the GO annotation (B; where IDA means "Inferred from Direct Assay").

- Select Gene Ontology module
  - Select "GO electronic annotation (IEA)" if you aim to use GO annotation inferred from electronic annotation.
  - Select "Gene Ontology Tree network" to view linked GO terms
- Select Signal peptide module to use the basic cutoff value of prediction
- Select Protein protein interaction module
  - Select "Protein protein interaction custom analysis"
  - Select "Extend PPi research using protein outside of the dataset", if wanted
  - Select "Human species" to analyse PPi using data available in Human, for example
  - Select either the 3 most used databases (IntAct, UniProt, and BioGrid as used in the basic analysis) or from 1 to 31 databases (PPi are daily updated in each database)

Alternatively, user can load automatically the same settings as those already used in a previous custom analysis, by clicking on the "pre-set" button.

At the completion of the custom analysis, the "ID Resume" web page provides the same information than the basic analysis (Figure 3).

Within the GO module, the choice to use the electronic annotation option has increased both the number of annotated proteins (132 rather than 128 without IEA in the basic analysis) and the number of annotations by around 40% (1080 unique GO terms rather than 624 in the basic analysis). Thanks to IEA option, ProteINSIDE correctly retrieved the 33 expected proteins related the glycolytic process and the 79 proteins related to a hormone activity (Table 1). The GOTree network linked 570 GO terms. A link between 2 terms is represented by an "is a" relation: "Diuretic hormone activity" linked to "Hormone activity" means that "Diuretic hormone activity" is a "Hormone activity" pathway. The network can be sorted by ontology group, by p-value range (to select and to link only the most enriched GO terms), by the number of directly linked terms or also by the number of GO terms linked together (to select group of GO terms involved in the same biological function). From our sample dataset, we have chosen to illustrate the GO tree of the "Molecular Function" group (Figure 10). In this visualisation, squares with dark red colour were GO terms which have annotated the highest number of proteins. Among them and as expected the GO:0005179 with the best *p*-value and the darkest red colour was "Hormone activity", in agreement with the over representation of hormones in our sample dataset.

The "Secreted Proteins" module has predicted the same 85 proteins as the basic analysis as being secreted. By comparison with the basic analysis, the use of IEA option has allowed to confirm this prediction for 82 proteins that were also annotated with GO terms related to a "secretion" function.

By comparison with the basic analysis, the settings selected within the "Protein Interaction" module provided PPi within the dataset (between proteins of the dataset, as the basic analysis) and PPi between proteins from the dataset and outside of the dataset. For the extended network, ProteINSIDE retrieved 688 PPi made by 500 proteins. Among them, 61 proteins were from our uploaded sample dataset. By using PPi outside of the dataset in Human species, we got 95% more PPi that involved 60% more proteins from the sample dataset than the PPi recovered with the basic analysis. The extended network (Figure 11) highlighted major subnetworks related to the respiratory chain (Figure 11-A), hormone activity such as signalization pathways of adipokines (Figure 11-B), growth hormone (Figure 11-C), thyroid hormones (Figure 11-D), glycolysis (not highlighted), and carbohydrate metabolism (not highlighted). This is consistent with the over selection of proteins from glycolysis, TCA or hormones or adipokines. Betweenness and closeness centralities were used to sort the most central proteins of this extended network (Figure 11-E). By this way, we identified 22 highly central proteins, 13 of them coming from the uploaded sample dataset and involved in respiratory chain and glycolysis as protein complexes.

# DISCUSSION

Currently, most genomic and proteomic studies increasingly generate data which have to be gathered, filtered, and analysed using one or more softwares [44-46]. The major and widely used strategies to systematically study proteins [47] and genes [48] in a cell are based on functional annotation, proteins interactions and pathways analysis. The literature describes many tools for genomic and proteomic data analysis [4]. Scientists have to select appropriate tools among those for either the GO annotation [15, 30, 49, 21, 29, 50], the prediction of secreted proteins [51, 52, 39, 53], or the search of protein - protein interactions [54, 55, 36, 56, 37, 57, 58].

ProteINSIDE is not just an additional resource since it was designed to provide efficient and original strategies to run in a single query, biological knowledge gathering, GO terms annotation, secreted protein prediction, and protein interaction. The DAVID [59], ToppGene [12] or Babelomics [60] software resources are often mentioned for the biological knowledge gathering, functional annotation using GO terms or searches for proteins interactions. By comparison to these tools, added-values of ProteINSIDE have to be highlighted.

ProteINSIDE provides a functional annotation using a monthly updated GO terms database and enrichment calculation. Indeed, the list of GO terms is in constant evolution and GO terms could become redundant or obsolete the next month [15]. This could induce bad information in the results of an analysis if the database is not often updated. Each result of the annotation is easily readable thanks to dynamic tables and diagrams which can be sorted with many options and can be downloaded to work offline. The GO tree visualization of the most often associated GO terms with a list of IDs, is another added-value of ProteINSIDE. Tree networks of GO terms are also done by AmiGO or QuickGO to get an ancestor chart of a single term. However, ProteINSIDE is the only tool which highlights biological pathways of a dataset using linked GO terms and their representativeness rate (using *p*-values and number of annotations). This network visualization is also easy to use thanks to the friendly user interface that gives access to the sort options. For the PPi research and visualization, ProteINSIDE uses only interactions that are based on experimental observations. The drawback is that the number of PPi identified by ProteINSIDE could be lower than those proposed by other resources that also list predicted interactions inferred from literature mining. Furthermore, ProteINSIDE is also capable to draw large interaction networks thanks to the use of the powerful graphical Cytoscape application. ProteINSIDE provides different options to filter large networks, making it as easy to use as the widely used resource STRING [57], and efficient to select keys proteins in a network. Moreover, to analyse locally the networks, files (e.g. .cys, xgmml, graphml) are ready to be open by a network viewer like Cytoscape (and its numerous plugins) and are downloadable from the PPi page result. To our knowledge, among the tools to mine genomics data from mammals, ProteINSIDE is the only resource that allows a very simple view and analysis of network, and prepares data for their further download and analysis by other network viewer software as Cytoscape. These features may be valuable for biologists without a strong bioinformatics background. For the less informed species, ProteINSIDE allows searching PPi in well-informed species thanks to homologous IDs. For this, ProteINSIDE automatically selects homologous IDs from its database for the wanted species. Nevertheless, user can choose to run a local Blastp to select the species with the highest sequence homology with the proteins of the input dataset, and then ProteINSIDE proceed to the selection of orthologous IDs for this species. A functional annotation of all proteins from an extended network (PPi between proteins within and outside of the dataset) is done by clicking a button on the network visualisation. Results of this annotation are available as a new analysis. In addition to biological knowledge gathering, GO annotation, and analysis of PPi, ProteINSIDE also proceeds to an in silico secretome analysis [40]. For this purpose, ProteINSIDE merges four strategies of analysis: signal peptide [16] and cellular location [22] predictions, as well as a review of GO term annotation and cellular location recorded in UniProt. This four-step analysis provides a reliable prediction of proteins secreted thanks to a signal peptide. To our knowledge, ProteINSIDE is the unique all-in-one tool that predicts secretome from a list of gene or protein IDs [40].

Scientists are dependent on the species of study when they choose among resources available for their genomic and proteomic data analysis. Indeed, many tools are dedicated to only one species such as BioMyn for the Human [9] or DroPNet for the Drosophila [7]). Moreover, many tools are dedicated to diseases studies such as NetPath [13] and ToppGene [12]. ProteINSIDE has been first tool designed for genomic and proteomic data analysis in ruminant species namely cattle, sheep, and goat. However, the lack of information on these species required us to add human, rat, and mouse species to do homologous analysis. Thus, IDs from these species are perfectly recognized and analysed by ProteINSIDE. To our knowledge, ProteINSIDE is the only resource that allows the user to recover biological knowledge from well-known species (human, rat or mouse) using IDs from ruminant species. This avoids losing information

since many sequences or annotations remain to be stored in public databases for ruminant species and especially for goat. To our knowledge, only AgBase [61], a manually curated gene annotation database for farm species, including cattle and sheep, is available for functional annotation. However, AgBase does not perform analysis of PPi or prediction of secreted proteins.

In this article we have presented the performances of ProteINSIDE, a new powerful workflow which gathers tools and public databases to retrieve biological information of genes or proteins lists from 6 species (Bovine, Ovine, Caprine, Human, Rat, and Murine). We have reported a tutorial to describe how to get and interpret the results of a basic and a custom analysis with ProteINSIDE. Currently, there is no tool that performs in one query the analyses proposed by ProteINSIDE. ProteINSIDE offers a friendly-user interface where user can view, work, and download the results of an analysis. ProteINSIDE gives also a single file containing all results of an analysis. Thus, ProteINSIDE offers a great support to analyse efficiently a large quantity of data from genomic and proteomic studies to gather and interpret results necessary to construct a new research hypothesis or answer to a single question.

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### **AUTHOR CONTRIBUTIONS**

Conceive and design ProteINSIDE web service: NK, JT, MR, and MB. Perform experiments and analysed the data: NK and MB. Write the paper: NK, MB, and BP.

# CONFLICT OF INTEREST DECLARATION

The authors declare no conflict of interest.

### SUPPLEMENTARY DATA

High resolution files of the main figures on the paper are available for download at <u>Genomics and</u> <u>Computational Biology online</u>.

### **ABBREVIATIONS**

BC: betweenness centrality

- CC: closeness centrality
- GO: gene ontology

ID: identifier

PPi: protein-protein interaction

Proteins without transmembrane segments (secreted outside the cell)					Re	search area	Downloa	d result	S	
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P01189	COLI_HUMAN	POMC	noTM	Secreted		GO:0005576 GO:0005615 GO:0030141 GO:0034774			4 Signa	al peptide
P02766	TTHY_HUMAN	TTR	noTM	Secreted		GO:0005576 GO:0005615 GO:0030198 GO:0070062			4 Signa	al peptide
Q15848	ADIPO_HUMAN	ADIPOQ	noTM	Secreted		GO:0005576 GO:0005615 GO:0034383 GO:0070062		4 Signa	al peptide	
Proteins wi	ith functional an	notation related to	to a secretio	on pathway • Name	Targe Cho	Presults ose a value v 10	Download	oad res	ults	B
Proteins	Protein ID	Gene Name	Subcell	ular location		GO related to secretion		#GO terms	TargetP resu	ults 🔻
P60174	TPIS_HUMAN	TPI1		- 1	<u>30:00056</u>	O:0005615 GO:0070062			Other secretory	pathway
P53621	COPA_HUMAI	N COPA		Secreted 9	GO:00056	15 GO:0006890 GO:0030126 GO:0030157 GO:004820	5 GO:0070062	6	-	
P06744	G6PI_HUMAN	I GPI		Secreted 9	GO:00700	<u>62</u>		1	-	
				27 - 27 - 27 - 27 - 27 - 27 - 27 - 27 -						

**Figure 6. Prediction of secreted proteins.** Proteins potentially secreted are listed as two or three downloadable dynamics tables. (A) The first table lists proteins predicted as secreted by SignalP. The column "Peptide" provides the results for a positive identification of a signal peptide on a protein sequence as provided by SignalP. Identified peptides can be "noTM" (not transmembrane) or "TM" (transmembrane), only "noTM" are listed in the first table. The column "Subcellular location" provides the location of the protein declared in the UniProt database. The column "TargetP" provides the prediction of the subcellular location of the protein by TargetP software, and GO related to secretion are also listed to improve the prediction. A second table lists proteins with the "TM" prediction of SignalP, not shown in the figure since there was no result with the sample dataset. (B) A third table lists proteins, and subcellular location are also selected to improve the prediction. In this table, GO terms, TargetP prediction, and subcellular location are also selected to improve the prediction.



Pi results table	Nesearchare			a dura a d	
Gene Name A	Detection method	Gene Name B Detection method Nu Choose a value • 10	29 by 28 protein(s	HUMAN	В
Interactor A	Gene Name A	Detection method	Interactor B	Gene Name B	Database source
075489	NDUFS3	anti bait coimmunoprecipitation	<u>075306</u>	NDUFS2	IntAct
075489	NDUFS3	blue native page	<u>075306</u>	NDUFS2	IntAct
P00338	LDHA	two hybrid	P00338	LDHA	IntAct
P01225	FSHB	x-ray crystallography	P01215	CGA	IntAct
P01241	GH1	anti bait coimmunoprecipitation	P01241	GH1	IntAct
P04075	ALDOA	two hybrid	P04075	ALDOA	IntAct
P04075	ALDOA	two hybrid	P05062	ALDOB	IntAct
P04406	GAPDH	two hybrid	P04406	GAPDH	IntAct
P06733	EN01	two hybrid fragment pooling approach	P04075	ALDOA	IntAct
P06733	ENO1	two hybrid fragment pooling approach	P52789	HK2	IntAct

**Figure 7. PPi results and visualisation.** Results for PPi are summarised as a downloadable table and a diagram. (A) Main results are downloadable as table and network file that can be visualized using a network viewer (as Cytsocape). An online network view (made using the Cytoscape web application) is also proposed from this page result. A pie diagram indicates the number of PPi identified with the different detection methods. (B) A dynamic table lists linked proteins within the dataset, the detection method used to identify the interaction, and the database source of the interaction.



Figure 8. Network visualization of the PPi results. (A) This menu provides options to filter the network by: detection method, number of interactions for a protein, type of layout, protein ID, or the values of centralities. The centralities values are useful to sort large networks and to view only a central subnetwork. The betweenness centrality quantifies how frequently a node is on the shortest path between every pair of nodes for detecting bottlenecks in a network. The closeness centrality quantifies how distant minimal paths are from a given node to all others, a large closeness indicates that a node is close to the topological center of the network. (B) The network view is a dynamic image where user can access to a protein data by clicking on a node (name, function, statistic results, and database source and link of the protein).

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3] 🕑 Use IEA	A evidence	for GO																
Prediction of	fsignal	peptides (power	red with Signa	IP 4.1):	Selec	t the	e prediction	of secreted n	rote	ins and the	option							
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	8	GeneMANIA	<u>More th</u> GeneMA data. As domain	n 100k in NIA finds sociation imilarity	other genes th data include pr	nat are otein a	related to a set of ir ind genetic interactio	nput genes, using a ver ons, pathways, co-expr	/ large s ession,	set of functional ass co-localization and	sociation protein	PPi	-	-	PRED		-	÷
		12D-IMEx	Less the To facilia integrate	n 100k in ite experin d known,	teractions: mentation and experimental a	integra and pre	ted computational a dicted PPIs for five	nalysis with model orga model organisms and h	nism Pl uman ir	PI networks, we ha n the I2D database.	ive	PPi	NPi	SPi	-	CURE	IMEx	-
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**Figure 9. Setting up a custom analysis.** Firstly, user has to enter a name for the analysis, select the species of study, and directly paste the input IDs (Figure 2). User has to select settings of the analysis: the setting followed by "software" mention activates the corresponding module in the workflow, and then user can select options for chosen module(s).





**Figure 10. GOTree network visualization**. Linked GO terms which annotate the dataset are linked using ancestor chart method. Each edge means that a term A is a subtype of a term B (is\_a). Information about a GO is obtained by clicking on the GO or the node. Red colour is only for the GO terms relative to the Molecular Function. The degree of colour saturation is related to the number of proteins annotated by a GO (dark and clear for high and low numbers, respectively).



**Figure 11. Extended network of PPi with proteins outside of the dataset.** This network is made of PPi retrieved by querying the BioGrid, UniProt, and IntAct databases and using PPi with human proteins outside of the dataset. Grey squares are for proteins outside the dataset; white proteins are from the dataset. We have highlighted linked proteins that are involved in pathways such as: (A) glycolysis, (B) hormone activity, (C) the growth hormone signalling, and (D) thyroid hormones signalling. (E) We have used high values of betweenness and closeness centralities (BC: 3600; CC: 0.2) to get the most central proteins of this extended network.

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# ENDNOTES

- <sup>a</sup> <u>http://www.cbs.dtu.dk/services/SignalP/performance.php</u>
- http://www.cbs.dtu.dk/services/TargetP/instructions.php