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Computational Knowledge Integration in Biopharmaceutical Research

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(Article begins on next page)

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Computational knowledge integration in biopharmaceutical research

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Abstract

An initiative to increase biopharmaceutical research productivity by capturing, sharing and computationally integrating proprietary scientific discoveries with public knowledge is described. This initiative involves both organisational process change and multiple interoperating software systems. The software components rely on mutually supporting integration techniques. These include a richly structured ontology, statistical analysis of experimental data against stored conclusions, natural language processing of public literature, secure document repositories with lightweight metadata, web services integration, enterprise web portals and relational databases. This approach has already begun to increase scientific productivity in our enterprise by creating an organisational memory (OM) of internal research findings, accessible on the web. Through bringing together these components it has also been possible to construct a very large and expanding repository of biological pathway information linked to this repository of findings which is extremely useful in analysis of DNA microarray data. This repository, in turn, enables our research paradigm to be shifted towards more comprehensive systems-based understandings of drug action.

INTRODUCTION

Private companies undertaking pharmaceutical research normally take stringent measures to protect their research results as private intellectual property (IP), until they achieve a marketable compound. An unintended consequence of treating research as IP is that many of the research teams within a single company cannot effectively share their research results with each other, because they do not contribute to the normal public academic discourse of peerreviewed journals and its accompanying widely-available repositories of scientific information. This public system, to which they do not routinely contribute, is the only system for widespread knowledge dissemination available internally.

Pharmaceutical company researchers are predominately consumers, rather than suppliers, of content for the published literature and its knowledge bases. Consequently, while their peers in academia and among commercial competitors are blocked from access to important proprietary discoveries, so are their (non-competing) colleagues elsewhere in the same company. Likewise the company may lack an effective historical memory of its research, outside laboratory notebooks and personal computer files, if it does not attempt to fill the gap left by opting out of public discourse. We call this gap the 'IP shadow', and we believe it creates scaling limits to productivity in private pharmaceutical research — possibly even creating negative economies of scale experienced as an 'innovation deficit'.^{1,2}

The authors believe most pharmaceutical company researchers can, from their own experience, readily supply examples of wasted effort, lost opportunities and diminished productivity at their companies, owing to inability to

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Organisational memory

Framework

communicate, integrate and share research findings across intra-company organisational and geographical boundaries. A systematic approach to resolving these issues may become increasingly vital in the context of 'extended enterprises', which now appear to be the mode in the pharmaceutical industry.³

This paper describes a comprehensive initiative, *MyBiology*, designed to resolve the problem of 'IP shadow' using computational knowledge management, lightweight process re-engineering and web deployment. At the same time, this initiative attempts to:

- create a more sophisticated infrastructure than is currently available to the public researcher;
- build in process design and process adoption support from the beginning, realising that adoption and use are critical success factors; and
- develop advanced capabilities for computational systems biology as part of the infrastructure.

MyBiology deals, in part, with a version of the 'organisational memory' (OM) problem in knowledge management. Previous OM efforts have been primarily concerned with 'business process', that is, with better recollection and dissemination of *how* the enterprise does its work. Ontology construction has been seen as central to achieving this goal, which in turn is presumed to influence enterprise productivity and excellence of product.^{4–6}

Scientific knowledge in a biopharmaceutical enterprise, however, *is itself a product*. Therefore, managing this knowledge is actually part of the production process, and failure to manage it properly contributes to wrong decisions and attrition in the pharmaceutical development pipeline.

MyBiology is a collaboration between Millennium Pharmaceuticals and

Ingenuity Systems and is implemented at Millennium across multiple research sites on two continents. Initially focused mainly on results relatively far 'upstream' in the research and development process, it is rapidly being extended to have productivity impact all the way to the clinic. The system is described below and results and prospects after one year of the programme are assessed.

THE PUBLIC KNOWLEDGE MANAGEMENT INFRASTRUCTURE

Public academic researchers share information and collaborate on a large scale through the well-organised system of peer-reviewed academic journals. Over 4,600 of these journals are continuously indexed and organised into MEDLINE^{®7} by the US National Library of Medicine using a formal ontology, the Medical Subject Headings (MeSH[®]).⁸ This constitutes a knowledge base (KB). Many other databases have been linked to this fundamental resource, including whole genomes for various organisms of clinical interest. These resources are readily searchable on the web via PubMed[®],⁹ providing a critical means of information sharing for the scientific community. Furthermore, there are numerous computational methods available to analyse research data, such as DNA sequences, against the organised findings of other researchers.¹⁰

Taken as a whole, these facilities constitute an open, public framework for scientific knowledge management. The functionality of this framework must at least be matched, if not bettered, within private research organisations, to achieve parity with public efforts in sharing scientific knowledge.

The public framework is composed of:

- a peer-reviewed publication system;
- a knowledge base (MEDLINE);
- an ontology by which the KB is organised (MeSH);

Infrastructure

- methods for searching the KB (PubMed);
- computational methods for analysing new research data against the KB;
- portals that collect search methods and computational tools in a single site (NCBI Entrez¹¹);
- a transparent electronic framework providing information accessibility, identity and interoperability (internet, W3C web protocols, registry systems).^{12,13}

As private companies choose to replicate these elements, they must decide whether to adopt the public technology or to improve upon it. They also encounter the challenge of creating 'semipermeable' links back to the public system.

Semi-permeable links

THE MYBIOLOGY INITIATIVE

MyBiology was designed to eliminate the 'IP shadow' at Millennium and within Millennium collaborations by:

- providing one or more curated internal electronic journals of key research results;
- extracting computable findings from the internal journal into a KB founded on a 'deep' ontology;
- integrating internal findings with external public research using the same ontology;
- analysing transcriptional profiling data against biochemical pathways described in the KB using novel algorithms;
- providing both text-based KB query and graphical interface to pathway analysis.

MyBiology integrates Millennium's proprietary information with information

from public literature to help scientists build an evolving picture of complex biological systems. As shown in Figure 1, *MyBiology* consists of several related components. Millennium scientists are an integral part of *MyBiology*, both as contributors and customers.

The major components of *MyBiology* today are:

- an ontology;
- Millennium's research scientists;
- knowledge intermediaries (KiMs);
- Journal of Millennium Science (J. Mill.);
- external and internal databases;
- MyTargetValidation (MyTV);
- the Life Science Knowledge Base (LSKB) and query interface;
- the public scientific literature;
- Pathway Analysis Research Information System (PARIS).

An *ontology* is critical to organisation of a KB, just as unique object identifiers and foreign key relationships are fundamental for various kinds of databases. Recent work on scientific ontologies has attempted to provide specialised annotation and organisation for genomics and genome-related research databases (eg the Gene Ontology Consortium, GO¹⁴) and computational service classification and organisation¹⁵ via fundamental expressive formalisms relevant for both purposes and suitable for deployment on the internet.¹⁶

For our basic ontology, we selected a system from Ingenuity Systems, Inc., together with annotated content, to which significant proprietary and inlicensed content were added. The Ingenuity ontology has many advantages, among which are the ability to distinguish

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Ontology

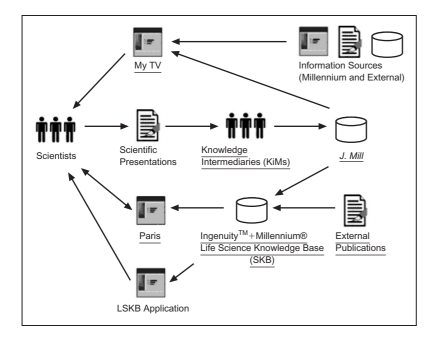


Figure 1: The MyBiology system at a high level component view

Repository	between objects and processes. The resulting richly-connected KB means it is possible, for example, to extract metabolic and cell-signalling pathways automatically and combine them with other information sources for computational analysis. <i>Research scientists</i> are both providers and consumers of information. They present their experimental methods, results and
Experimental findings	interpretations at regular research team meetings throughout the company. Finding ways to capture the information in these presentations and to make it readily available to scientists across the
Encoding	company is a critical component of the <i>MyBiology</i> effort. <i>Knowledge intermediaries</i> work with researchers to capture and encode the
Computation	information found in research scientists' presentations. KiMs are PhD-trained biologists and chemists with additional training in computational science. The system operates effectively with a very small number of KiMs supporting more than 1,000 researchers. <i>The Journal of Millennium Science</i> is an electronic journal constructed upon the Documentum ECM Platform ¹⁷ using customised interfaces. Scientific presentations, typically in PowerPoint, are

stored with metadata in *J. Mill.* Metadata include an abstract, author and publication information and unique identifiers for targets, compounds or other reagents referenced in the presentation.

MyTargetValidation is a scientific portal to pertinent information on a given biopharmaceutical target (gene/protein). It integrates a large number of internal and external databases using a Web Services Architecture.¹⁸ It also enables direct access to assay results in lead optimisation for molecules active against a target, through integration with an internal drug discovery database of compounds, assay results and basic computed compound properties.

The *Life Science Knowledge Base* is a highly-structured KB that combines selected findings from the public literature with internal Millennium research results in computable form. Ingenuity employs a large number of 'content scientists', to read articles from 33 highly cited scientific journals, curate the results and interpretations, and add them to the KB. They also use natural language processing (NLP) to add information from MEDLINE abstracts dating back to 1980. The KB contains more than 1 million findings, approximately 800,000 derived by content scientists.

Lifesciences is a web-based application from Ingenuity that provides textual query access to the LSKB including a comprehensive overview of all findings related to a given gene, its products and biological relationships.

The Pathway Analysis Research Information System¹⁹ analyses the results of a transcriptional profiling experiment against the biological pathway information in LSKB. It enables deeper interpretation of cellular processes affected in an experiment, by interpolating nontranscriptionally regulated components of relevant pathways.

Several stages of scientific research in support of pharmaceutical discovery and development are supported by these components. For example, target identification and validation are enabled PARIS

Integration

Target validation

by MyTV and *J. Mill.* as individual researchers evaluate results from genomescale experiments and seek to better understand the role that individual genes and their products play in the system under study. In characterising individual compounds, PARIS is often used better to understand the mechanism of action of the chemical, as it provides important context to understanding the large numbers of genes that are typically affected in a single experiment. The PARIS pathway browser is further used to explore the interconnections between proteins identified in an experiment.

Furthermore, there has been great value in extending J. Mill. into areas outside biology; several different groups of researchers have approached the KiMs about including reports and other scientific results in J. Mill. so that they are more broadly available and are integrated with the rest of the Millennium knowledge integration platform. PARIS is having a dramatic impact on the analysis of clinical trial data in the form of evaluating patient responses to candidate pharmaceuticals. Through its pathway mapping and interpretation, it provides analysts with context sufficient to direct further research experiments in support of the therapeutic goal.

The most fundamental goals of this initiative are to enable scientists to employ all available scientific information public and private; to select targets and compounds most likely to succeed in clinical development; and to conduct more effective clinical trials. Ultimately, we wish to shift our research paradigm away from a dyadic compound \rightarrow target view and towards a more holistic compound \rightarrow pathway \rightarrow disease understanding. Increases in productivity here should result in faster availability of improved therapies to patients.

MYTARGETVALIDATION

Researchers seeking to understand the biology of a system under study seek information from many sources, including public literature and external websites with information on gene and protein families (eg NCBI²⁰ and GO). Also, researchers at Millennium have typically interacted with core technology groups via the web to request experiments and/ or access the results. This led to a proliferation of individual unconnected websites and to redundant requests for identical experiments from different researchers. Researchers expressed enthusiasm for a single point of entry for access to multiple sources of information from both external sources and internal resources.

MyTV is a scientific portal (see Figure 2) to all available information on a given target, whether external or internal. It begins to answer the question: 'What do we know about this target?'

Access to information in MyTV is valuable in many different contexts of pharmaceutical development including target discovery, target validation, assay development, lead optimisation and analysis of clinical data. At a glance, a scientist can quickly assess what research has been performed at Millennium, by whom and in what stage of the pipeline; what types of experimental data are available; and what information is available from the public domain regarding biological targets and their annotation.

Public information is acquired through mappings to identifiers provided by Millennium's Gene Catalog system — a non-redundant catalogue of all genes constructed from both public and proprietary DNA sequences and annotations. Public sources include: SWISS-PROT,²¹ LocusLink,^{22,23} OMIM,²⁴ GO, InterPro²⁵ and ENZYME.²⁶ MyTV also links targets to Ingenuity's LifeSciences GeneView, providing access to key findings from the public literature and internal findings captured in *J. Mill.* and curated into Ingenuity's KB.

Millennium's other internal sources include:

• GeneCatalog for information on

My Target Validation: MG10655.5 TOP1_HUMAN DNA topoisomerase I (EC 5.99.1.)	2).	Search:
Gene Summary	Expression Analysis	Results and Interpretations
Mine Number MG10655.5 (was MG1287497, MG1324269, MG1548436,) Mine Classification	TaqMan Expression Explorer Oncology General Phase II Panel Human Phase 1 MLN 944-Treated Colon Cell Line Panel	Journal of Millennium Science Further studies of MLN944 mechanism of action: Not a topoisomerase inhibitor Laura Rudolph-Owen, 1/20/03
topole eukaryotic DNA topoisomerase I	(Run 1)	<u>ML944: Studies on mechanism of action</u> Darshan Sappal, 9/20/02
Curated Sequences (BasePerfect) No Curated Sequence (Base Perfect	<u>MLN 944-Treated Colon Cell Line Panel</u> (Run 2)	Defining mechanism of action for MLN944 and MLN576. Laura Rudolph-Owen, 7/17/02
Sequence) available	Synopsis Exp. 3888 (Up in Th2 Ohr vs Th1 Ohr)	Suggest new J. Mill entry
SwissProt Q9UI54 P628_HUMAN Protein PRO0628. P11387 TOP1_HUMAN DNA topoisomerase I (EC 5.99.1.2).	Exp. 3888 (Up in Th2 Unr Vs Th1 Unr) p53ER late downregulated p53 transient up MPMx 30K Ovarian NOE vs Tum (POOF	Target Advancement First Pass Reports No First Pass Assessment files found.
LocusLink 7150 topoisomerase (DNA) I	1.50 to 1.85) Exp. 4428 (Up in Mast cells stim with IL9 and IgE compared to no IL9)	TAPAS (Target Partnership) Reports No TAPAS files found.
Chromosome Human chromosome 20	<u>Show more</u> (20 items in all)	Comments
Orthologs Saccharomyces cerevisiae, Rattus rattus,	Experiments by Gene Folder (TaqMan) No old (pre-TAQEE) TaqMan files found.	Add new comment
<u>Mus musculus</u>	Inflammation TaqMan Folder No Inflammation TaqMan files found.	Gene Ontology GO Annotation
Gene Variation No SNP data found for MG10655.5	Neurobiology TagMan Folder	DNA topoisomerase I

Figure 2: Example MyTV page for DNA topoisomerase I

sequences, protein families, chromosome location and orthologues.

Content aggregation

Caching

Productivity

Web services

• Reagents and experimental systems that include information on expression vectors, expressed proteins, microarrays and experiments done in molecular pathology including TaqEE, a warehouse of TaqMan expression data.

- Scientists' analysis and interpretations are surfaced through links to Synopsis, a system for capturing interesting sets of genes and their biological context; *J. Mill.* for key scientific presentations and findings; summary reports from the Target Advancement and Assay Development groups.
- MyDrugDiscovery, Millennium's portal for information on highthroughput screening of targets and other assay information about compounds and lead series.

MyTV uses a pull model to query multiple systems for summary information about each target. Each system to be queried implements a simple web service returning an XML²⁷ fragment adhering to the MyTV XML schema.²⁸ The system then constructs an XML file for each target. This XML content is transformed into HTML^{29} by the presentation layer.

The system uses on-demand instantiation: target content is not aggregated until a scientist requests it. All requested information is then cached on the file system. Cache is periodically updated to refresh the information.

The system was initially released in a basic configuration in May 2002. Frequent releases were made, integrating new systems as requested, adding significant functionality over time, and continuously improving performance. Within four months, MyTV developed a user base of over 100 scientists. On a typical day, 20-25 individuals use the system. After about five months of use, users were surveyed to assess their perception of MyTV's impact on their work. Scientists strongly agreed that the system improved their productivity (they spend less time finding information) and quality of decision making by providing more complete and relevant information.

JOURNAL OF MILLENNIUM SCIENCE

J. Mill. is a collection of software, processes and semi-structured data. It serves as a central repository of experimental results and interpretations. J. Mill. is designed to maximise the connectedness of research teams across

Documentum [©]	therapeutic disciplines, create an	Documentum Foundation Classes
	organisational memory of key scientific	(DFCs) ³² to access Documentum's
	findings with links to supporting data and	eContent Server, which stores J. Mill.
	serve as a staging area for findings export	documents and presentation metadata
	to the highly structured KB. It was	manages the document life cycle. In
	considered helpful to use a framework	addition, several open source framewo
	similar to one that researchers are familiar	and components are used, including
	with, that of publication in a scientific	Jakarta's Struts ³³ which enforces a mod
	journal.	view-controller (MVC) ³⁴ architecture
Scientific	J. Mill. relies on the fact that most	within the application. The presentation
communication	scientific communication occurs in a	layer is written using JavaServer Pages
	group setting and is supported by	(JSPs) ³⁵ and the Struts tag libraries to
	electronic documents such as	format the presentation for the user.
	PowerPoint ³⁰ presentations containing	Authentication of users is accomplis
	scientific results and interpretations. Since	via a web-based single-sign-on (SSO)
	these documents form the core of the	web agent.
	communication between scientists on a	J. Mill. makes itself available to othe
	project team, if they were archived with	applications through the publication o
	some associated metadata, a record would	simple object access protocol ³⁶ (SOAF
	be available of the important activities	interface that allows other applications
	undertaken by the project team.	execute queries against J. Mill. content
	By providing a central location for	The SOAP interface requires the same
	scientific results, it was hoped to	user authentication as the web applicat
	encourage information sharing and reuse	and uses the same SSO interface to
Repository	among Millennium scientists and to	control access.
	prevent critical information from being	J. Mill. was initially put into produc
	lost or overlooked. J. Mill. is accessed via	in June 2002. Since that time,
	a web-based application which allows	approximately 100 articles by Millenni
	scientists to browse the repository and	authors have been published internally
	KiMs to publish entries in the repository.	the future, J. Mill. will provide self-
	Published J. Mill. entries are also available	publication facilities to authors in orde
	through MyTV.	expand its ability to collect and publish
	An entry consists of documents	content more rapidly to our scientists.
	describing an internal Millennium	
Metadata	presentation and curated metadata about	LIFESCIENCES FOR
	that presentation. The documents are	MILLENNIUM
	typically Microsoft Word ³¹ or	The Ingenuity LifeSciences Suite (Fig
	PowerPoint files that have been presented	3) is a user interface to the Ingenuity
GeneView	within Millennium along with metadata	Pathways KB. It includes GeneView,
	about the presentation curated by a	which catalogues the functions of a give
	Millennium scientist. The metadata	gene and gives scientists access to
	include information such as the author,	functional assertions between a specific
	gene and compound names and identifiers	gene and other genes, small molecules
	and some key scientific findings distilled	cellular processes (Figure 3), as well as
	from the presentations.	specific biological experiments suppor
		each assertion. Most features of
	J. Mill. application	GeneView run dynamically off the KE
	The J. Mill. application is a three-tier web	for example, statements regarding
	application. Its user interface for browsing	biological experiments are generated
	is deliberately reminiscent of PubMed,	using natural language algorithms appl

cuments and presentation metadata and mages the document life cycle. In dition, several open source frameworks d components are used, including carta's Struts³³ which enforces a modelew–controller (MVC)³⁴ architecture thin the application. The presentation er is written using JavaServer Pages Ps)³⁵ and the Struts tag libraries to mat the presentation for the user. Authentication of users is accomplished a web-based single-sign-on (SSO)

I. Mill. makes itself available to other plications through the publication of a nple object access protocol³⁶ (SOAP) erface that allows other applications to ecute queries against J. Mill. content. e SOAP interface requires the same er authentication as the web application d uses the same SSO interface to ntrol access.

J. Mill. was initially put into production June 2002. Since that time, proximately 100 articles by Millennium thors have been published internally. In future, J. Mill. will provide selfblication facilities to authors in order to pand its ability to collect and publish ntent more rapidly to our scientists.

FESCIENCES FOR ILLENNIUM

e Ingenuity LifeSciences Suite (Figure is a user interface to the Ingenuity thways KB. It includes GeneView, nich catalogues the functions of a given ne and gives scientists access to nctional assertions between a specific ne and other genes, small molecules and lular processes (Figure 3), as well as the cific biological experiments supporting ch assertion. Most features of eneView run dynamically off the KB; example, statements regarding ological experiments are generated using natural language algorithms applied to structured information stored in the knowledge server. Each new piece of

with which scientists are already familiar.

J. Mill. is written in Java and uses the

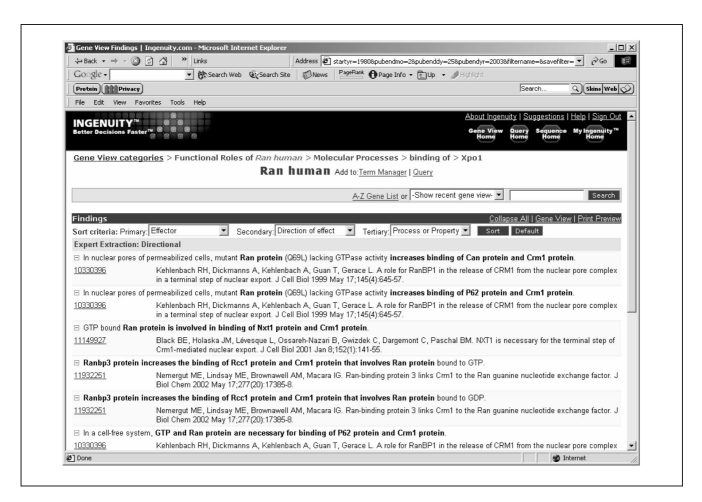


Figure 3: A GeneView page displaying relationship between Xpo1 and Ran provides the user with a natural language display of the underlying relationships and properties of that object, shown here. Note that findings were also returned containing the 'Crm1' synonym of Xpo1. This synonym capability resides in the Ingenuity ontology

information is hyper-linked to other relevant findings in the KB. Alternatively, scientists using the Query **KB** Query application may pose a specific question regarding relationships between biological entities; for example, 'What apoptotic effects does cytochrome c exert in neurons?' The fields in Query, which Findings reflect the perturbations and results recorded in most biological experiments, enable scientists to construct these questions efficiently and intuitively. Query intelligently leverages the underlying ontology to empower the biologist: for example, a query about neurons returns findings regarding superior cervical ganglion cells, as the ontology recognises the ganglion cells as a type of neuron (Figure 4).

Two methods are used to capture findings. In findings capture through expert knowledge acquisition, qualitycontrolled operational processes enable Ingenuity-trained, PhD-level scientists to structure information from full text articles with a high degree of semantic richness. This method was applied to tens of thousands of articles from the most highly cited biological journals. The second method, semi-automated knowledge acquisition, employs algorithmic capture of knowledge from the abstracts of hundreds of journals spanning more than a decade of published research. In both approaches, every finding structured into the Ingenuity KB was quality-checked by a PhD-level scientist.

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Google -	▼ 😚 Search Web	Search Site News	ageRank 🚯 Page Info 🔹 🔁 Up 🔹 🥒 Highlight		
Protein Privac	y)			Search Q Skins W	ab 🕤
File Edit View	Favorites Tools Help				1.*
			About In	<u>genuity Suggestions Help Sign (</u>	Dut
INGENUITY Better Decisions Fai			Gene V		
better becisions ra	0 0 0 0		Hom		
Ouery Results >	> Query Findings				
	,				
Findings				<u>apse All Query Results Print Previ</u>	ew
Sort criteria: Prim	ary: Effector 🗾 Se	condary: Direction of effect	Tertiary: Process or Property	ort Default	
Expert Extraction	: Directional				
⊡ In cytosol, mou	se <u>Casp9</u> protein is necessary f	or the cytochrome C protei	n-enhanced apoptosis of sympathetic neu	r on deprived of Ngf protein.	
12223555	Deshmukh M, Du C, Wang J Neurosci 2002 Sep 15;22		smac induces competence and permits casp	ase activation in sympathetic neuror	IS.
Anti-rat <u>Cytochi</u> protein.	rome c [Cycs] protein antibody d	ecreases the apoptosis of r	at superior cervical ganglion neurons that is	increased by depletion of Ngf	
9744886	Neame SJ, Rubin LL, Philp (6):1583-93.	ott KL. Blocking cytochrome	c activity within intact neurons inhibits apopto	sis. J Cell Biol 1998 Sep 21;142	
□ Rat Cytochrome	e c [Cycs] protein causes little c	or no change in the rate of	apoptosis of rat superior cervical ganglion n	eurons.	
<u>9744886</u>	Neame SJ, Rubin LL, Philp (6):1583-93.	ott KL. Blocking cytochrome	c activity within intact neurons inhibits apopto	sis. J Cell Biol 1998 Sep 21;142	
□ Rat <u>Cytochrome</u> depletion of Ngf		or no change in the rate of	apoptosis of rat superior cervical ganglion n	eurons that is increased by	
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🗉 Cytochrome C	protein causes little or no char	ige in the apoptosis of neu	rons that involves Ngf protein.		
<u>12223555</u>	Deshmukh M, Du C, Wang J Neurosci 2002 Sep 15;22		smac induces competence and permits casp	ase activation in sympathetic neuror	IS.
🗉 in cytosol, pCP	T-cAMP decreases the cytochro	me C protein-enhanced ap	ptosis of sympathetic neuron deprived of N	Vgf protein.	
12223555	Deshmukh M, Du C, Wang J Neurosci 2002 Sep 15;22		smac induces competence and permits casp	ase activation in sympathetic neuror	IS.
🗉 in extosol nota	ssium chloride decreases the o	otochrome C protein-enhar	ced apoptosis of sympathetic neuron dep	ived of Naf protein	

Figure 4: Query result after search for apoptotic effects of cytochrome C on neurons. Note that superior cervical ganglion neurons were recognised as a type of neuron

Frame-system

SEMANTIC INTEGRATION: THE MYBIOLOGY KB

To build a platform for biological computation — as well as for intelligent search and query - it was thought necessary to integrate the large corpus of literature-based findings in the Ingenuity KB with key experimental results from Millennium's research programmes into a single, computable structure. This posed multiple challenges; for example, semantic inconsistency in the literature makes it difficult to recognise when genes are identical or distinct. A need for a high degree of accuracy impedes the rate at which experimental findings can be formally represented. Finally, more sophisticated computations will probably require detailed context regarding functional relationships - for example, in

what cell type one protein phosphorylates another.

Frame-based knowledge representation systems offer a versatile and powerful approach to structuring knowledge while addressing these critical needs. Generally, frame-based systems allow definition of abstract concepts and their relationships with other concepts.³⁷ These systems define three types of formal objects to represent knowledge in a given domain: concepts, properties and instances. Concepts (classes) are descriptions of particular categories of objects. Properties are attributes that describe the concept itself or relate one concept to another. An instance is a real-world example of a concept. Once information is represented in this manner, most frame-based knowledge representation systems support basic inference capabilities such as classification ('Is this protein a type of kinase?') and declaration of axioms ('If loss-of-function mutations in a gene lead to a DNA repair phenotype, that gene participates in DNA repair'). Axioms in particular impose semantic constraints on the KB that help maintain the consistency and integrity of the data. Finally, framebased knowledge representation systems provide basic query capabilities for retrieving stored data.

Using a frame-based knowledge representation, one can build an ontology (Figure 5): a taxonomy and formal description of the concepts and relationships germane to a particular domain. Ontologies can be used in many different ways. By defining a class structure and formalising relationships between those classes, ontologies (i) allow more efficient browsing of concepts through the taxonomy, (ii) provide a standard framework for exchange of data, and (iii) provide the basis for representing domain knowledge. Building the ontology for a frame-based system is analogous to building the schema for a relational database. A KB consists of an ontology populated with instances of real data. In academia, frame-based representations of biological data have proven useful in several biological domains across a range of organisms.^{38–41}

Ingenuity has developed a frame-based system with the goal of representing hundreds of thousands of functional relationships between genes for the purpose of enabling computations that provide critical path insights to scientists involved in therapeutic discovery. Toward this end, Ingenuity has constructed a proprietary ontology of more than 300,000 distinct classes including, but not restricted to, genes proteins, small molecules, cellular

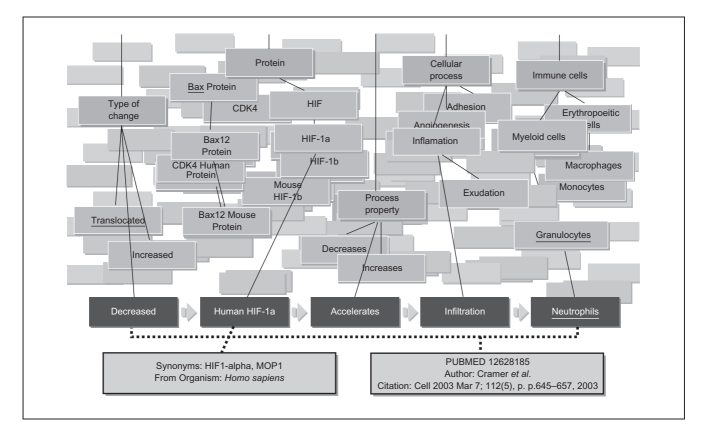


Figure 5: A conceptual model showing how a biological finding is formally represented in the Ingenuity Pathways KB using objects structured to capture meaning and context from the scientific experiment. Actual findings contain more semantic detail than displayed here, and the actual ontology is composed of more than 300,000 classes

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Biological ontology

	components, cells, tissues and biological	(
	processes. Ingenuity has utilised this	i
	ontology to formally represent, in]
	machine computable form, more than a	1
	million biological experimental findings	1
Experimental findings	from the public domain literature,	
	specifically focused on human, mouse and	
	1 · · · ·	3
	rat genes. The Ingenuity KB can be used	
	to structure not only knowledge from the	
	public domain, but also a pharmaceutical]
	organisation's proprietary scientific	1
	findings, creating a leveraged, integrated	ŝ
	knowledge asset. This project has	1
Scalability	required scalability for both the ontology	1
	and the back-end not previously	4
	demonstrated in the smaller, academic	ę
	efforts at frame-based systems described]
	above. Ingenuity currently has operational	1
	processes capable of incorporating, per	1
	month, more than 5,000–10,000	1
	concepts into its proprietary ontology,	
	and more than 40,000 biological findings	
	into the KB.	
]
	Applications have been developed to	(
	leverage the Ingenuity KB in support of a	é
	diverse array of complifations enabling	
	diverse array of computations, enabling	
Findings capture	the discovery process for end-user	
Findings capture	the discovery process for end-user biologists — including both systems	
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capturing and curating Millennium's internal research results for entry into the KB was needed. The challenge, of course, lay in the various forms that such results may take in a modern pharmaceutical company (notebook entries, annotated gene lists, PowerPoint presentations, etc.).

In approaching this problem, the premise was that demands may only be made on scientists (eg to contribute scientific findings to an electronic repository) in proportion to the value they perceive from using the system. The 'bootstrapping' problem was how to get scientists to contribute as 'producers' of knowledge before there existed a critical mass of content within the repository for them to draw upon as 'consumers.' As an initial strategy, a small number of dedicated 'curation' roles were created in order to initially minimise the burden of participation upon scientists. These dedicated roles are currently being moved away from.

SCIENTIFIC FINDINGS CAPTURE

The scientific findings capture (SFC) process was developed as a systematic method for capturing presentations and their associated findings for publication Millennium-wide in both J. Mill. and the Ingenuity LSKB. The process is composed of several discrete steps and metrics gathering has been incorporated throughout. Initially, this work was started with the dedicated support of KiMs. KiMs served to assist individual research scientists with the process of knowledge sharing.⁴² SFC KiMs were PhD scientists with knowledge of the internal and external computational tools used by Millennium scientists. SFC has several steps, which are outlined below.

Encoding the presentation

Scientists serving as KiMs are invited to laboratory and departmental meetings where scientists present the results of their work to their colleagues. These presentations are typically prepared in PowerPoint and are excellent sources of key experimental results and interpretations. Using the *J. Mill.* web application (see above), an individual laboratory scientist's presentation is separated into components resembling a journal article. A title and abstract are prepared using background material, and hyperlinks to internal and external resources are also included within the application.

Publication

After initial creation in *J. Mill.*, the completed entry is returned via e-mail to the presenting scientist for final approval. In this manner, the role of the presenting scientist is similar to that of the author of a scientific manuscript in that he/she is ultimately responsible for its contents. Upon approval, the presentation is automatically published Millennium-wide through MyTV.

TRANSFERRING EXPERIMENTAL RESULTS FROM MILLENNIUM TO INGENUITY

To generate a combined KB of internal Millennium content and external Ingenuity content, a transfer system was devised to facilitate the transfer of Millennium experimental results (MERs) to a private copy of the Ingenuity KB.

All scientific presentations contain potential MERs to be represented in the Ingenuity KB. Those findings determined to be useful in the KB are queued for entry. All proposed novel terms and genes are examined and validated by both teams to verify accuracy. Also, clarification of findings from the KiMs is a necessity to ensure that the translation from the written sentence in J. Mill. to the Ingenuity KB is performed consistently. Each MER is modelled into the Ingenuity ontology by the Ingenuity content scientists and displayed in LifeSciences within the context of the findings derived from the external literature. Further, a hyperlink to the original presentation in J. Mill. is also available for easy reference as

well as increased searchability through Ingenuity's LifeSciences Query application.

TECHNICAL INTEGRATION: THE MYBIOLOGY ARCHITECTURE

A third but equally important aspect of integration in the *MyBiology* project was the technical integration of MyTV, *J. Mill.* and the *MyBiology* KB into Millennium's informatics platform within a coherent architecture, despite the rather complex technical environment.

The hardware and network for deployment were primarily Millennium's internal network but there was also a need to include some form of connectivity to the Ingenuity LSKB and Ingenuity's LifeSciences application. End-users used both Microsoft Windows and Apple Macintosh computers.

The *MyBiology* software environment included four applications (MyTV, PARIS, LifeSciences and *J. Mill.*). These applications had to be able to link to several other existing web-based applications. Each of the applications, existing and new, was targeted at different user communities and had been customised for their use.

Within this context, we also had the following requirements for the *MyBiology* architecture:

- User experience. Users must have a common user experience when using any of the *MyBiology* suite of applications, and it must be easy to navigate between applications.
- User access. The applications need to be accessible from both Microsoft and Macintosh computing platforms.
- Decoupled development. Asynchronous releases across the four development teams improved productivity.

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Electronic publication

Architecture

Integration

Knowledge transfer

Loose coupling

SOAP

Intracellular processes

MODEL AND DISCUSSION

To make transitions between applications seamless, the *MyBiology* software architecture (Figure 6) focused on the interconnections and transitions between the applications. To maintain loose coupling between the applications and databases, connections to application databases were limited solely to the applications that owned them. Connections between the applications are permitted at two levels. Low-level connections were made between applications using web services. User interface level connections were made between applications via HTML links. At the lower level, application-to-application connections were made using web services via the SOAP interface. A single standardised SOAP interface was exposed by all applications. These SOAP connections were used to pass data between the applications and to provide URL links that the client application then used to provide links between the applications at the user interface level. These SOAP connections were also used to connect to other applications outside the MyBiology initiative.

The exception to this form of

connectivity is the LifeSciences application. In this case, since the application was hosted outside Millennium, the applications are connected via periodic file transfers, over a secure network, of the data to be shared with Ingenuity's KB and the links back to the Millennium applications.

The architectural approach used in *MyBiology* has been successful in its primary goals of preserving loose coupling between the applications for ease of development while simultaneously presenting a unified seamless view of the applications to the end users.

THE PATHWAY ANALYSIS AND RESOURCE INFORMATION SYSTEM

The Ingenuity KB integrates large amounts of biological knowledge. Because this information is encoded in a structured format that enables computation, the Ingenuity KB can be used to solve one of the more difficult challenges in drug discovery: the identification of the intracellular processes perturbed or activated in a biological experiment.

The PARIS database of protein functional relationship integrates data

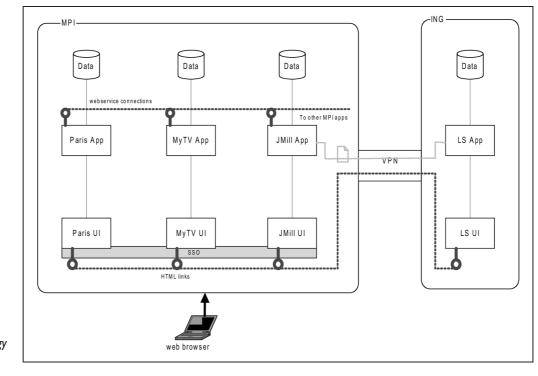


Figure 6: The MyBiology architectural model

	from both public sources (CSNDB, ⁴³
Graph	$LIGAND^{44}$) and the Ingenuity KB,
	including Millennium scientific findings
	encoded in the KB. This integration has
Pathway database	given Millennium a human pathway
Fatliway Uatabase	database much larger than any publicly-
	available resource.
	The detection of perturbed pathways is
	required to understand the function of a
	novel gene, the mechanism of action of a
	compound or the different responses of
	patients to treatment. Transcript profiling
Microarray	(TP) using microarrays is a powerful
	technology to perform such
	investigations: in a single assay the
	transcriptional activity of the entire
	genome can be measured. ⁴⁵
	Translating hundreds of observed
	mRNA changes in terms of perturbed
	pathways is work too complex to be
Neighbourhood	optimally performed by hand, however.
	Help can be provided to the researcher by
	using computation. ^{46,47} Such
	computation combines the experimental
	data with large amounts of pathway
	knowledge and extracts the portion of the
	knowledge relevant to the observed
	transcriptional activity. Besides helping
	the scientist deal with large amounts of
	data, appropriate computation can also
	significantly improve the sensitivity of TP
	data interpretation. Indeed, it is now well
	established that functionally-related genes,
	eg pathway neighbours, have a significant
Regulation	tendency to be simultaneously regulated
	at the mRNA level. ^{48–50} Therefore, even
	if the individual gene expressions stand in
	the range of experimental noise, the
	coordinated differential expression of
	genes belonging to a same pathway can
	indicate its perturbation.
	Computational interpretation of TP
	data requires large quantities of pathway
	knowledge. The coverage provided by
	current public sources ^{51,52} is relatively
	low. Moreover, to enable computation
	the data have to be consolidated into one
	format. The Ingenuity KB is an ideal
	source of pathway knowledge

Consequently, Millennium uses Ingenuity to maintain a system called the Pathway Resource and Information System (PARIS). The knowledge contained in PARIS can be summarised in a graph where vertices are genes and edges represent their functional relationships (Figure 7). Those relationships are: membership in the same complex, protein binding, posttranslational modifications, transcriptional regulation and neighbouring of enzymes in metabolic pathways. Currently the network contains about 5,000 genes and 22,000 interactions.

Millennium's computational biologists have developed algorithms that use this network to analyse TP experiments.¹⁹ The algorithms identify the portions of the graph that are significantly regulated in a given experiment (Figure 8). For each gene the average regulation in its pathway neighbourhood is computed. This local density of transcriptional activity is then compared with those obtained when randomly reassigning the individual gene expressions.

Figure 8 shows an example of application of the PARIS system to identify pathways perturbed upon p53mutation in ovarian tumour samples.⁵³ The PARIS system has extracted a set of genes having significant density of differential expression in their functional neighbourhood when comparing transcript profiling of p53 null samples against p53 wild-type samples. Those genes define pathways related to cell cycle and proliferation, tumour growth factor beta signalling and cellular adhesion.

The PARIS system, based on pathway knowledge extracted from the Ingenuity KB, is used at Millennium to analyse TP experiments performed at several stages of the drug discovery process. The computationally extracted pathways can point to the cellular processes downstream of a target, which are perturbed by a compound, responsible for drug resistance or involved in the different responses of patients to treatment.

The PARIS web application contains hyperlinks to MyTV and other sources to enable scientists to rapidly learn about genes regulated in their experiments.

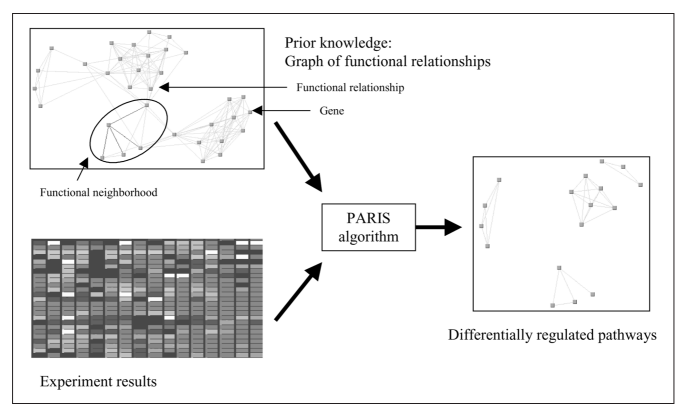


Figure 7: Interpretation of transcript profiling experiments using the PARIS system. Knowledge extracted from the Ingenuity knowledge base is represented as a simple undirected graph where vertices are genes and edges represent their interactions (eg binding). During a TP experiment analysis, each gene of the graph is scored for the density of differential expression observed in its neighbourhood. Genes at the centre of significantly regulated neighbourhoods (eg P < 0.05) induce a subgraph representing the regulated pathways

CONCLUSION

Biopharmaceutical-scientific knowledge is acquired through a long and expensive series of directed experiments, in which the supporting data can become quite voluminous. New experimental data must be posited against the conclusions from previous experiments. Integration and synthesis with public knowledge must be complete, but knowledge developed inside the enterprise must remain shielded so long as it is treated as intellectual property. Keeping knowledge hidden from competitors may result in it being hidden from the broader organisation. Our goal was to resolve these problems using a suite of technologies and organisational process change.

What was achieved? The system described in this paper is currently in production at Millennium and is being actively extended. It provides not only a long-term organisational memory for key science in the biopharmaceutical enterprise, with enhanced analytical capability for experimental data, but also the fullest record of inter-linked data and information to enable more informed decisions about the project pipeline.

Experiences with this system to date show that:

- (1) The semi-structured and lightweight *J. Mill.* KB is much more significant than was originally thought.
- (2) The availability of public domain, *highly structured* findings provides substantial value even in the absence of internal experimental results.
- (3) The ability to directly posit experimental data from transcriptional profiling upon the Ingenuity KB is extremely productive and has led to

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Transcript profiling
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Decisions

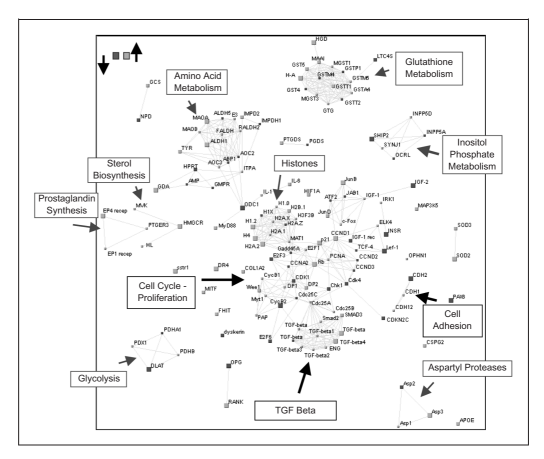


Figure 8: PARIS analysis of p53 null v. wild-type clinical ovarian tumours.⁵³ The graph shows the pathways that were computationally extracted using the PARIS system when comparing p53 null ovarian tumour samples with tumour samples that have not lost p53. Each gene is represented by a square whose size and shade reflects the differential expression between groups of tumours. Dark grey means more expression in p53 null samples and light grey less expression. A line between two genes indicates the existence in the PARIS system of a functional relationship (eg binding) between their products. The position of the genes on the picture tries to preserve their respective distances in the full network

key insights concerning possible drug mechanisms of action.

(4) It is important to direct the content acquisition of the KB, and to align it with the research.

J. Mill. was originally conceived mainly as a staging area for capture of the highly structured findings. As it turns out, it has become valuable in its own right as a primary repository for enterprise research findings in human-readable form. This is due both to the rich context provided by the underlying presentations and to the familiar PubMed usage pattern where scientists can skim a title and abstract before going to the original source material. While Ingenuity's highly structured KB is invaluable for computation, links to the original source material (both *J. Mill.* and public sources) provide critical additional background and context vital to correct interpretation.

The linkage from MyTV to *J. Mill.* also contributed to the effective use of *J. Mill.* and proved to be highly beneficial for gaining new insight into future experiments. One example case occurred when a scientist reviewing target information in MyTV encountered a link to related experiments in *J. Mill.* An examination of the presentation led to a change in future experimental design and a potential new research avenue.

The ability to capture and structure

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Semi-structured findings

Insight

Compound	scientific findings in the KB is crucially important, because proper structure allows computation across a very large corpus of knowledge. At present our internally generated knowledge constitutes only a tiny fraction of the KB — consequently most of its computational power comes from public knowledge, as expressed in the formal language of the KB's ontology. In this	 For comp further d has prove and exple mechanis one com different that had landscape
PARIS algorithms	context, its single most valuable application has been found to be in combination with the PARIS algorithms,	• Applying clinical d insights i
Clinical	in analysing transcriptional profiling experiments. In part, this is a reflection of the state of experimental technology and the power of genome-scale expression analysis. We conclude that the most useful knowledge <i>currently</i> going into the KB is therefore that concerning pathways. Accordingly we have accelerated our	between responde relevant s the two p to guide compour pharmace
Decisions	 work in capturing pathway-related findings from public literature and databases. Unlike browsing by humans, computational approaches can leverage the consistency, structure and quantity of information in the knowledge base. The PARIS approach has proven successful because it enables scientists to apply this prior knowledge to genome-scale expression data and quickly gain insight into biological processes relevant to the experimental context. It is expected that PARIS will be the first of many computational approaches that will be able to exploit Ingenuity's KB. The value of computational pathways analysis is increased by its broad application across the pharmaceutical pipeline: 	Lastly, it system to di and to estab mechanismi systems are feedback wi understandi by the scient Better oper be enabled, captured. A record should com and deserve organisation enough resc experiment this context judgments i resources. F will begin t
Model systems	• In the area of target discovery, analysis of model systems and their perturbations was enabled our scientists	discovery p Acknowledg
Target	to identify new pathways containing potential therapeutic targets. The relevance of these pathways was not anticipated and represents new knowledge about the biology of the model systems.	The authors as Millennium's Laura Rudolp Harshwardhar and Robert St Manchester D Peter Szolovit

pounds that are in-licensed for evelopment, pathway analysis en to be valuable for validating oring the supposed sms of these compounds. For pound, the analysis suggested a mechanism of action - one a more favourable competitive е.

g pathway analysis to our lata has provided valuable nto the molecular differences responders and noners and suggested biologically signatures for distinguishing populations. This is being used development of follow-on nds and biomarkers for ogenomics.

is vitally important in such a irect the content acquisition olish control and feedback s. If the process and software working, proper direction and ill advance the scientific ing and this will be perceptible ntific staff and management. rational decision-making will and this in turn should be

of operational decisions plement scientific knowledge es an equal place in the nal memory. There are never ources to do all the desired s. Operational knowledge in t really concerns scientific in the context of limited Future versions of *MyBiology* to capture such higher-level ffecting the entire drug ipeline within the enterprise.

gments

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