# The *BioLexicon*: a Large-Scale Domain-Specific Lexical Resource for Biomedical Text Mining

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# Outline

#### The BioLexicon

- Who
- Why
- How
- What
- Where from
  Focus on Verbs
  Representation
  How many entries
  Evaluation
  Distribution
- Conclusions

### The BioLexicon: who

Joint and collaborative work of the following teams in the framework of the European **BOOTStrep** project (FP6 - 028099)

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Boot

Strep

European Bioinformatics Institute is an Outstation of the European Molecular Biology Laboratory.

BB

MANCHESTER

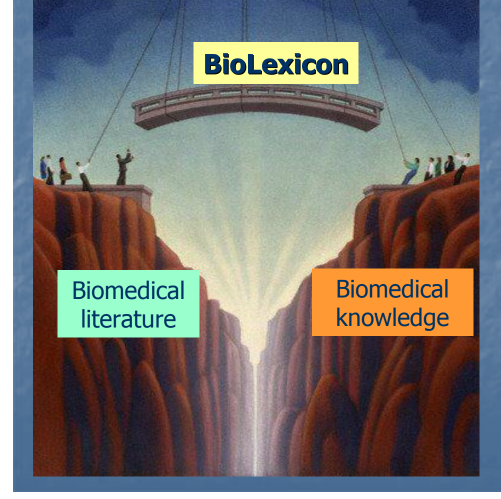
## The BioLexicon: why

Text Mining needs information about words

the lexical component still remains a major bottleneck

- TM systems in the biomedical domain must be provided with a substantial lexicon covering a realistic vocabulary and providing the kinds of linguistic information appropriate to grasp the knowledge embedded in texts
  - Biomedical term variants (orthographic, semantic, geographical, ...)
    - better information retrieval
  - Terminological verbs and their combinatorial properties (subcategorization frames and predicate-argument structure)
     better information extraction and question answering
  - Word derivations
    - to reach similar meaning expressed in different ways (e.g. activation vs activate)

## The BioLexicon as a key ingredient to **bridge the gap** between text and knowledge



To perform such a key role the BioLexicon MUST reflect the actual usage of words in biomedical texts be continuously updated with new word synonyms emerging from texts include rich linguistic information on the behavioural properties of nouns and verbs

#### The BioLexicon: how

#### General Requirements

Modularity, extensibility, conformity to standards, reusability

Biomedical Domain Specific Requirements Gene names, protein names, bio-events and participants, ...

#### Linguistic/Terminological Requirements term variants, source identifiers, acronyms, syntactic and semantic

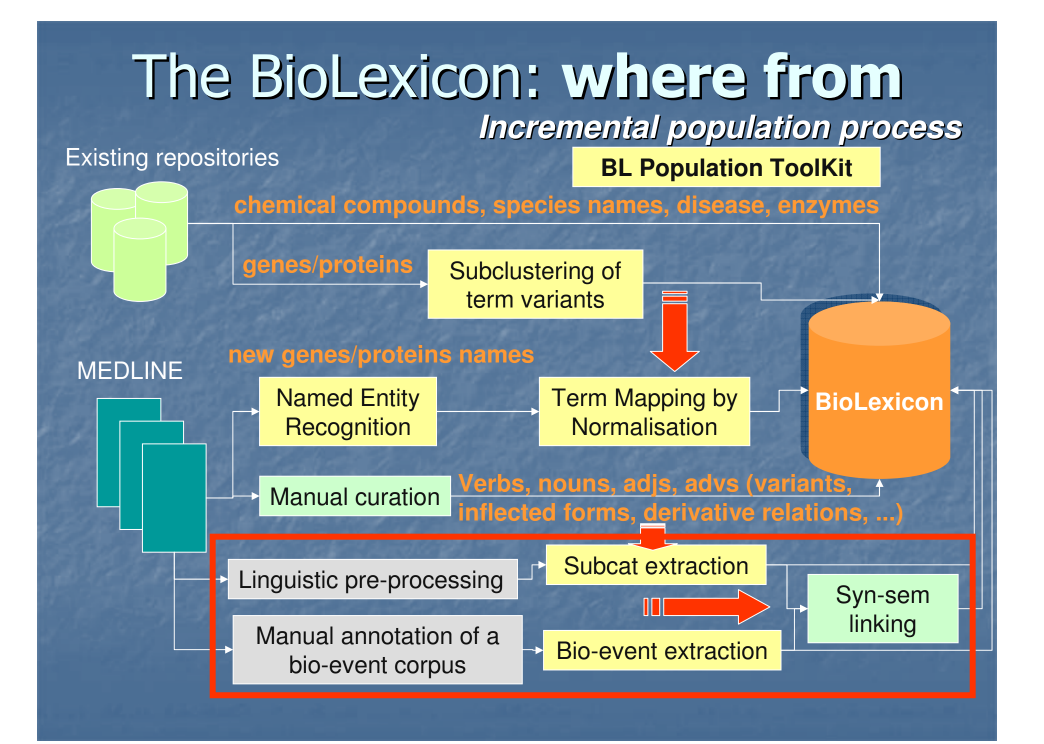
properties of terms, ...

Text Mining / Machine Learning Requirements Confidence scores for automatically extracted info (e.g. variants, subclusterizations, subcat frames, ...)

#### The BioLexicon: what

integrated lexical-terminological resource of ~2.2M lexical entries for bio-text mining with information about

- nouns, verbs, adjectives, adverbs
- both domain-specific and general language words
- populated with terms gathered from
  - available biomedical sources
  - texts (biomedical literature)
- including rich linguistic information ranging over different linguistic descriptions levels
  - e.g. derivational morphology, subcategorization patterns, predicate argument structure, syntax-semantics linking
- combining features of both terminologies and open-domain computational lexicons
- conforming to international lexical representation standards (the ISO/DIS 24613 "Lexical Mark-up Framework")
- providing links to the Gene Regulation Ontology



#### The BioLexicon: focus on verbs

Accurate TM applications focused on event extraction require lexical resources providing an exhaustive account of the semantic and syntactic combinatorial properties of lexical units conveying event information

- Several exist for the general language domain, e.g. FrameNet, VerbNet, PropBank
- Specialist domains such as *biology* require **domain**specific resources
  - use of different predicates to describe events

**E.g.** *methylate, phosphorylate* 

general language predicates may have different properties

E.g. <u>the patient presented with influenza to the</u> <u>doctor</u> vs the patient presented the doctor with influenza

#### Current biomedical lexical resources

- A number of attempts have been made to produce domainspecific extensions of general-purpose lexical semantic resources providing information on predicate-argument structure
  - BioFrameNet and PASBio
    - corpus-based
    - small-scale
  - SPECIALIST lexicon
    - extension of a large lexicon of general English
    - not corpus-driven
    - syntactic complementation patterns only
  - Creation of resources focussed on predicate-argument structure can be a major bottleneck
    - Mostly manually created by lexicographers
    - Limited coverage
    - Time-consuming to port to new domains
- Automatic or semi-automatic acquisition methods more promising and increasingly viable
  - Advances in NLP and machine learning technology
  - Availability of corpora

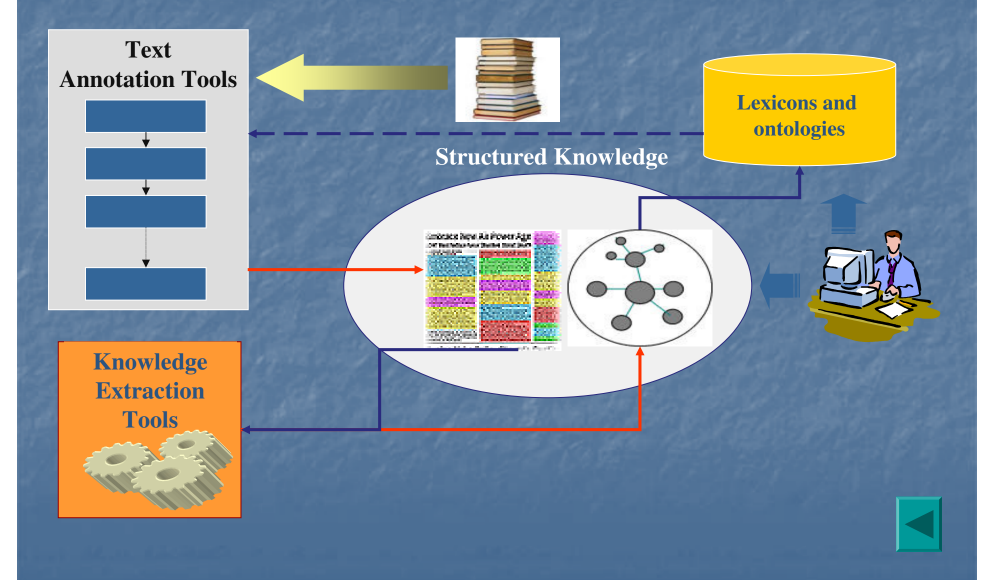
Current biomedical lexical resources: the need

To our knowlege, there is currently no large-scale domain-specific lexical resource providing predicate-argument information
 based on domain-specific corpora
 containing both syntactic and semantic information

#### The **BioLexicon** as a possible answer

The BioLexicon: our approach towards acquisition of verb information **bootstrapped** from biomedical corpora the most relevant verbs are included in the lexicon their encoded behaviour is domain-specific contains *both* syntactic and semantic information syntactic subcategorization (e.g. act ARG1#PP-as#) **semantic event frame information** (e.g. *bind* AGENT#THEME#LOC#) **explicit link** between the two (e.g. *express* AGENT>ARG1#THEME>ARG2#LOC>PP-in#) built semi-automatically by combining NLP and Machine Learning techniques syntactic frames are extracted through unsupervised learning on dependency-annotated text semantic frames are based on manual annotation of gene regulation bio-events by domain experts link between syntactic and semantic information is manually added

## From Text to Knowledge: NLP and Knowledge Extraction



# The BioLexicon: our approach to subcat extraction

 acquired through unsupervised automatic acquisition techniques from linguistically pre-processed domain corpora

the starting point: shallow or deep syntactic annotation?

- particular requirements for Subcategorization Frames (SCFs) in biomedical language
  - SCFs should also include strongly selected modifiers (such as location, manner and timing), as these are deemed to be essential for the correct interpretation of texts

average number of arguments in SCFs higher than general language

"discovery" approach to SCF acquisition based on a looser notion of SCFs, which includes typical verb modifiers in addition to strongly selected arguments

- no a priori knowledge about the set of possible SCFs
- no distinction between argument/modifier

to meet this basic requirement, SCF induction operates on a deep level of syntactic annotation

#### The BioLexicon: the **subcat** extraction process

SCFs extracted from a corpus of MEDLINE abstracts and full papers made up of 6 million word tokens
 The induction process was performed through:

- syntactic annotation of the acquisition corpus with Enju syntactic parser (v2.2, http://www-tsujii.is.s.u-tokyo.ac.jp/enju/) (adapted to biomedical texts)
- <u>extraction of the observed dependency sets</u> (ODSs) for each verbal occurrence
  - each ODS represented as a set of dependencies described in terms of relation type (e.g. ARG1, ARG2, PP-in, PP-across, that-CL, etc.)
  - order of dependencies in each ODS is normalised

#### induction of relevant SCF associated with a given verb

- for each ODS type, the conditional probability given the verb type  $\nu$  was computed
- weighted thresholds used to filter out noisy frames
  - an ODS type with an associated probability score beyond a certain threshold is selected as eligible SCF for that verb type
- each SCF has been extracted for one *normalised verb token*, i.e. the extraction process makes abstraction from the passive usages

#### Overproduction of TorR in a torT strain **resulted** in partial constitutive expression of the torA'-'lacZ fusion, **suggesting** that TorR **acts** downstream from TorT.

ROOT	ROOT	ROOT	-1	ROOT	resulted	result	VBD	7
resulted	result	VBD	7	verb_arg1	Overproduction	overproduction	NN	0
a	a	DT	4	det_arg1	strain	strain	NN	6
the	the	DT	13	det_arg1	fusion	fusion	NN	15
torT	tort	NN	5	noun_arg1	strain	strain	NN	6
partial	partial	JJ	9	adj_arg1	expression	expression	NN	11
torA'-'lacZ	tora'-'lacz	NN	14	noun_arg1	fusion	fusion	NN	15
constitutive	constitutive	JJ	10	adj_arg1	expression	expression	NN	11
acts	act	VBZ	20	verb_arg1	TorR	torr	NN	19
of	of	IN	1	prep_arg12	Overproduction	overproduction	NN	0

#### act ARG1=torr@NN 0 0 0 0 PP-from=tort@NNP 0 0 0 0 0 0 0 0 0 0 VBZ 0 result ARG1=overproduction@NN 0 0 0 0 PP-in=expression@NN 0 0 0 0 MOD=suggest@VBG 0 0 0 0 suggest ARG1=UNKNOWN@UNKNOWN 0 0 0 0 0 0 0 0 0 0 0 0 0 that-CL=act@VBZ 0 0 VBG 0

	of	of	IN	12	prep_arg12	expression	expression	NN	11	
	of	of	IN	12	prep_arg12	fusion	fusion	NN	15	
	that	that	IN	18	comp_arg1	acts	act	VBZ	20	
	downstream	downstream	RB	21	adj_arg1	acts	act	VBZ	20	
100	in	in	IN	8	prep_arg12	resulted	result	VBD	7	
	in	in	IN	8	prep_arg12	expression	expression	NN	11	
	from	from	IN	22	prep_arg12	acts	act	VBZ	20	
	from	from	IN	22	prep_arg12	TorT	tort	NNP	23	
	suggesting	suggest	VBG	17	verb_mod_arg12	resulted	result	VBD	7	
	suggesting	suggest	VBG	17	verb_mod_arg12	UNKNOWN	UNKNOWN	UNKNOWN	-1	
	suggesting	suggest	VBG	17	verb_mod_arg12	acts	act	VBZ	20	

# The BioLexicon: **subcat** extraction results

#### 136 different induced SCFs

• vs SPECIALIST Lexicon: very limited number of complementation patterns

Verb	SCF	P(subcat v)	Pass
abolish	ARG1#ARG2#	0.8669767	0.1437768
abolish	ARG1#ARG2#MOD@VBG#	0.0390697	0.1904761
abolish	ARG1#ARG2#PP-in#	0.0939534	0.7029702
accumulate	ARG1#ARG2#	0.2940677	0.0403458
accumulate	ARG1#	0.4627118	0
accumulate	ARG1#ARG2#PP-in#	0.1084745	0.140625
accumulate	ARG1#PP-in#	0.1347457	0

# The BioLexicon: acquired SCFs and strongly selected modifiers

- many of the strongly selected modifiers spread over different SCFs
  - radically underestimated role
- SCFs complemented with information about individual dependencies of verbs
  - typical verbal dependencies, corresponding to either arguments or strongly selected modifiers, detected through the II association score
  - 44 induced dependency types

v	SCF	v_freq		SCF_	_freq	p(\$	SCFIv)	<b>•</b>	assive ages		
methylate	ARG1#ARG2#	422			294		0.6967		0.1258		
methylate	ARG1#ARG2#PP-in#		V	7	DE	Р	all_ dep	dep_ freq	p(deplv)	11	% passive usages
methylate	ARG1#ARG2#PP-at#		methy	ylate	ARG	2#	1406	410	0.2916	778.5146	0.25
			methy	ylate	PP-at	#	1406	29	0.0206	57.9113	0.31
		methy	ylate	PP-in	#	1406	45	0.0320	18.2749	0.60	

# The BioLexicon: an example of stored subcat information for the verb *acquire*

v	SCF	p(SCF v)	% pass
acquire	ARG1#ARG2#	0.5461	0.1284
acquire	ARG1#ARG2#PP-in#	0.0886	0.0833
acquire	ARG1#ARG2#PP-from#	0.0406	0.1818
acquire	ARG1#ARG2#PP-by#	0.0406	0.0000
acquire	ARG1#ARG2#PP-during#	0.0295	0.3750

v	DEP	ll	% pass	
acquire	ARG2#	579.96392	0.1512915	
acquire	WH-when#	25.703417	0.1	
acquire	PP-from#	22.716082	0.3333333	
acquire	PP-by#	13.626654	0	
acquire	PP-in#	13.416025	0.1666667	

Prepositionbased parsing

Full parsing

# The BioLexicon: contained verb subcategorization information

complementarity between Verb-Dep and Verb-SCF associations
 both information types included into the BioLexicon

subcat frame information acquired for 759 different verbs, corresponding to 658 different base forms
 e.g. the occurrences of *colocalize, colocalise, co-localize* and *co-localise* recorded under *colocalize*

the BioLexicon was augmented with:

1410 Verb-SCF associations, involving 136 different subcat frame types

3040 Verb-Dep associations, involving 44 slot types

#### The BioLexicon: bio-event frames

Extracted from a corpus of 677 MEDLINE abstracts manually annotated by biologists

- semantic parsers not mature enough to provide the starting point (as opposed to dependency parsers)
- manual semantic annotation carried out on top of shallow syntactic annotation ("chunking")
  - consistency of marked text spans helped by annotating syntactic chunks

[NP The narL gene product ] [VP activates ] [NP the nitrate reductase operon ] [PP in ] [NP Escherichia coli ]

# Bio-Event annotated corpus: incremental annotation approach

Domain specific annotation

Domain independent annotation Bio-Event Linguistic Annotation

Syntactically "chunked" text

Morpho-syntactically tagged text

Tokenised text

Raw domain corpora



**KNOWLEDGE** 

### The BioLexicon: **bio-event frames** annotation

Annotation consisted of:

- Identifying relevant gene regulation events centred on verbs and nominalised verbs (e.g. expression)
- Finding all semantic arguments of an identified event
  - Within-sentence annotation
- Assigning a semantic role to each argument
- Assigning named entity types to semantic arguments (where appropriate)
  - A hierarchy of NEs, specially tuned to gene regulation, was created
  - Organised into five entity-specific super-classes

Thompson P., Cotter P., McNaught J., Ananiadou S., Montemagni S., Trabucco A., Venturi G. 2008. Building a Bio-Event Annotated Corpus for the Acquisition of Semantic Frames from Biomedical Corpora. In Proceedings of LREC 2008

#### Bio-Event annotated corpus: semantic roles

- Aim for a set of **verb-specific event frames** Use of frame-independent semantic roles
  - annotation of all sublanguage semantic arguments, using a set of domain-specific and domainindependent roles
  - The proposed set of 12 event-independent semantic roles includes:
    - two domain-specific semantic roles, i.e. CONDITION and MANNER;
    - semantic roles particularly important for the precise definition of complex biological relations, even though not necessarily specific to the field, i.e. LOCATION and TEMPORAL;
    - semantic roles widely traceable across all domains

### Bio-Event annotated corpus: list of semantic roles (1)

AGENT	Drives/instigates event	<b>The narL gene product</b> <i>activates</i> the nitrate reductase operon		
THEME	<ul> <li>a) Affected</li> <li>by/results from event</li> <li>b) Subject of events</li> <li>describing states</li> </ul>	<b>recA protein</b> was <i>induced</i> by UV radiation <b>The FNR protein</b> <i>resembles</i> CRP		
MANNER	Method/way in which event is carried out	cpxA gene <i>increases</i> the levels of csgA transcription by <b>dephosphorylation</b> of CpxR		
INSTRUMENT	Used to carry out event	We have <b>isolated</b> a strain with the aid of <b>the Casadaban Mud phage</b>		
LOCATION	Where <i>complete</i> event takes place	Phosphorylation of OmpR <i>modulates</i> expression of the ompF and ompC genes <b>in Escherichia coli</b>		
SOURCE	Start point of event	A transducing lambda phage was <b>isolated</b> from <b>a strain</b> harboring a glpD''lacZ fusion		
DESTINATION	End point of event	Transcription of gnfT is activated by <i>binding</i> of the cyclic AMP (cAMP)-cAMP receptor protein (CRP) complex to <b>a CRP binding site</b>		

### Bio-Event annotated corpus: list of semantic roles (2)

TEMPORAL	Situates event w.r.t another event	The Alp protease activity is <i>detected</i> in cells <b>after</b> <b>introduction</b> of plasmids carrying the alpA gene
CONDITION	Environmental conditions/changes in conditions	Strains carrying a mutation in the crp structural gene fail to <i>repress</i> ODC and ADC activities in response to <b>increased cAMP</b>
RATE	Change of level or rate	marR mutations <i>elevated</i> inaA expression by <b>10- to</b> <b>20-fold</b> over that of the wild-type.
DESCRIPTIVE- AGENT	Descriptive information about AGENT	It is likely that HyfR <i>acts</i> as <b>a formate-dependent regulator</b> of the hyf operon
DESCRIPTIVE- THEME	Descriptive information about THEME	The FNR protein <i>resembles</i> CRP.
PURPOSE	Purpose/reason for the event occurring	The fusion strains were <b>used to study</b> the regulation of the cysB gene by assaying the fused lacZ gene product

# Named Entity Superclasses

NE class	Definition
DNA	Entities chiefly composed of nucleic acids and their structural or positional references. This includes the physical structure of all DNA-based entities and the functional roles associated with regions thereof.
PROTEIN	Entities chiefly composed of amino acids and their positional references. This includes the physical structure and functional roles associated with each type.
EXPERIMENTAL	Both physical and methodological entities, either used, consumed or required for a reaction to take place.
ORGANISMS	Entities representing individuals or collections of living things and their component parts.
PROCESSES	A set of <i>event</i> classes used to label biological processes described in text.

#### **Event Annotation example**

A promoter has been identified that directs AGENT Verb

PROCESS relA gene transcription towards THEME

DNA the pyrG gene DESTINATION

in a counterclockwise direction MANNER

DNA on the E. Coli chromosome LOCATION

#### The extraction process of Event Frame

Input sentence: Agent Theme transfer operon expresses F-like plasmids DNA DNA Syntactic analysis:

[Agent:NN:DNA] [Verb:VBZ:express] [Theme:NN:DNA]

**Extracted event frame:** 

express(Agent=>DNA, Theme=>DNA)

#### The BioLexicon: acquired **bio-event** frames

đ	verb	Bio-event frames
	activate	Agent#Theme#
		Agent#Theme#Condition#
		Agent#Theme#Location#
		Agent#Theme#Manner#
		Agent#Theme#Source#
		Theme#
		Theme#Condition#

200	verb	Bio-event frames with NE types
	activate	Agent-DNA#Theme-DNA#
		Agent-Organisms#Theme-Protein#
		Agent-Protein#Theme-DNA#

## The BioLexicon: Syntax-Semantics Linking (1)

#### The starting point:

- acquired subcategorization frames
- verbal bio-event frames based on corpus annotation
- acquired using different techniques and corpora of different size
- Linking concerned 168 verbs for which both syntactic and semantic information was available
- Linking process carried out manually by a linguist
  - Different information types were taken into account, i.e.
    - literature regarding hierarchies of semantic roles and grammatical functions
      - given a thematic role hierarchy (agent>theme ...) and a syntactic functions hierarchy (subject>object ...), the mapping usually proceeds from left to right
    - a list of 'prototypic' syntactic realisations of semantic arguments
    - exploitation of general language repositories of semantic frames containing both syntactic and semantic information (as possible benchmarks)

### The BioLexicon: Syntax-Semantics Linking (2)

Linking process resulted in 668 linked frames

Different types of mapping were performed:

- full mapping (239 frames)
  - arity of the subcategorization and bio-event frames is the same (ISO)

#### partial mapping

 semantic frame contains more slots than corresponding subcategorization frame (123 frames) (AUG)

e.g. AGENT>ARG1#THEME>ARG2#LOCATION>PP-in#CONDITION>0

2) subcategorized slots do not have a semantic counterpart in the corresponding bio-event frame (166 frames) (RED)

e.g. 0>ARG1#THEME>ARG2#DESTINATION>PP-into

3) a combination of cases 1) and 2) above (140 frames)

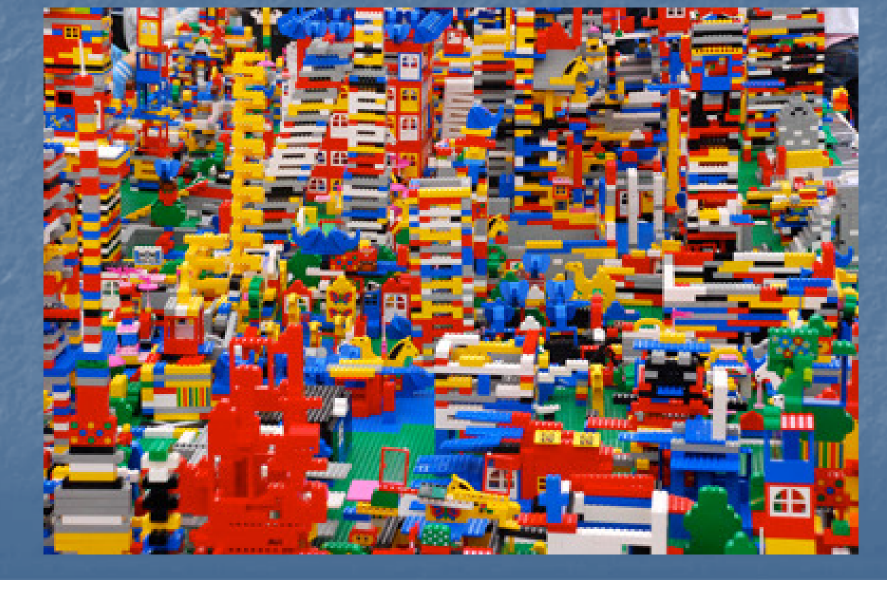
e.g. 0>ARG1#THEME>ARG2#SOURCE>PP-from#CONDITION>0

The BioLexicon:										
syntax-semantics linking										
3638	Theme	ARG2	0	ARG1			RED			
	Agent	ARG1	Theme	ARG2	Conditio n	PP-in	ISO			
	Agent	ARG1	Theme	ARG2	Manner	PP-by	ISO			
	Agent	ARG1	Theme	ARG2	Location	PP-in	ISO			
activate	Agent	ARG1	Theme	ARG2	Source	0	AUG			
1.49	Theme	ARG2	Condition	PP-in	0	ARG1	RED			
1	Agent	ARG1	Theme	ARG2			ISO			
	Theme	ARG1					ISO			
	Agent	ARG1	Theme	ARG2	Manner	PP-in	ISO			

Useful information for mixed syntax-semantics approaches

G. Venturi, S. Montemagni, S. Marchi, Y. Sasaki, P. Thompson, J. McNaught, S. Ananiadou, 2009, "Bootstrapping a Verb Lexicon for Biomedical Information Extraction", in Proceedings of the CICLing-2009 conference, Mexico.

# From bricks of biolexical knowledge to a computational BioLexicon



### The BioLexicon: representation model

The BL model is conformant to ISO-LMF (ISO 24613:2008)

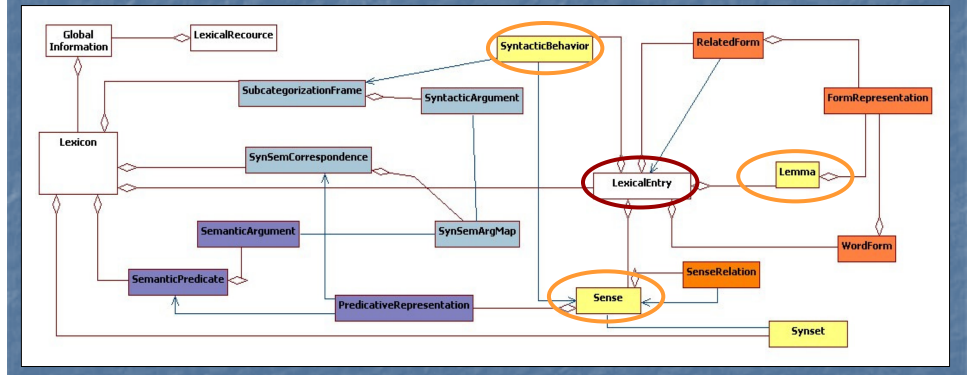
- high-level objects: the meta-model, i.e. a set of independent lexical objects with relations among them
- Iow-level objects: a set of Data Categories, i.e. linguistic constants in the form of attribute-value pairs (either drawn from the ISO-12620 or defined for the special domain)

#### XML DTD for the entire lexicon

- The implementation consists of a flexible, extensible relational MySQL database
- Automatic population procedures relying on a dedicated input data structure, the BioLexicon XML Interchange Format (XIF)
- An XML LMF conformant export function is available

#### The BioLexicon: representation model

#### The BioLexicon Model: High-level objects, lexical objects

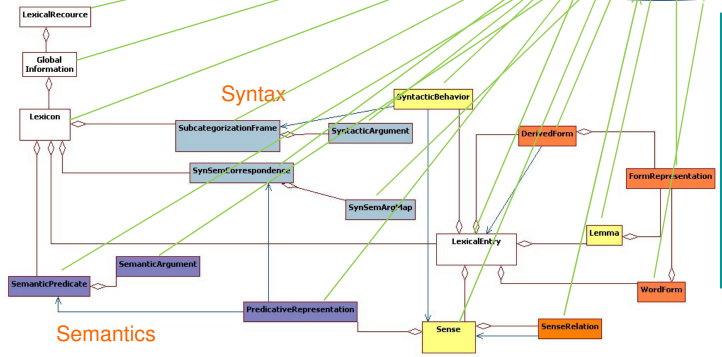


## The BioLexicon: representation model

#### BioLexicon Model: Data Categories

DC selection

e.g. <feat att="POS" val="VVZ"> <feat att="ConfScore" val="0.9"> <feat att="source" val="UNIPRO"



Partially
drawn by the
ISO Data
Category
Registry
Partially
integrated
with domain-specific DCs

# The BioLexicon: the starting point

Semantic	Resources		Semantic type	Resources
туре			NucleicAcid	Sequence
Cell	Cell ontology		Region	Ontology :Region
Cell Component	Gene Ontology GO:0005575 cellular component		Operon	RegulonDB, ODB (Operon DataBase)
Chemical	CHEBI, IMR:0000947		Organism	NCBI Species
	chemical		Transcription	Sequence
Disease	OMIN		Factor-BindingSite	Ontology
Enzyme	Enzyme commission		Protein	BioThesaurus
Gene	BioThesaurus		Protein Complex	Corum database
Ligand	IMR - INOH Protein name/family name ontology		Protein Domain	InterPro
			Transcription Regulator	RegulonDB, TransFac, Gene
Nuclear Receptor	5			Ontology Annotation

## The BioLexicon by numbers

#### Entries and variants by semantic type

Sem. Type	# Entries	# Variants
Gene/Prot	1640608	1408312
Gene/Prot (synsets)	358335	936126
Organisms	482992	182610
Enzymes	4016	4164
Protein Domains	16940	15412
Protein Compl.	2104	418
Chemicals	19637	77475
Diseases	19457	11314
Molecular Roles	8850	29831
Cell	842	512
Trans. Factors	160	129
Operons	2672	368
Sequences	1431	741

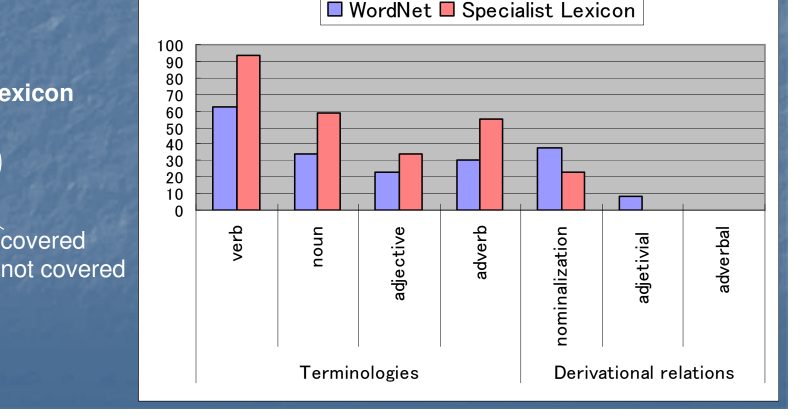
En	tries by spee	part of ch		
POS		# entries		
Nouns		2231574		
Adjectives		3428	Verbs	
Verbs		1154		
Adverbs		550	domain- specific	general
	Sec.		658	496
	inflected forms		15274	
	absorb -> absorber/	ntries (e.g. > absorption/N, /N, absorbing/J, le/J, absorbent/J, tly/R)	2764	-
30	verb-SCF associations		3040	-
verb-SLOT associations		1710	-	
bio-event frames			856	-
syntax-semantics mappings (concerned with 168 verbs)			668	-

### The BioLexicon: intrinsic evaluation

Comparison with two existing large-scale dictionaries WordNet: General English Thesaurus NLM Specialist Lexicon: Biomedical Lexicon

Coverage evaluation

covered





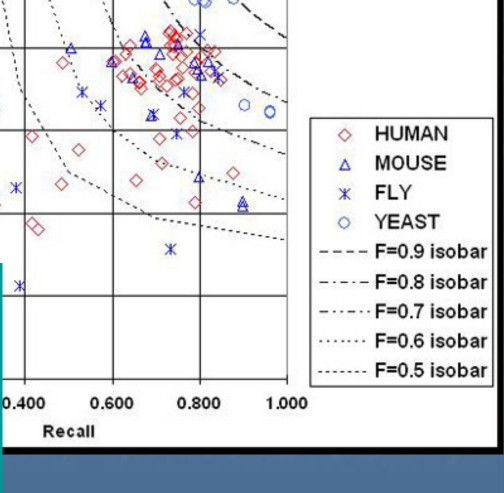
**BioLexicon** 

#### BioLexicon in BioCreAtIve II, GN

**Applied methods:** 

- Abner for gene identification
- Statistical features (BioLexicon) for filtering of non-relevant terms
- Classification on BioCreAtlve II data
- Only human concept ids
- $\Rightarrow$  Baseline results
- $\Rightarrow$  Highly reproducible
- $\Rightarrow$  Available as Whatizit module

(BioLexHuman)



# The BioLexicon: extrinsic evaluation

Task-based evaluation (still ongoing)

Task	Data	Тоој
IR	TREC Genomics Track 2007	<ul><li>BLTagger</li><li>NeMine (NER)</li></ul>
IE	UoM Gene Regulation Corpus	<ul> <li>BLTagger</li> <li>NeMine (NER)</li> <li>Enju with the BL</li> </ul>

NeMine (http://text0.mib.man.ac.uk/~sasaki/bootstrep/nemine.html)

Y. Sasaki, P. Thompson, J. McNaught, S. Ananiadou, 2009, "Three BIONLP tools powered by the BioLexicon", in Proceedings of EACL 2009.

# BioLexicon distribution

- The BioLexicon (MySQL version) is distributed through the European Language Resources Association (ELRA)
  - <u>http://www.elra.info</u> or <u>http://www.elda.org</u>

#### Benefits

- Servicing of bug reporting through ELRA
- Organisational embedding into other lexical resources
- Long-term availability
- Support to European language infrastructures
- Different licence types for
  - Commercial use
  - Research use by commercial organisations
  - Research use by academic organisations



## Conclusions

A unique resource amongst large-scale computational lexicons within the biomedical domain in terms of coverage and typology of contained information

Including both domain-specific and general language words

Rich linguistic information ranging over different linguistic descriptions levels Designed to meet **bio-Text** Mining requirements

> Populated with info from available **biomedical** resources and texts

Semi-automatically populated from corpora: Population toolkit available

Conformant to international lexical representation standards



