

# Potential of biological control based on published research. 1. Protection against plant pathogens of selected crops

Philippe C. Nicot, Marc Bardin, Claude C. Alabouvette, Jürgen Köhl, Michelina Ruocco

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WPRS International Organisation for Biological and Integrated Control of Noxious Animals and Plants: West Palaearctic Regional Section

**SROP** Organisation Internationale de Lutte Biologique et Integrée contre les Animaux et les Plantes Nuisibles: Section Régionale Ouest Paléarctique

# Classical and augmentative biological control against diseases and pests:

# critical status analysis and review of factors influencing their success



Edited by Philippe C. Nicot

2011





The content of the contributions is the responsibility of the authors

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# Preface

One of the Research Activities (RA 4.3) of the European Network for Durable Exploitation of crop protection strategies (ENDURE<sup>\*</sup>) has brought together representatives of industry and scientists from several European countries with experience ranging from fundamental biology to applied field work on biological control against pests and diseases. The unique diversity of expertise and concerns allowed the group to set up very complementary approaches to tackle the issue of the factors of success of biocontrol.

The initial part of the work accomplished by this group consisted in a thorough review of scientific literature published on all types of biological control. Although it had to be focused on selected key European crops and their major pests and pathogens, this review is unique in the scope of the topics it covered and in the comprehensive inventories it allowed to gather on the potential of biocontrol and factors of success at field level.

In parallel with identifying knowledge gaps and key factors from published research, information was gathered on aspects linked to the production and commercialization of biocontrol agents.

These results, complemented by the views of experts in the field of biocontrol consulted at the occasion of meetings of IOBC-wprs, allowed the identification of majors gaps in knowledge and bottlenecks for the successful deployment of biocontrol and lead to the proposition of key issues for future work by the research community, the field of development and prospects for technological improvement by industry.

Avignon, June 2011 Philippe C. Nicot

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# For Chapter 1

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# **Chapter 1**

# Potential of biological control based on published research. **1.** Protection against plant pathogens of selected crops

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#### **Evolution of the scientific literature**

The scientific literature published between 1973 and 2008 comprises a wealth of studies on biological control against diseases and pests of agricultural crops. A survey of the CAB Abstracts® database shows a steady increase in the yearly number of these publications from 20 in 1973 to over 700 per year since 2004 (Figure 1).

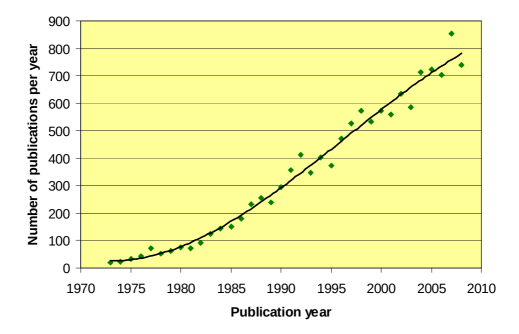


Figure 1: Evolution of the yearly number of publications dedicated to biological control of plant diseases based on a survey of the CAB Abstracts® database.

This survey was further refined by entering keywords describing some of the major plant pathogens/diseases of cultivated crops in Europe, alone or cross-referenced with keywords indicating biocontrol. Among studies published in the period between 1973 and 2008 on these plant pathogens and pests, the percentage dedicated to biological control was substantial, but unequally distributed (Table 1). It was notably higher for studies on soil-borne (9.5%  $\pm$  1.6% as average  $\pm$  standard error) than for those on air-borne diseases (2.8%  $\pm$  0.7%).

Table 1: Scientific papers published between		on biological	control against major
plant diseases (from CAB Abstracts <sup>®</sup> d	latabase).		

Disease or plant pathogen	Total number of references	References on biological control		
			%	
Soil-borne:				
Fusarium	34 818	1 925	5.5	
Rhizoctonia	10 744	1 278	11.9	
Verticillium	7 585	592	7.8	
Pythium	5 772	821	14.2	
Sclerotinia	5 545	456	8.2	
Air-borne:				
rusts	29 505	360	1.2	
powdery mildews	18 026	251	1.4	
Alternaria	12 766	415	3.3	
anthracnose	12 390	351	2.8	
Botrytis	9 295	705	7.5	
downy mildews	8 456	80	1.0	
Phytophthora infestans	5 303	61	1.1	
Monilia rot	1 861	81	4.3	
Venturia	3 870	104	2.7	

# Inventory of potential biocontrol agents (microbials, botanicals, other natural compounds)

The scientific literature described above was further examined to identify biocontrol compounds and microbial species reported to have a successful effect. Due to the great abundance of references, it was not possible to examine the complete body of literature. The study was thus focused on several key diseases selected for their general importance on cultivated crops, and in particular on those crops studied in the case studies of the European Network for Durable Exploitation of crop protection strategies (ENDURE<sup>\*</sup>).

# Methodology

Three steps were followed. The <u>first step</u> consisted in collecting the appropriate literature references for the selected key diseases/plant pathogens to be targeted by the study. The references were extracted from the CAB Abstracts<sup>®</sup> database and downloaded to separate files using version X1 of EndNote (one file for each target group). The files were then distributed among the contributors of this task for detailed analysis.

In the <u>second step</u>, every reference was examined and we recorded for each:

- the types of biocontrol agents (Microbial, Botanical or Other compounds) under study and their Latin name (for living organisms and plant extracts) or chemical name
- the Latin name of the specifically targeted pathogens,
- the crop species (unless tests were carried out exclusively in vitro),
- the outcome of efficacy tests.

Two types of efficacy tests were distinguished: Controlled environment tests (including tests on plants and *in vitro* tests), and field trials. The outcome of a test was rated (+) if significant effect was reported, (0) if no significant efficacy was shown and (-) if the biocontrol agent stimulated disease development.

<sup>\*</sup> EU FR6 project 031499, funded in part by the European Commission

To allow for the analysis of a large number of references, the abstracts were examined for the presence of the relevant data. The complete publications were acquired and examined only when the abstracts were not sufficiently precise.

The data were collected in separate tables for each type of key target pest. For each table, they were sorted (in decreasing order of priority) according to the type and name of the biocontrol agents, the specifically targeted pest, and the outcome of efficacy tests.

In the <u>third step</u>, synthetic summary tables were constructed to quantify the number of different biocontrol compounds and microbial species and strains reported to have successful effect against each type of key pathogen/disease or pest target.

#### Results

A total number of 1791 references were examined for key airborne diseases including powdery mildews, rusts, downy mildews (+ late blight of Potato/Tomato) and *Botrytis* and *Monilia* rots, together with soilborne diseases caused by *Fusarium oxysporum* (Table 2). Based on the examination of these references, successful effect in controlled conditions was achieved for all targets under study with a variety of species and compounds (Appendices 1 to 6, Table 3).

Target disease / plant pathogen			Period of publication examined	
Botrytis	OR, FV, GR* (postharvest)	880	1998-2008	
Powdery mildews	all	166	1998-2008	
Rusts	AC, FV, OR	154	1973-2008	
Downy mildews + Phytophthora infestans	FV, GR, PO, TO	349	1973-2008	
Monilinia rot	OR	194	1973-2008	
Fusarium oxysporum	FV. TO	48	2007-2009	

 Table 2:
 Numbers of references on biocontrol examined per group of disease/plant pathogen.

\*AC: Arable Crops; FV: Field Vegetables; GR: Grapes; OR: orchard; PO: Potato; TO: Tomato

Concerning **airborne diseases and pathogens**, the largest number of reported successes was achieved with microbials, but there is a growing body of literature on plant and microbial extracts, as well as other types of substances (Table 3). On average, reports of success were far more numerous for experiments in controlled conditions (*in vitro* or *in planta*) than for field trials.

Very contrasted situations were also observed depending on the type of target disease/pathogen, with rare reports on the biocontrol of rusts and mildews compared to *Botrytis*, despite the fact that the literature was examined over a 35 year period for the former diseases and only over the last 10 years for the latter.

In total in this review, 157 species of micro-organisms have been reported for significant biocontrol activity. They belong to 36 genera of fungi or oomycetes, 13 of yeasts and 25 of bacteria. Among them, 29 species of fungi/oomycetes and 18 bacteria were reported as successful in the field against at least one of the five key airborne diseases included in this review (Table 4).

# Nicot *et al*.

IIIOIIIatic		0			ented in Appe	
Target plant pathogen /	Botanicals		Microl	pials <sup>y</sup>	Others <sup>z</sup>	
disease	laboratory tests <sup>x</sup>	field trials	laboratory tests <sup>x</sup>	field trials	laboratory tests <sup>x</sup>	field trials
Botrytis						
in vitro	26	-	31 b, 21 f	-	7	-
legumes	4	2	10 b, 12 f	3b, 9 f	0	0
protected vegetables	0	1	22 b, 24 f	8 b, 9 f	5	1
strawberry	0	0	14 b, 21 f	2 b, 13 f	7	1
field vegetables	0	0	5 b, 15 f	2 f	0	0
grapes	1	3	5 b, 27 f	5 b, 13 f	0	1
pome/stone fruits	1	0	12 b, 35 f	2 b, 6 f	4	0
others	3	0	15 b, 25 f	6 b, 6 f	0	0
Powdery mildews						
Grape	1	1	4b; 10f	2b; 12f	3	2
Arable crops	1	0	2b;9f	1b	5	0
Strawberry	0	0	4b; 6f	0	0	0
Cucurbitaceae	4	0	14b; 22f	4b; 9f	9	1
Pome/stone fruits	0	0	3f	1f	0	0
Pepper	1	0	4f	0	1	0
Tomato	5	0	4b; 5f	1f; 1b	0	0
Various	2	0	2b; 10f	1b; 1f	5	0
Rusts						
arable crops	0	0	5 b, 6 f	2 b	2	0
others	0	0	8 b, 13 f	0	1	0
Downy mildews + late						
blight						
grapes	2	4	2 f	3 b, 2 f	2	3
field vegetables	0	0	4	0	4	6
potato	9	1	8 b, 10 f	5 b, 4 f	3	1
tomato	2	1	5 b, 5 f	4 b	12	1
Monilia rot						
in vitro	0	-	8	-	1	-
pome fruit	0	0	7	0	0	0
stone fruit	0	1	23b, 19	7b, 7f	2	2
others	0	0	1b	2b, 1f	0	0

Table 3:Numbers of different biocontrol compounds and microbial species reported as having<br/>successful effect against key airborne pathogens/diseases of selected crops. Detailed<br/>information and associated bibliographic references are presented in Appendices 1 to 5

<sup>x</sup> tests conducted *in vitro* and/or *in planta* in controlled conditions

<sup>y</sup> b: bacteria; f: fungi / oomycetes / yeasts

<sup>z</sup> including culture filtrates and extracts from microorganisms

Table 4:Microbial species of fungi/oomycetes, yeasts and bacteria reported to have a significant<br/>effect against five main types of airborne diseases or pathogens in laboratory conditions<br/>or in the field (yellow highlight). Bibliographic references are presented in Appendices 1<br/>to 5.

	Target disease / pathogen					
Microbial species	Botrytis	Powdery mildew	Rust	Downy mildew, late blight	Monillia rot	
Acremonium spp.			others			
Acremonium alternatum		cereals, <mark>protected</mark> vegetables				
A. cephalosporium	<mark>grapes</mark>					
A. obclavatum			others			
Alternaria spp.	grapes	cereals				
A. alternata			others	grapes		
Ampelomyces quisqualis		fruits, grapes, strawberry, protected vegetables, others,				
Aspergillus spp.		,	others	tomato		
A. flavus				others		
Beauveria sp	protected vegetables					
Botrytis cinerea non- aggressive strains	legumes					
Chaetomium cochlioides	grapes					
C. globosum	legumes					
Cladosporium spp.	flowers		others			
C. chlorocephalum				others		
C. cladosporioides	flowers, legumes	others				
C. oxysporum	flowers	others	others			
C. tenuissimum		strawberry	field vegetables, others			
Clonostachys rosea	flowers, legumes, others, strawberries, field vegetables, protected vegetables,					
Coniothyrium spp.	grapes					
C. minitans	field vegetables					
Cylindrocladium	others					
Drechslera hawaiinensis		others				
Epicoccum sp	flowers, grapes, field vegetables					
E. nigrum	legumes, strawberries				plum, <mark>peach</mark>	
E. purpurascens					apple, cherry	
Filobasidium floriforme	fruits				· · · ·	
Fusarium spp.	flowers		others			
F. acuminatum		cereals				
F. chlamydosporum			others			
F. oxysporum		cereals		tomato		
F. proliferatum				grapes		
Galactomyces geotrichum	fruits					

Gliocladium spp.	<mark>grapes</mark> , protected vegetables <mark>, others</mark>				
G. catenulatum	protected vegetables, legumes				
G. roseum	flowers, <mark>grapes</mark> , legumes, <mark>others</mark>	others			blueberry
G. virens	strawberries, field vegetables			<mark>potato</mark> , <mark>others</mark>	
G. viride	protected vegetables				
Lecanicillium spp.		protected vegetables			
L. longisporum		protected vegetables			
Meira geulakonigii		protected vegetables			
Microdochium dimerum	protected vegetables, protected vegetables				
Microsphaeropsis ochracea	<mark>field vegetables</mark>				
Muscodor albus	fruits, grapes				peach
Paecilomyces farinosus		cereals			
P. fumorosoroseus		protected vegetables			
Penicillium spp.	fruits, field vegetables		others	potato, tomato	
P. aurantiogriseum	legumes			potato	
P. brevicompactum	legumes				
P. frequentans	C 11				plum, <mark>peach</mark>
P. griseofulvum	legumes, field vegetables				
P. purpurogenum					peach
P. viridicatum				potato	
Phytophthora cryptogea				potato	
Pseudozyma floculosa		<mark>grapes</mark> , protected <mark>vegetables</mark> ,			
Pythium oligandrum	protected vegetables				
P. paroecandrum	grapes				
P. periplocum	grapes				
Rhizoctonia	flowers			potato	
Scytalidium	grapes				
S. uredinicola			others		
Sordaria fimicola					apple
Tilletiopsis spp.		grapes			
T. minor	~ ~ ~	others			
Trichoderma spp.	flowers, <mark>grapes</mark> , legumes, strawberries, <b>protected vegetables</b> , others			potato	
T. asperellum	strawberries				
T. atroviride	legumes, strawberries				peach
T. hamatum	flowers, <mark>legumes</mark>				
T. harzianum	flowers, grapes, legumes, strawberries, field vegetables, protected vegetables, others	others, strawberry, <mark>protected</mark> <mark>vegetables</mark> ,	others	grapes, potato, tomato, field vegetables, others	cherry, peach
T. inhamatum	flowers				
T. koningii	strawberries, field vegetables				peach
T. lignorum				others	

T. longibrachiatum	strawberries				
T. polysporum	strawberries				apple
T. taxi	protected vegetables				
T. virens	grapes				
T. viride	fruits, grapes, <mark>legumes,</mark> strawberries, field vegetables, <mark>others</mark>	others	others	<mark>potato</mark> , others	peach
Trichothecium	grapes				
T. roseum	grapes, <mark>legumes</mark>				
Ulocladium sp.	grapes, field vegetables				
U. atrum	flowers, grapes, strawberries, field vegetables, protected vegetables				
U. oudemansii	grapes and a second s				
Ustilago maydis	protected vegetables				
Verticillium	grapes		legumes		
V. chlamydosporium			cereals		
V. lecanii	strawberries	cereals, protected vegetables, others	legumes, others		

#### B. Yeasts

	Target disease / pathogen							
Microbial species	Botrytis	Powdery mildew	Rust	Downy mildew, late blight	Monillia rot			
Aureobasidium pullulans	fruits, grapes, strawberries, protected vegetables				apple, cherry			
Candida spp.				tomato	peach			
C. butyri	fruits							
C. famata	fruits							
C. fructus	strawberries							
C. glabrata	strawberries							
C. guilliermondii	grapes, protected vegetables				cherry			
C. melibiosica	fruits							
C. oleophila	fruits, grapes, strawberries, protected vegetables							
C. parapsilosis	fruits							
C. pelliculosa	protected vegetables							
C. pulcherrima	fruits, strawberries							
C. reukaufii	strawberries							
C. saitoana	fruits							
C. sake	fruits							
C. tenuis	fruits							
Cryptococcus albidus	fruits, strawberries, protected vegetables							
C. humicola	fruits							
C. infirmo-miniatus	fruits				cherry			
C. laurentii	fruits, strawberries, protected vegetables				cherry, peach			
Debaryomyces hansenii	fruits, grapes				cherry, peach			
Hanseniaspora uvarum	grapes							
Kloeckera spp	grapes							
K. apiculata	fruits				cherry, peach			
Metschnikowia fructicola	fruits, grapes, strawberries							
M. pulcherrima	fruits				apple, apricot			
Pichia anomala	grapes, fruits							

P. guilermondii	fruits, strawberries, protected vegetables			
P. membranaefaciens	grapes			peach
P. onychis	field vegetables			
P. stipitis	fruits			
Rhodosporidium diobovatum	protected vegetables			
R. toruloides	fruits			
Rhodotorula				peach
R. glutinis	flowers, fruits, strawberries, protected vegetables	field vegetables,		
R. graminis	flowers			
R. mucilaginosa	flowers			
R. rubra	protected vegetables			
Saccharomyces cerevisiae	fruits	protected vegetables		
Sporobolomyces roseus	fruits			
Trichosporon sp.	fruits			
T. pullulans	fruits, grapes, protected vegetables			

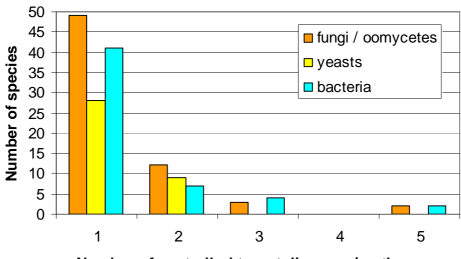
#### C. Bacteria

	Target disease / pathogen							
Microbial species	Botrytis	Powdery mildew	Rust	Downy mildew, late blight	Monillia rot			
Acinetobacter lwoffii	<mark>grapes</mark>							
Azotobacter				other				
Bacillus spp.	grapes, strawberry, protected vegetables, others	<mark>protected</mark> vegetables	others	<mark>potato</mark> , field vegetables	apricot			
B. amyloliquefaciens	arable crops, <mark>flowers,</mark> fruits, field vegetables, protected vegetables				peach			
B. cereus	<mark>flowers</mark> , legumes		others	tomato				
B. circulans	protected vegetables							
B. lentimorbus			others					
B. licheniformis	fruits, strawberry, protected vegetables							
B. macerans	legumes							
B. marismortui	strawberry							
B. megaterium	legumes, others							
B. pumilus	fruits, strawberry			tomato, <mark>others</mark>				
B. subtilis	flowers, fruits, <mark>grapes</mark> , legumes, strawberry, field vegetables, protected vegetables	cereals, grapes, strawberry, protected vegetables, others	<mark>legumes</mark>	<mark>grapes</mark> , potato, others	apricot, <mark>blueberry</mark> , <mark>cherry, peach</mark>			
B. thuringiensis	strawberry							
Bakflor (consortium of valuable bacterial physiological groups)	protected vegetables							
Brevibacillus brevis	field vegetables, protected vegetables	grapes, protected vegetables		grapes				
Burkholderia spp.				tomato				
B. cepacia	protected vegetables				cherry			
B. gladii					apricot			
B. gladioli	flowers							
Cedecea dravisae			others					
Cellulomonas flavigena				tomato				

	grapes, protected				
Cupriavidus campinensis	vegetables, others				
Enterobacter cloacae		protected vegetables		potato	
Enterobacteriaceae	strawberry				
Erwinia	fruits, others				
Halomonas sp.	strawberry, protected vegetables				
H. subglaciescola	protected vegetables				
Marinococcus halophilus	protected vegetables				
Salinococcus roseus	protected vegetables				
Halovibrio variabilis	protected vegetables				
Halobacillus halophilus	protected vegetables				
H. litoralis	protected vegetables				
H. trueperi	protected vegetables				
Micromonospora coerulea	protected vegetables				
Paenibacillus polymyxa	strawberry, protected vegetables				
Pantoea spp.	grapes, protected vegetables				
P. agglomerans	<mark>fruits</mark> , <mark>grapes</mark> , <mark>legumes</mark> , strawberry		legumes		apple, apricot, blueberry, cherry, peach, plum
Pseudomonas spp.	flowers, fruits, grapes, field vegetables		others	potato, tomato, field vegetables	<mark>apricot</mark>
P. aeruginosa	protected vegetables				
P. aureofasciens		<u>cereals</u>			cherry
P. cepacia	strawberry				peach
P. chlororaphis P. corrugata	strawberry				cherry
P. fluorescens	fruits, grapes, legumes, strawberry, protected vegetables, others	cereals, , protected vegetables others	legumes	<mark>grapes</mark> , <mark>potato</mark> , tomato, <mark>others</mark>	peach blueberry, cherry
P. putida	<mark>flowers</mark> , legumes, protected vegetables, others		cereals		
P. syringae	<mark>fruits</mark> , strawberry, field vegetables	grapes			apple, peach
P. reactans		strawberry			
P. viridiflava	fruits				
Rhanella spp.				potato	
R aquatilis	fruits				
Serratia spp.	flowers			potato	
S. marcescens S. plymuthica	protected vegetables				
Stenotrophomonas maltophilia			legumes		
Streptomyces spp.				tomato	
S. albaduncus	legumes			tomato	
S. ahygroscopicus	protected vegetables				
S. exfoliatus	legumes				
S. griseoplanus	legumes				
S. griseoviridis	protected vegetables, others				
S. lydicus	protected vegetables				
S. violaceus	legumes				
Virgibacillus marismortui	strawberry				
Xenorhabdus bovienii				potato	
X. nematophilus		protected vegetables			

One striking aspect of this inventory is that although the five target diseases / pathogens included in our review are airborne and affect mostly the plant canopy, the vast majority of cited biocontrol microorganisms are soil microorganisms. The scarcity of biocontrol agents originating from the phyllosphere could be due to actual lack of effectiveness, or it could be the result of a bias by research groups in favour of soil microbes when they gather candidate microorganisms to be screened for biocontrol activity. This question would merit further analysis as it may help to devise improved screening strategies. As "negative" results (the lack of effectiveness of tested microorganisms, for example) are seldom published, the completion of such an analysis would in turn necessitate direct information from research groups who have been implicated in screening for biocontrol agents, or the development of a specific screening experiment comparing equal numbers of phyllosphere and of soil microbial candidates.

Another striking aspect is that most of the beneficial micro-organisms inventoried in this study (49 fungi/oomycetes, 28 yeasts and 41 bacteria) are cited only for biocontrol of one of the five types of airborne diseases included in the survey (Figure 2). However, several species clearly stand out with a wide range of effectiveness, as they were successfully used against all five types of target diseases on a variety of crops. This includes the fungi *Trichoderma harzianum* and *Trichoderma viride* (2 of 12 species of *Trichoderma* reported as biocontrol-effective in the reviewed literature) and the bacteria *Bacillus subtilis* and *Pseudomonas fluorescens*.



Number of controlled target diseases / pathogens

Figure 2: Range of efficacy of 157 microbial biocontrol agents against five main types of airborne diseases. Detailed data are presented in Table 4.

Concerning *Fusarium oxysporum*. A data base interrogation with the key words "Fusarium oxysporum AND biological control" provided 2266 for the period 1973-2009. Using these key words we did not select only papers regarding biological control of diseases induced by *F. oxysporum* but also all the paper dealing with the use of strains of *F. oxysporum* to control diseases and weeds. There are quite many papers dealing with the use of different strains of *F. oxysporum* to control Broom rape (orobanche) and also the use of *F. oxysporum* f. sp. *erythroxyli* to eradicate coca crops.

We decided to limit our review to the two last years and to concentrate on references for which full text was available on line. Finally we reviewed 48 papers. All these papers were dealing with the selection and development of micro-biological control agents; only two were considering others methods. One was addressing the use of chemical elicitors to induce resistance in the plant; the

other was aiming at identifying the beneficial influence of non-host plant species either used in rotation or in co-culture. Based on this very limited number of papers the *formae speciales* of *F*. *oxysporum* the most frequently studied was *F.o. f. sp. lycopersici* (17 studies). Other included *f. spp. melonis, ciceris, cubense, niveum* and *cucumerinum*. The antagonists studied included *Bacillus* spp and *Paenibacillus* (16 papers), *Trichoderma* spp. (14 papers), fluorescent Pseudomonads (7 papers), Actinomycetes (5 papers), non pathogenic strains of *F. oxysporum* (5 papers), mycorrhizal fungi and *Penicillium*.

Most of the publications (28) reported on *in vitro* studies. Among them a few concerned the mechanisms of action of the antagonists, the others just related screening studies using plate confrontation between the antagonists and the target pathogens. In most of these papers (22) the *in vitro* screening was followed by pot or greenhouse experiments aimed at demonstrating the capacity of the antagonist to reduce disease severity or disease incidence after artificial inoculation of the pathogen. Finally only 9 publications report results of field experiments. Most of these papers concluded on the promising potential of the selected strains of antagonists able to decrease disease incidence or severity by 60 to 90%. Generally speaking, this limited literature review showed that most of the lab studies are not followed by field studies. There is a need for implementation of biological control in the fields.

# Identified knowledge gaps

Several types of knowledge gaps were identified in this review. They include:

- the near absence of information on biocontrol against diseases of certain important European crops such as winter arable crops.
- the scarcity of reports on biocontrol against several diseases of major economic importance on numerous crops, such as those caused by obligate plant pathogens (rusts, powdery mildews, downy mildews)
- the still limited (but increasing) body of detailed knowledge on specific mechanisms of action and their genetic determinism. The little knowledge available at the molecular level is concentrated on few model biocontrol agents such as *Trichoderma* and *Pseudomonas*.
- the still very limited information on secondary metabolites produced by microbial biocontrol agents
- the lack of understanding for generally low field efficacy of resistance-inducing compounds
- the lack of knowledge on variability in the susceptibility of plants pathogens to the action of BCAs and on possible consequences for field efficacy and its durability.

# References

Due to their high number, the references used in this chapter are presented, together with summary tables, in Appendices 1 to 6.

# **Chapter 2**

# Potential of biological control based on published research. 2. Beneficials for augmentative biocontrol against insect pests. The grapevine case study

#### **Massimo Giorgini**

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# Bibliographic survey on augmentative biological control against arthropod pests in selected crops

We carried out a preliminary bibliographic survey to quantify the literature on augmentative biological control of pests published from 1973 to 2008. The survey was restricted to crops relevant to case studies of ENDURE. They included grapevine; orchards: apple and pear; arable crops: corn and wheat; field vegetables: carrot and onion. Augmentative biological control (Van Driesche & Bellows, 1996) comprises of inoculative augmentation (control being provided by the offspring of released organisms) and inundative augmentation (control expected to be performed by the organisms released, with little or no contribution by their offspring).

Our bibliographic survey was conducted by using the CAB Abstracts database by entering the name of each crop and one key word selected from the following list in order to retrieve the maximum number of references. For each selected crop, the key words used for the bibliographic survey were: a) augmentative biological control; b) augmentation biological control; c) inoculative biological control; d) inundative biological control. The survey with these key words produced a very low number of results all of which were examined. For this reason we added two key words that were more general: e) insects biological control; f) mites biological control. For the searching criteria a to d, total records will be examined. In this case, given the extremely high number of records, only references within the period 1998-2008 were examined to select only the publications concerning the augmentative biological control. The results of this survey are reported in Appendix 7.

The analytical review of the scientific literature on augmentative biological control, presented in the rest of this chapter, was then focused on grapevine.

# Status of researches on augmentation of natural enemies to control arthropod pests in grapevine

The references extracted from the CAB Abstracts database, following the criteria described in the previous paragraph, were examined to identify those concerning the use of natural enemies in augmentation biological control in grapevine. The abstracts of 607 references were examined and only 70 papers reported data on application and efficiency of augmentative biocontrol (Table 5).

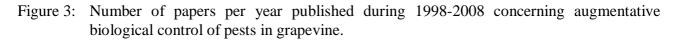
 Table 5:
 References extracted from the CAB Abstracts database and examined for reviewing augmentation biological control in grapevine.

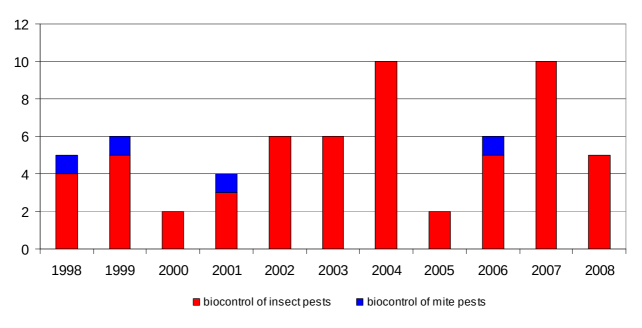
Key words	Total records	1998-2008
	(1973-2008)	
Augmentative biological control	7	6
Augmentation biological control	10	6
Inoculative biological control	4	1
Inundative biological control	7	3
Insects biological control		373
Mites biological control		190
Total references examined	28	579
Total references showing data on	70	0
augmentative biocontrol		

The survey includes records for grapevine, grape and vineyard.

# Results

Very few papers (62) on augmentative biocontrol in grapevine have been published during the period 1998-2008, with an average of 5.6 publications per year. Most references (93.5%) showed data on biological control of insects and only 4 papers on the biological control of mites were published (Figure 3).





The data extracted from the abstracts of the selected references were collected analytically in separate tables for each group of biocontrol agents (Appendix 8) and references were sorted chronologically (starting from the eldest). For each species of biocontrol agent, target species of pest, Country, type of augmentation (inundative, inoculative), type of test (laboratory, field), efficacy of biocontrol, additional information and results were reported.

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Data reported in Appendix 8 were summarized in Table 6, Table 7, Table 8, Figure 4 and Figure 5. A list of the biocontrol agents used in augmentative biological control in grapevine is reported in Table 6 and Figure 4. A list of groups and species of the targeted pests and the antagonists used for their control is reported in Table 7 and Figure 5; the efficacy of biocontrol agents is reported in Table 8.

The group of pests on which the highest number of researches on augmentative biocontrol has been carried out is Lepidoptera (60% of total references) with the family Tortricidae representing the main target (55%) (Figure 5) including the grape berry moths key pests *Lobesia botrana* and *Eupecilia ambiguella* (Table 7). *Bacillus thuringiensis* has resulted the most frequently used biocontrol agent against Lepidoptera by achieving an effective control of different targets in different geographic areas (Table 7, Table 8, Appendix 8.7). We sorted 28 references (39% of the total citations) dealing with the use of *B. thuringiensis* of which 23 references were referred to the control of *L. botrana*. The augmentation of egg parasitoids of the genus *Trichogramma* (Hymenoptera: Trichogrammatidae) resulted the alternative strategy to *B. thuringiensis* to control Lepidoptera Tortricidae (13 references, 16% of total citations) (Table 7, Table 8). Field evaluations indicated *T. evanescens* as a promising biocontrol agent of *L. botrana* (El-Wakeil *et al.*, 2008 in Appendix 8.1).

Fewer researches were carried out on augmentative biocontrol of other group of pests. First in the list were mealybugs (Hemiptera: Pseudococcidae) (9 references, 13% of the total citations). In field evaluations (4 papers) parasitoid wasps of the family Encyrtidae have resulted extremely active and promising to be used in augmentative biocontrol of mealybugs (Appendix 8.2).

Antagonists used in augmentative biocontrol in grapevine were mainly represented by insect pathogens (59% of the total citations), including the bacterium *B. thuringiensis*, fungi and nematodes (Figure 4, Table 6). Beside the efficacy of *B. thuringiensis*, promising results were obtained from researches in the control of the grape phylloxera *Daktulosphaira vitifolie*, a gallforming aphid, by soil treatments with the fungus *Metarhizium anisopliae* (Table 8, Appendix 8.5). Once controlled by grafting European grape cultivars onto resistant rootstocks, the grape phylloxera has gone to resurgence in commercial vineyards worldwide and new biological control strategy could be necessary to complement the use of resistant rootstocks and to avoid the distribution of chemical insecticides in the soil.

Entomophagous arthropods, including parasitoid wasps and predators represented 41% of the total citations (Figure 4, Table 6). Best results were obtained from researches on parasitoids (18 references), namely the use of Trichogrammatidae and Encyrtidae in augmentative biocontrol of grape moths (Tortricidae) and mealybugs (Pseudococcidae) respectively (Table 7, Table 8, Appendix 8.1 and 8.2). Among predators, augmentation of Phytoseiidae mites has produced some positive results in controlling spider mites and eriophyid mites on grape (Table 7, Table 8, Appendix 8.3).

#### **Brief considerations**

Key pests of grapevine like *L. botrana* and *E. ambiguella* can be controlled effectively with augmentative strategies that rely on the use of *B. thuringiensis*. To date, formulations of *B. thuringiensis* are currently used in IPM strategies. The specificity of *B. thuringiensis* could be a problem in those vineyards where other pests can reach the status of economically importance, if not controlled by indigenous and/or introduced natural enemies. Researches on augmentative biocontrol should be implemented in order to develop new strategies to solve problems related to emerging pests and alternatives to *B. thuringiensis* if resistant strains should appear in target species.

# References

Due to their high number, the references for this chapter are presented in Appendix 8.

 Table 6:
 Biocontrol agents evaluated in researches on augmentative biological control of pests in grapevine.

Target pests and biocontrol	References before 1998	<b>References</b> 1998-2008	Number of citations	
<b>BIOLOGICAL CONTROL OF I</b>				
Bacteria [1 specie - Bacillus thuringiensis	s: 2 subspecies]	0	28	28
(subsp. kurstaki, subsp. aizaw				
Fungi	[5 species]	0	10	
- Metarhizium anisopliae				7
- Beauveria bassiana				2
- Beauveria brongniartii				1
- Verticillium lecanii				1
- Clerodendron inerme				1
Nematodes	[5 species]	1	3	_
- Steinernema spp.	2 <i>spp</i> .			2
- Heterorabditis spp.	3 spp.	•		3
Parasitoid Hymenoptera	[15 species]	2	16	10
- Trichogramma spp. (Trichogram				13
- Coccidoxenoides spp. (Encyrtidae	· · · ·			2
- Anagyrus spp. (Encyrtidae)	2 spp.			3
- Muscidifurax raptor (Pteromalida		2	4	1
Predators	[5 species]	2	4	2
- Chrysoperla (Neuroptera: Chryso	pidae) 3 spp.			3 2
- Cryptolaemus montrouzieri (Coleoptera: Co	a a sin allida a)			2
- Nephus includens (Coleoptera: Co				1
	,			1
<b>BIOLOGICAL CONTROL OF N</b>	MITES			
Predators (Acari: Phytoseiidae)	[4 species]	2	4	
- Typhlodromus pyri	[+ species]			5
- Kampimodromus aberrans				2
- Amblyseius andersoni				1
- Phytoseiulus persimilis				1

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 Table 7:
 Number of references on augmentative biocontrol agents per group and species of target pest in grapevine.

Pest	References	Bacillus thuringiensis (2 subspecies)	Trichogramma (10 species)	other parasitoids (5 species)	Predators of mites Acari: Phytoseidae (4 species)	Predators of insects Coleoptera: Coccinellidae (2 species)	Predators of insects Neuroptera: Chrysopidae (3 species)	Fungi (5 species)	Nematodes (5 species)
Lepidoptera: Tortricidae	39								
<i>Lobesia botrana</i> (grape berry moth)	28	23	5						
(grape berry moth) Eupoecilia ambiguella (grape berry moth)	б	3	3						
(grape berry mon) <i>Epiphyas postvittana</i> (light brown apple moth)	3		3						
(South American tortricid moth)	3	1	2						
Bonagota cranaodes (Brasilian apple leafroller)	2		2						
<i>Endopiza viteana</i> (grape berry moth)	2		2						
Sparganothis pilleriana (grape leafroller)	1	1							
<i>Epichoristodes acerbella</i> (South African carnation tortrix)	1	1							
Lepidoptera: Pyralidae	1								
<i>Cryptoblabes gnidiella</i> (honey moth)	1	1							
Lepidoptera: Arctiidae	1								
<i>Hyphantria cunea</i> (fall webworm)		1							
Lepidoptera: Sesiidae	2								
Vitacea polistiformis	2								2

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# Table 7 (continued)

Hemiptera:	9					
Pseudococcidae						
Planococcus ficus	6	4	2			
		Encyrtidae				
Pseudococcus maritimus	1		1			
Pseudococcus longispinus			1			
Maconellicoccus hirsutus		1			1	
		Encyrtidae				
Hemiptera:	3					
Cicadellidae						
Erythroneura variabilis	3			3		
Erythroneura elegantula	3			3		
Hemiptera:	5					
Phylloxeridae						
Daktulosphaira vitifoliae					4	1
(grape phylloxera)						
Diptera:	1					
Tephritidae						
Ceratitis capitata	1	1				
		Pteromalidae				
Coleoptera:	2					
Scarabeidae						
Melolontha melolontha	2				1	1
Thysanoptera:	3					
Thripidae						
Frankliniella occidentalis	2				2	
grape thrips	1				1	
Acari:	6					
Tetranichidae						
Panonychus ulmi	5	5				
Tetranychus urticae	1	1				
Tetranychus kanzawai	1	1				
Eotetranychus carpini	2	2				
Acari:	2		-			
Eriophyidae						
Calepitrimerus vitis	1	1				
Calomerus vitis	1	1	 			

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Groups of Pests	Biocontrol agents	Total number of references	Number of references reporting dat on efficacy in pest and related damage control <sup>*</sup> Laboratory assays Field evaluation			
			Laboratory assays	Field evaluation		
Lepidoptera: Tortricidae	Bacillus thuringiensis	26	2 +	16 + 1 -		
	<i>Trichogramma</i> spp. parasitoids	13	1 -	9 + 1 -		
Lepidoptera: Pyralidae	Bacillus thuringiensis	1		1 +		
Lepidoptera: Arctiidae	Bacillus thuringiensis	1		1+		
Lepidoptera: Sesiidae	Nematodes	2	2 +	1 + 1 + (greenhouse)		
Hemiptera: Pseudococcidae	Encyrtidae parasitoids	5		4 +		
	Coccinellidae	3		1 + (greenhouse)		
	Fungi	1		1 +		
Hemiptera: Cicadellidae	Chrysopidae	3		2 -		
Hemiptera: Phylloxeridae	Nematodes	1	1 +			
i nynoxei iuae	Fungi	5	1 +	2 + 1 -		
Diptera: Tephritidae	Pteromalidae parasitoids	1	1 +	1+		
Acari: Tetranichidae	Phytoseidae	6		4 +		
Acari: Eriophyidae	Phytoseidae	2		1 +		
Coleoptera: Scarabeidae	Nematodes	1		1 +		
	Fungi	1		1 +		
Thysanoptera: Thripidae	Fungi	3	1 +	2 +		

Table 8:Number of references reporting data on the efficacy of augmentative biocontrol of<br/>pests in grapevine.

\* + means effective, - means not effective biocontrol agent

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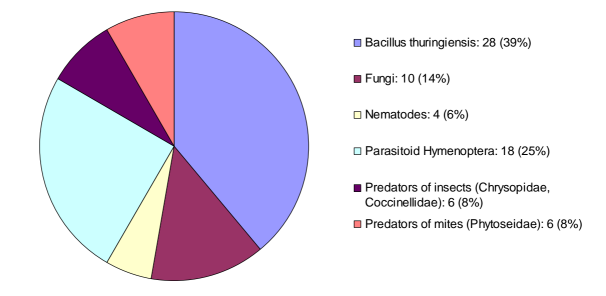


Figure 4: Groups of biocontrol agents investigated in augmentative biological control researches in grapevine. Number of references for each group is reported.

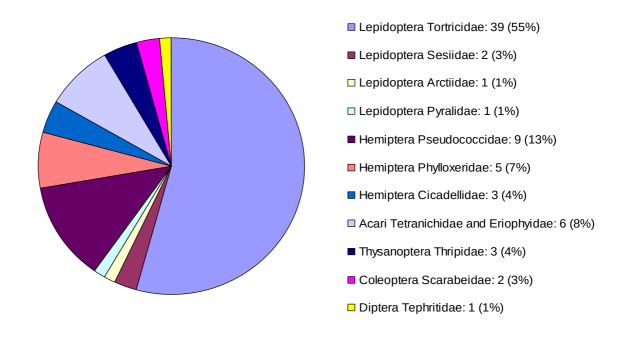


Figure 5: Groups of target pests investigated in augmentative biological control researches in grapevine. Number of references for each group is reported.

# **Chapter 3**

# Potential of biocontrol based on published research.3. Research and Development in classical biological control with emphasis on the recent introduction of insect parasitoids

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#### Scope of the review

Defined as "the intentional introduction of an exotic, usually co-evolved, biological control agent [hereafter BCA] for permanent establishment and long-term pest control', classical biological control [hereafter CIBC] is a pest control strategy that has crystallized numerous studies since more than one century and provided numerous efficient solutions for pest control. The main advantages and risks of this strategy can be summarized as follows. In a context of the globalisation of international trade and human mobility, an ever growing number of exotic pests emerge locally. Such species can rapidly pullulate and jeopardize cultural practices. This general trend can also be favoured by global climatic changes that may allow the development of agronomic pests beyond their initial distribution area and increase their demography. Within this context, CIBC appears often to be the first way to try to regulate such pest populations. Moreover, when successful, ClBC appears to be very economic insofar as financial costs are only associated with the identification, evaluation and initial releases of exotic BCA. Contrary to other pest control strategy, the implication of practitioners and other costs are not necessary after the establishment of the BCA. The overall financial costs of such operations are consequently rather limited with regard to the durability of the pest control, in particular when the local introduction of a new BCA benefits from the previous experiences in other countries. Nevertheless, at least two kinds of risks are usually associated with ClBC. First of all, the average success rate of ClBC varies between 10 and 30% according to the authors for a total of more than 5000 introductions worldwide during the last century. As consequence, such operations may also appear too risky to be funded. Another risk is those associated with the non-target effects. Although few cases have been reported, their echoes may have contributed to a more harmonized approach and in some countries to more or less stringent regulations.

As consequences, classical biological programmes are at the crossroad of several concerns:

- agronomic; insofar as each introduction of exotic BCA is obviously an hope for the producers;
- scientific; CIBC namely questions both ecologist and evolutionist in order to optimize the probability of establishment while minimizing the non-target effects. Their implication on such issues nevertheless depends on their own interest (in term of scientific question and/or possibility or publishing);
- political; since the introduction of BCA may depend on regulation or homologation decided at national or international levels;

- financial; since the development of CIBC is relying on various sources of funding (agronomic partners, scientific partners, politic institutions) with various interests and rationale (more or less short-term results, scientific excellence versus applied objectives).

Within this context, global evaluations of CIBC programmes are necessary to better understand the evolution of this practice and try to improve its use and efficiency. This has been repeatedly achieved during the last years either through reviews or meta-analysis. Based on a large (but probably not exhaustive) bibliographic survey, the present work aims to give a complementary point of view with the willingness to portray a realistic "state of the art" of Research and Development programmes of CIBC against arthropods. This chapter also firstly gives a broad temporal survey of the publication and a more precise survey of the literature for the decade [1999; 2008]. Biocontrol programmes against arthropods were then more precisely detailed with the objectives to give qualitative cues about the main pests and the types of related studies. Finally, a particular emphasis has been put on recent introductions of exotic insect parasitoids.

Based on these data, we also address some more or less important subjective recommendations based on our own opinion.

# Method

A large bibliographic survey has been conducted with the CAB abstracts. Several combinations of key-words were used with various successes. Too broad (e.g; cases for which discussion about CIBC are marginal) or unprecise (e.g. cases for which a pest is not precised) publications were excluded. A total of 764 publications were found using the keywords "classical biological control" or "classical biocontrol". 452 papers were published during the period [1999-2008] but about 30% were not relevant with regard to the purpose of this survey and have been discarded. Using the more complex combinations ["biological control" AND "exotic" AND "introduction"], 329 CIBC-related publications were obtained but only 253 dressed precisely questions related to classical biological control. 117 were published during the selected temporal frame but only 81 were relevant with regard to our objectives. Additionally, 47 CIBC-related publications were obtained using the more keywords association ["biological control" AND "exotic" AND "importation"] with 17 papers for the last ten years. Most of this literature was dedicated to the risk or regulatory aspects associated with the importation of exotic BCA so that only 7 relevant publications with regard to our objectives. Finally, 130 publications were found using "acclimatization" AND "biological control" for only one relevant publication for the targeted period. A total of 358 publications were also obtained which is probably for far from being exhaustive. For instance, 37 new references about BCA introductions were found in addition to the first 35 references found with the previous key-words combinations (see Table 9). Additional bibliographic research were also realised for some taxa (see below)

[Remark: Although the terms "classical biological control" or "classical biocontrol" may be not as explicit as others ("introduction", "importation"), the generalization of their use in titles, key-words or abstracts should be nevertheless used in order to improve the efficiency of bibliographic survey]

# **General trends**

The temporal survey shows a quite regular increase of CIBC related publications with a mean of about 45 hits / year for the last ten years (Figure 6). Within this period, we observe a relative stability between the different combinations of pests and BCA (Figure 7). The main part of the publications (56%) of the cases deals with the biocontrol of phytophagous arthropods on which we will focus here. 42% of the papers deal with the biocontrol of weed. In this case, BCA are for 57% of the cases phytophagous insects and for 41% fungi (data not shown).

More than 70 arthropod pests were listed which cover 7 orders and approximately 40 families. As shown in Figure 8, Hemiptera and Lepidoptera were the two main orders with a total of 66% of the pest species and 70% of the publications. If the citation rate / order is highly correlated with the number of pests / order, this trend hides a great variability at the infra-order level. Indeed, the citation rate highly differs with regard to the pest species with a median of 2 papers / pest species and a range from 1 to 13 citations. The 13 most cited pests are listed in Figure 9. Two main observations can be drawn from this short list.

 $\Rightarrow$  Firstly, this list is quite equally composed of either very specialist pests like *Phyllocnistis citrella* (on Citrus species), *Mononychellus tanajoa* (on cassava) or *Toxoptera citricida* (on Citrus species) or more generalist taxa like *Homalodisca vitripennis*, *Lymantria dispar* or *Pseudococcus viburni*. All of them are phytophagous pests whose damage are linked either to their herbivory, consumption of sap or virus transmission except the particular case of the fire ant *Solenopsis invicta* which is responsible for direct nuisance on farmers or indirect ecological modifications in the agrosystems.

 $\Rightarrow$  The second observation is that the percentage of CIBC related publications / pest is negatively correlated with the corresponding total number of references (including also studies on other pest control strategies and/or various biological topics). For instance, 22% of the 32 references focusing on *H. vitripennis* explicitly deal with classical biological control while this percentage falls down to only 1% to 3% for well documented species like L. dispar, *S. invicta* or *D. virgifera virgifera*. This may be explained by the fact that CIBC is mainly considered as a "pionneer" pest control strategy that are developed either soon after the emergence of a new invasive pest or on "non biological model" for which the investigations on other biological aspects are limited.

[Remark: Although Classical Biological Control can be perceived as a "pioneer" pest control strategies on non "biological models", substantial investments are required on several biological aspects (e.g. community ecology, population genetics)]

# **Biocontrol agents used**

The biocontrol agents related to CIBC (hereafter CIBCA) against arthropod species were not detailed in only 12% of the papers. These are in most of the cases either prospective works (55%) such as faunistic inventories of natural enemies on "new" pests like *Diabrotica virgifera virgifera* or retrospective studies (35%) on advanced programmes that take into account several BCA (see Appendix 9.1). Among the documented cases, 76% of CIBC programmes were based on the use of insect parasitoids. Pathogens and nematodes on one side and predatory arthropods on the other side are equally represented with about 12% of the publications for each case.

#### Pathogens and Nematodes as candidate for CIBCA

The particular cases of pathogens and nematodes have been recently reviewed by Hajek and co-workers  $(62, 63^3)$ . Our own survey indicates that half of the papers actually deal with entomopathogenic fungi. Six pest species were identified including two mites (Aceria guerreronis and Mononychellus tanajoa) and two insects (Aphis gossypii and Coptotermes formosanus). However, except for the evaluation of Neozygites species against M. tanajoa (14, 39, 42, 43), other attempts seem to be rather limited. With regard to the catalogue of Hajek et al.(62), two other cases of entomopathogen fungi were missed in our own survey. These are the introductions of Entomophaga maigmaiga and Metarhizium anisopliae, against respectively the Lymantria dispar and the Curculionidae Otiorynchus nodosus for which the sources of Hajek and coworkers were mainly personal communications. The rather limited use of entomopathogenic fungi in CIBC was also confirmed by the review of Shah and Pell(156). The use of viruses as biocontrol agent for ClBC against arthropod pests were only documented fort three cases that are the Lepidoptera species Anticarsia gemmatalis (48, 127) and Lymantria dispar (16) and the Coleoptera Oryctes rhinoceros (81). Microspodia as candidate for CIBC were reported in only two studies (25, 165). The sole case of the use of nematodes is the study of Hurley et al. (79) who studied the extension of the use of parasitic nematode Deladenus siricidicola against the woodwasp Sirex noctilio.

#### Predatory arthropods as candidate for CIBCA

The literature about predatory arthropods is dominated by four case-studies. The first one is the classical biocontrol of the cassava green mites M. tanajoa by Typhlodromalus aripo and, to a lesser extent, T. manihoti. All these studies are the extension of a very large classical biocontrol programme at a continental scale; two main issues were addressed during the recent decade that are the introduction and field evaluation of T. aripo in Mozambique and Malawi (125, 194) and the ecological interactions with other species (14, 124, 193) or plants(55). The second case-study is those of the predatory ladybird *Harmonia axyridis* (19, 90, 91, 137). The main concern of these publications is nevertheless not the Research and Development in CIBC but rather the risks of non-intended effects and geographic spray of this insect that is now considered as a world-wide invasive species. Another case of the use of ladybird is those of Cryptolaemus montrouzieri and Scymnus coccivora which have been successfully used to control the hibiscus mealybug Maconellicoccus hirsutus (51, 86, 103) which is the extension of a worldwide use of these species. The fourth main case-study is the classical biocontrol programme of *Prostephanus truncatus*, a serious pest of stored maize beetle using *Teretrius* (formerly *Teretriosa*) nigrescens (73, 169, 170). The lasts reported uses of predatory arthropods as candidate for CIBC were those of the Coleoptera Laricobius nigrinus against the adelgid Adelges tsugae (197) and the phytoseid Neoseiulus baraki against the coconut mite A. guerreronis (119). Contrary to other cases which were the continuity of older programmes, these two studies are associated with new BCA inventories undertaken during the last ten years - see respectively (196) and (99).

#### Insect parasitoids as BCA

#### Related journals papers and categorization of the studies

In total, 125 publications were used for this analysis. Only 14% were associated to proceedings of meetings or other supports than journals. 43 different journals were identified but 50% of the publications were published only by five: Biological Control (21%), BioControl (8%), Biocontrol Science and Technology (7%), Florida Entomologist (7%) and

<sup>&</sup>lt;sup>3</sup> within this Chapter, numbers in parentheses refer to references listed in Appendix 9

Bulletin of Entomological Research (7%). Impact Factors are respectively 1.805, 1.957, 0.874, 0.886 and 1.415.

The types of the works were categorized according to the simplified sequential steps in R&D of biological programmes: BCA Inventories  $\Box$  BCA characterization (systematic, molecular tools) Pest or BCA rearing  $\Rightarrow$  BCA biology (life history traits, thermal biology, behavioural ecology)  $\Rightarrow$  Pre-release survey  $\Rightarrow$  BCA introduction  $\Rightarrow$  Post-release survey. Studies related to "non-target effects" (i.e. the direct or indirect impacts of the CIBCA on non-target species) as well as those related to the "biocontrol disruption" (i.e. the negative impacts of organisms on the CIBCA) (details in Appendix 9.3) were also categorized. As shown in Figure 10, most of the CIBC related publications logically deals either with BCA biology, BCA introductions or post-release surveys which are central steps of the CIBC programmes. A strong discrepancy nevertheless exists between the different types of work in term of scientific publication; highest Impact Factors are relied to studies linked to Non-intended effects, Biocontrol disruption or BCA Biology.

[Remark: The different steps of R&D in Classical Biological Control are currently unequally promoted with regard to "scientific criteria", with a clear emphasis on community ecology including non target effects. Such trend may be detrimental to the short-term development of less gratifying tasks and consequently on the whole dynamism of ClBC.]

### **BCA Introductions**

As shown in Table 9, 65 introductions were recorded during the period of 1991-2006. This list is probably not exhaustive insofar as "cryptic introductions" may have been missed. This list does not also cover all the R&D in classical biocontrol programmes since some programmes may have been interrupted before releases. A faunistic inventory of the natural enemies of the North American leafhopper *Scaphoideus titanus* has for instance been led by our lab in 2000-2002 but the rearing of BCA candidates (mainly dryinids and egg-parasitoids) were not successful.

 $\Rightarrow$  All these releases involve 55 different biocontrol agents (all hymenopteran except the *Pseudacteon* species used against the fire ant *Solenopsis invicta*) and 35 pests. 57% of these pests were *Hemiptera*, other being quite equally distributed between Lepidoptera, Diptera, Hymenoptera and Coleoptera.

 $\Rightarrow$  Most of these introductions were realized against pest found on orchards and in particular Citrus. Other targeted crops were mainly tropical productions, ornamental or forest.

 $\Rightarrow$  Most of the BCA introductions (42%) were realized in Europe or neighbouring countries (including Mediterranean Basin) and in North America (26%). The percentages of introductions in other geographical areas were: Australia-New Zealand and neighbouring islands (12%), South America (8%), sub-Saharan Africa (8%), Pacific Islands (3%), Asia (1%).

 $\Rightarrow$  The total number of released parasitoids and number of sites were highly variable ranging respectively from 456 to 660000 individuals and from 2 to 132 sites. The percentage of establishment was 83% and, when established, high parasitism was found in 42% of the cases. It is noteworthy that these values are relatively high compared to other estimates and we are currently unable to say if this is linked to an improvement of practices or methodological differences or biases.

[Remark: With regard to natural or other human-mediated introductions of exotic species, species flow associated with the ClBC seems to be rather limited. Although possible non-intended effects cannot be excluded (their studies having to be increased), we fear that too drastic regulations could severely disturbed R&D programmes]

[Remark: Estimating the success of ClBC is difficult because of methodological several biases ("cryptic introductions", barriers linked to languages and/or publishing). Shared international database should be necessary for more accurate estimation as well as an increasing traceability.]

[Remark: In parallel with the geographical expansion of their related pests, some biocontrol agents have been repeatedly released and established worldwide. Population genetics studies in such pest-BCA interactions should be particularly interesting to understand local adaptations, co-evolutionary processes and ultimately, the durability of Classical Biological Control.]

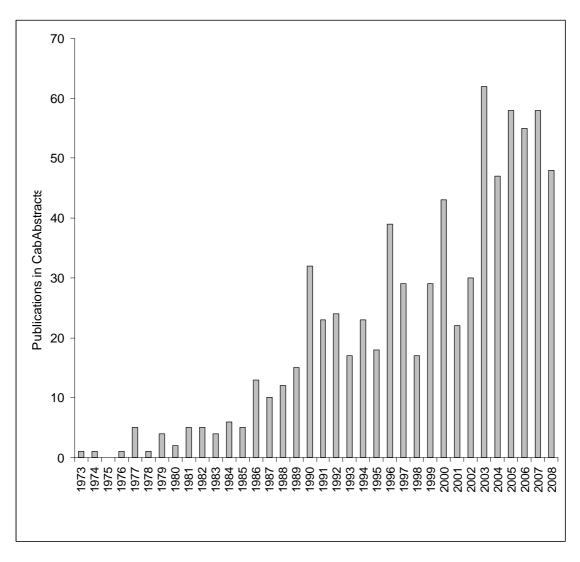


Figure 6: Large-scale temporal survey of the publications associated with classical biological control

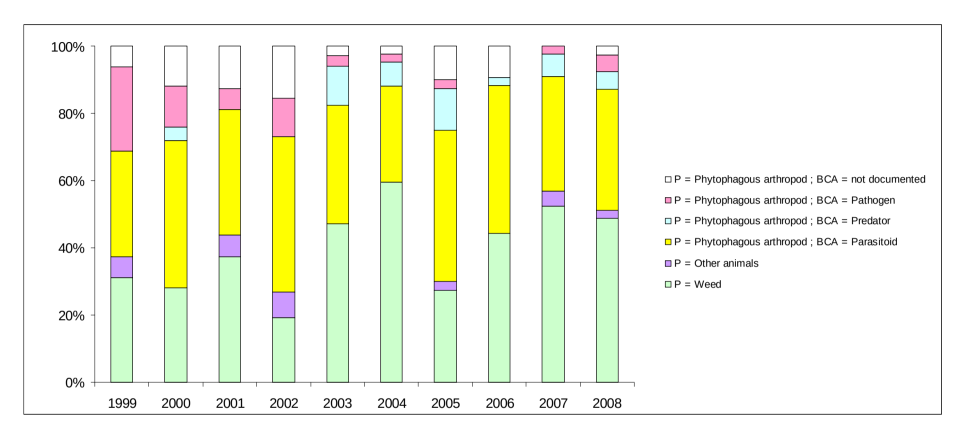


Figure 7: Relative importance of the different types of biocontrol during the temporal frame [1999-2008]

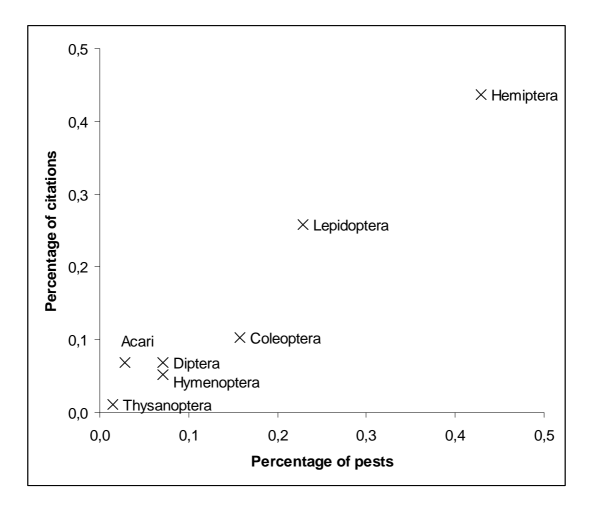


Figure 8: Number of pest species and related citation rate by orders during the period [1999; 2008]

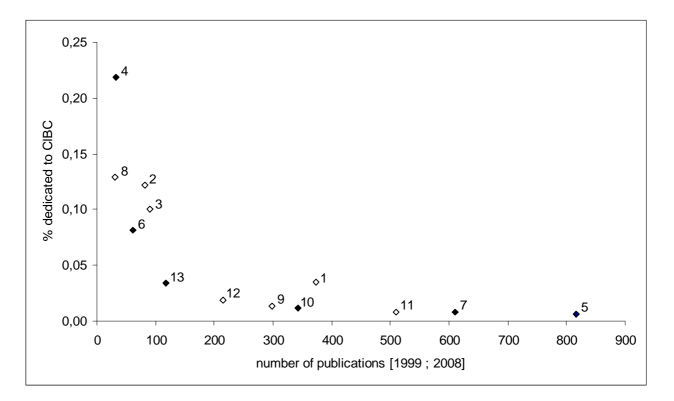


Figure 9: Relationships between the number of publications associated to the main pests and the relative percentage of CIBC related studies.

Pest species are ranked in the decreasing order in number of publications : 1 : *Phyllocnistis citrella* ; 2 : *Mononychellus tanajoa* ; 3 : *Toxoptera citricida* ; 4 : *Homalodisca vitripennis* ; 5 : *Lymantria dispar* ; 6 : *Pseudococcus viburni* ; 7 : *Solenopsis invicta* ; 8 : *Aleurocanthus spiniferus* ; 9 : *Bactrocera oleae* ; 10 : *Chilo partellus* ; 11 : *Diabrotica virgifera virgifera* ; 12 : *Diatraea saccharalis* ; 13 : *Maconellicoccus hirsutus*. Specialist and generalist pests are respectively indicated by white and dark diamonds.

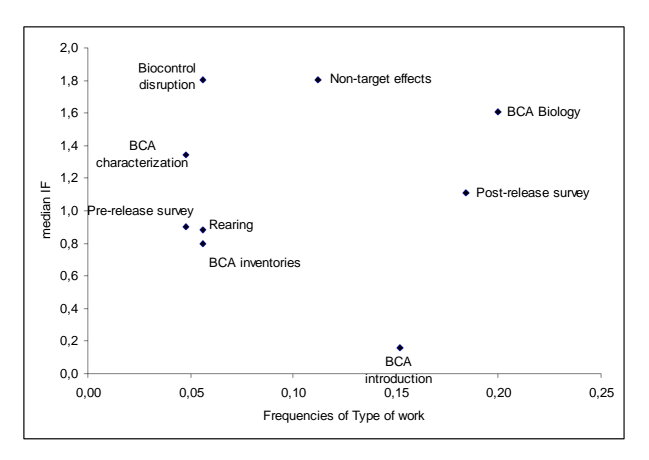


Figure 10: Frequencies of papers and associated median IF related to the different categories of work

Targeted pest	Сгор	BCA Name	Introduction Area	Introduction Date	Individuals (sites)	Outcome	References
Aleurocanthus woglumi	Citrus	Amitus hesperidum	Trinidad	2000	1600 (3)	Establishment High parasitism	(White et al., 2005)
Aleurodicus dispersus	Banana	Encarsia guadeloupae Lecanoideus floccissimus	Spain (Tenerife)	-	_	_	(Nijhof et al., 2000)
		Encarsia haitiensis	Australia	1992-1996	_	Establishment	(Lambkin, 2004)*
Aleurolobus niloticus	Orchard	Eretmocerus siphonini	Egypt	1998-1999	237000	Establishment High parasitism	(Abd-Rabou, 2002)
Aonidiella aurantii	Citrus	Aphytis lingnanensis	Spain	2000	_	Establishment	(Pina and Verdu, 2007)*
Aphis gossypii	Vegetable	Lysiphlebus testaceipes	Bulgaria	_	_	Establishment	(Dimitrov et al., 2008)*
Bactrocera dorsalis	Orchard	Fopius arisanus	French Polynesia	2003		Establishment High parasitism	(Vargas et al., 2007)*
Bemisia tabaci	Arable crops Vegetable	Eretmocerus hayati	Egypt	2000-2002	200700	Establishment	(Abd-Rabou, 2004)*
Ceratitis capitata	Orchards (incl. <i>Citrus</i> )	Diachasmimorpha krausii Fopius arisanus Fopius ceratitivorus Psyttalia concolor (complex)	Israel	2002-2004 2002-2004 2002-2004 2002-2004	75881 258750 58860 75881	Establishment ? Establishment ?	(Argov and Gazit, 2008)*
Ceroplastes rubens	Orchard (incl. <i>Citrus</i> )	Anicetus beneficus	Papua New Guinea	2002	2200 (2)	Establishment	(Krull and Basedow, 2005)
Chilo sacchariphagus	Sugarcane	Xanthopimpla stemmator	Mozambique	2001	5000 (5)	?	(Conlong and Goebel, 2002)
Cinara cupressivora	Forest Ornamenta	Pauesia juniperorum	Mauritius	2003-2004	1500	?_	(Alleck et al., 2006)
Coccus viridis	Citrus Coffee	Diversinervus sp. near stramineus	Australia	_	(4)	Establishment High parasitism	(Smith et al., 2004)*
Ctenarytaina eucalypti	Forest Ornamental	Psyllaephagus pilosus	Chile	2001	_	Establishment High parasitsm	(Rodriguez and Saiz, 2006)*
Diaphorina citri	Citrus	Diaphorencyrtus aligarhensis	USA	_	_	_	(Hoy, 2005)
		Tamarixia radiata	USA	_	_	_	
Diatraea saccharalis	Sugarcane	Cotesia flavipes	USA	2001-2002	(4)	Failure	(White et al., 2004)*
Dryocosmus kuriphilus	Forest Ornamental	Torymus sinensis	Italy	2005-2006	1100 (14)	Establishment	(Aebi et al., 2007)

# Table 9: Recent introductions of parasitoids as Classical Biocontrol agents

Legend : \_ : data not available ; ? : long-term establishment not sure ; \* : additional references

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# Table 9: Recent introductions of parasitoids as Classical Biocontrol agents (continued)

Hemiberlesia pitysophila	Forest	Coccobius azumai	China	2002		Establishment	(Wang et al., 2004)*
Homalodisca vitripennis	Wide range	Gonatocerus ashmeadi	Tahiti	2005	14000 (27)	Establishment High parasitism	(Grandgirard <i>et al.</i> , 2007a) (Grandgirard <i>et al.</i> , 2008) (Petit <i>et al.</i> , 2008)
Hypothenemus hampei	Coffee	Cephalonomia stephanoderis	Cuba	_	(2)	?	(Murguido Morales <i>et al.</i> , 2008)*
		Phymastichus coffea	Colombia	_	(41)	Establishment	(Aristizabal <i>et al.</i> , 2004)*
Lilioceris lilii	Ornamental	Diaparsis jucunda	USA	_	_	_	(Casagrande and Tewksbury, 2005)*
		Lemophagus errabundus		_	_	_	
		Tetrastichus setifer		2001	1700 (21)	_	(Tewksbury <i>et al.</i> , 2005)*
Liriomyza trifolii	Vegetables	Dacnusa sibirica	Egypt	_	90000	?	(Abd-Rabou, 2006)*
	-	Diglyphus isaea		_	90000	?	
Listronotus bonariensis	Pasture	Microctonus hyperodae	New Zealand	1991-1998	66000 (121)	_	(McNeill <i>et al.</i> , 2002) (Phillips <i>et al.</i> , 2008)
Maconellicoccus hirsutus	Wide range	Anagyrus kamali	North America			Establishment High parasitism	(Kairo <i>et al.</i> , 2000)
Metcalfa pruinosa	Wide range	Neodryinus typhlocybae	Greece	2006	_	Establishment	(Anagnou-Veroniki et al., 2008)
Ophelimus maskelli	Forest Ornamental	Closterocerus chamaeleon	Israel	2005-206	12000 (6)	Establishment High parasitism	(Protasov et al., 2007)*
		Closterocerus sp.	Italy	_	(5)	Establishment High parasitism	(Rizzo et al., 2006)*
Paracoccus marginatus	Wide range	Acerophagus papayae	Palau	2003-2004	_	Establishment High parasitism	(Muniappan et al., 2006)
		Anagyrus loecki		2003-2004	_	Establishment High parasitism	
		Pseudleptomastix mexicana		2003-2004	_	Failure	

Legend : \_ : data not available ; ? : long-term establishment not sure ; \* : additional references

Phyllocnistis citrella	Citrus	Ageniaspis citricola	Morocco	1995-1996	_	Failure	(Rizqi et al., 2003)
			USA	_	_	Establishment High parasitism	(Hoy, 2005)
			Italy	1995	_	Failure	(Siscaro et al., 2003)
			Italy	1996-1997	_	Establishment High parasitism	(Siscaro <i>et al.</i> , 1999)
			USA	1999	25000 (132)		(Paiva et al., 2000)
			Argentina Brazil	2001-2004 1999	25000	?	(Zaia <i>et al.</i> , 2006) (Paiva <i>et al.</i> , 2000)*
		Cirrospilus ingenuus	Morocco	_	_	_	(Rizqi et al., 2003)
		Cirrospilus quadristriatus [C. ingenuus]	USA	_	_	Establishement	(Hoy, 2005)
		Citrostichus phyllocnistoides	Italy	1995	_	Establishment	(Siscaro <i>et al.</i> , 2003)
			Morocco	2000	_	Establishment	(Rizqi et al., 2003)
			Spain	1996-1999	_	Establishment High parasitism	(Garcia-Mari <i>et al.</i> , 2004)*
		Quadrastichus sp	Morocco Italy	1995	_	_ Failure	(Rizqi <i>et al.</i> , 2003) (Siscaro <i>et al.</i> , 2003)
			Italy	1996-1997	_	Failure	(Siscaro <i>et al.</i> , 1999)
		Semielacher petiolatus	Morocco	1996-1997		Establishment	(Rizqi et al., 2003)
Pseudococcus viburni	Orchard	Pseudaphycus maculipennis	New Zealand	2001	_	_	(Charles, 2001)
Saissetia coffeae	Olive	Coccophagus cowperi	Egypt	2001-2003	300000	Establishment	(Abd-Rabou, 2005)*
Siphoninus phillyreae	Orchard	Eretmocerus siphonini	Egypt	1998-1999	237000	Establishment High parasitism	(Abd-Rabou, 2002)
Sirex noctilio	Forest	Ibalia leucospoides	South Africa	1998-2001	456	Establishment	(Tribe and Cillie, 2004) <sup>*</sup>
Solenopsis invicta	_	Pseudacteon curvatus	USA	2003	10100 (2)	Establishment	(Vazquez et al., 2006)
		Pseudacteon obtusus	USA	2006		?	(Gilbert et al., 2008)
		Pseudacteon tricuspis	USA	1999-2001		Establishment	
Tephritidae sp.	Orchard (incl. <i>Citrus</i> )	Diachasmimorpha longicaudata	Brazil	2002	34000 (2)	Failure	(Alvarenga et al., 2005)
Toxoptera citricida	Citrus	Lipolexis oregmae	USA	2000-2002	33500	Establishment	(Hoy, 2005) (Persad <i>et al.</i> , 2007)
Yponomeuta malinellus *	Orchard	Ageniaspis fuscicollis	Canada	1987-1997	_	Establishment	(Cossentine and Kuhlmann, 2007)*

# Table 9: Recent introductions of parasitoids as Classical Biocontrol agents (continued)

Legend : \_ : data not available ; ? : long-term establishment not sure ; \* : additional references

# **Chapter 4**

# **Registered Biocontrol Products and their use in Europe**

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### **Collection of information**

A small team formed by ACTA and IBMA conducted a survey on biological active substances approved in the European Union and on Biological Control Products (BC products) authorised in five European countries. The investigation focused on crops covered by ENDURE RA1case studies. The frame of the present survey was defined in a meeting on 9<sup>th</sup> January 2009 in Basle, and the work was performed during the period from April to September 2009.

To compile a list of registered biocontrol products, the online EU Pesticides Database was consulted on 21<sup>st</sup> April 2009. Data were retrieved and the list was reorganised and the information about use categories complemented with the help of the inclusion directives where necessary. Substances deemed suitable for biocontrol were identified and it was decided to distinguish four major groups: micro-organisms, semiochemicals (attractants), botanicals and "other plant protection substances of natural origin".

This study was complemented by an analysis of specific uses of products commercialized in four countries of the EU (France, Germany, Spain and the United Kingdom). A fifth country, Switzerland was included in the study for comparison, because it has not been restricted by the implementation of Directive 91/414/EEC (superceded in June 2011 by EC regulation No 1107/2011) until recently. For each country, official national online databases on authorised plant protection products (Table 10) were screened for authorised biocontrol active substances:

Country	Official source / website	Reference date
France	e-phy database of the Ministry of Agriculture & Fisheries	31/8/2009
France	http://e-phy.agriculture.gouv.fr	51/8/2009
	Online-Datenbank Pflanzenschutzmittel of the Federal Office of Consumer	
Cormony	Protection and Food Safety (BVL)	12 /8/2009
Germany	http://www.bvl.bund.de/DE/04_Pflanzenschutzmittel/01_Aufgaben/02_Zulassung	12/0/2009
	PSM/01_ZugelPSM/01_OnlineDatenbank/psm_onlineDB_node.html	
	Registro de productos Fitosanitarios of the Ministerio de Ambiente y Medio Rural	
Spain	y Marino	
	http://www.mapa.es/es/agricultura/pags/fitos/registro/menu.asp	
	Plant protection index ("Pflanzenschutzmittelverzeichnis") of the Federal Office	
Switzerland	for Agriculture (BWL, Fachbereich Pflanzenschutzmittel)	31/7/2009
	http://www.psa.blw.admin.ch/index_de_5_2_A.htm	
United	Pesticides Register of UK approved products under the responsibility of the	
Kingdom	Chemicals Regulation Directorate Pesticides	4/2009
Kinguolli	https://secure.pesticides.gov.uk/pestreg/ProdSearch.asp	

 Table 10:
 Consulted sources of information on authorized biocontrol plant protection products in five European countries:

The survey was limited to uses concerning seven crops or cropping groups which are subject to ENDURE case studies: pomefruit (apples and pears), grapevine, cereals, rape, maize, potatoes and tomatoes (greenhouse and field), the latter being extended to other vegetables where deemed of interest. Country lists of representative products (generally up to two) were created and sorted according to uses in crops, target pests and pathogens were identified by English and scientific names wherever possible.

## **Biocontrol substances registered on Annex 1 of the EU (Pesticides Database)**

The complete list compiled from data retrieved in April 2009 in the EU Pesticides Database is presented in Appendix 10. Excerpts concerning the four categories of substances compatible with biological control are presented in Table 11.

## **Botanicals**

Botanicals are plant-substances resulting from simple processing e.g. pressing or from extraction. By extension the definition applies to a small numbers of compounds or even single ones extracted from plants and purified e.g. laminarine.

Fourteen botanicals have been identified (Table 11) including two borderline cases for which single molecules identical to naturally occurring substances have been synthesised.

- Four botanicals are authorised as repellents only: Extract from the tea tree, garlic extract, clove oil (plant oils) and pepper.
- Six botanicals enter into the category of plant growth regulators.
- The phytohormones gibberellic acid and gibberelline are botanicals produced in fermenters acting on plant growth. Spearmint oil and sea-alga extract are listed for their effect on plant growth as well.
- The phytohormone ethylene is naturally present in plants and in soil and can be included here although it is typically produced in the petrochemical industry by steam cracking.
- Carvone is a terpene produced by aromatic plants in particular by the mint. It can also be classified among the botanicals. To obtain a pure grade it is generally synthesised. In plant protection it is used as a growth regulator.
- Laminarin is extracted from sea weed and is classified as elicitor. Rape seed oil enters into the category of insecticides/acaroids. Citronella oil is the only BCA approved as herbicide.
- Pyrethrins are extracted from Pyrethrum flowers, from cultivars of *Chrysanthemum cinerariaefolium*. By their origin they are botanicals but their structures are analogous and their properties are similar to those of synthetic pyrethroids. Due to their mode of action which is analogous to conventional insecticides and their toxicity for aquatic and other non target organisms, they are not typical biological substances although they are accepted in organic farming.

Substance	Category <sup>1, 2</sup>	List <sup>3</sup>	Inclusion Date	Expiry Date	Legislati on
Botanicals					
Extract from tea tree	RE	A 4	01/09/2009	31/08/2019	2008/127
Garlic extract	RE	A 4	01/09/2009	31/08/2019	2008/127
Gibberellic acid	PG	A 4	01/09/2009	31/08/2019	2008/127
Gibberellin	PG	A 4	01/09/2009	31/08/2019	2008/127
Laminarin	EL	С	01/04/2005	31/03/2015	05/3/EC
Pepper	RE	A 4	01/09/2009	31/08/2019	2008/127
Plant oils / Citronella oil	HB	A 4	01/09/2009	31/08/2019	2008/127
Plant oils / Clove oil	RE	A 4	01/09/2009	31/08/2019	2008/127
Plant oils / Rape seed oil	IN, AC	A 4	01/09/2009	31/08/2019	2008/127
Plant oils / Spearmint oil	PG	A 4	01/09/2009	31/08/2019	2008/127
Sea-algae extract (formerly sea-algae extract and	PG	A 4	01/09/2009	31/08/2019	2008/127
seaweeds)					
Botanicals copied by synthesis (s) or excluded (e)					
Carvone (s)	PG	С	01/08/2008	31/07/2018	<u>2008/44/</u> <u>EC</u>
Ethylene (s)	PG	A 4	01/09/2009	31/08/2019	2008/127
Pyrethrins (e)	IN	A 4	01/09/2009	31/08/2019	2008/127
Microbials					
Ampelomyces quisqualis strain AQ10	FU	С	01/04/2005	31/03/2015	05/2/EC
Bacillus subtilis str. QST 713	BA, FU	С	01/02/2007	31/01/2017	07/6/EC
Bacillus thuringiensis subsp. aizawai (ABTS-1857 and GC-91)	[IN]	A 4	01/01/2009	31/12/2018	2008/113
Bacillus thuringiensis subsp. israelensis (AM65-52)	[IN]	A 4	01/01/2009	31/12/2018	2008/113
Bacillus thuringiensis subsp. kurstaki (ABTS 351, PB	[IN]	A4	01/01/2009	31/12/2018	2008/113
54, SA 11, SA12 and EG 2348)					
Bacillus thuringiensis subsp. tenebrionis (NB 176)	[IN]	A 4	01/01/2009	31/12/2018	2008/113
Beauveria bassiana (ATCC 74040 and GHA)	IN	A 4	01/01/2009	31/12/2018	2008/113
Coniothyrium minitans	FU	С	01/01/2004	31/12/2013	03/79/EC
Cydia pomonella granulosis virus (CpGV)	[IN]	A 4	01/01/2009	31/12/2018	2008/113
Gliocladium catenulatum strain J1446	FU	С	01/04/2005	31/03/2015	05/2/EC
Lecanicillimum muscarium (Ve6) (former Verticillium lecanii)	IN	A 4	01/01/2009	31/12/2018	2008/113
Metarhizium anisopliae (BIPESCO 5F/52)	IN	A 4	01/01/2009	31/12/2018	2008/113
Paecilomyces fumosoroseus Apopka strain 97	[IN]	С	01/07/2001	30/06/2011	01/47/EC
Paecilomyces lilacinus	[IN]	C	01/08/2008	31/07/2018	<u>2008/44/</u> EC
Phlebiopsis gigantea (several strains)	FU	A 4	01/01/2009	31/12/2018	2008/113
Pseudomonas chlororaphis strain MA342	FU	C	01/10/2004	30/09/2014	04/71/EC
Pythium oligandrum (M1)	FU	A 4	01/01/2009	31/12/2018	2008/113
Spodoptera exigua nuclear polyhedrosis virus	FU	C	01/12/2007	30/11/2017	07/50/EC
Streptomyces K61 (K61) (formerly Streptomyces	FU	A 4	01/01/2009	31/12/2018	2008/113
griseoviridis)	10		01/01/2009	51/12/2010	2000/112
<i>Trichoderma aspellerum</i> (ICC012) (T11) (TV1) (formerly <i>T. harzianum</i> )	FU	A 4	01/01/2009	31/12/2018	2008/113
<i>Trichoderma atroviride</i> (IMI 206040) (T 11) (formerly <i>Trichoderma harzianum</i> )	FU	A 4	01/01/2009	31/12/2018	2008/113
Trichoderma gamsii (formerly T. viride) (ICC080)	FU	A 4	01/01/2009	31/12/2018	2008/113
Trichoderma harzianum Rifai (T-22) (ITEM 908)	FU	A 4	01/01/2009	31/12/2018	2008/113
Trichoderma narzanam Khar (1-22) (11EM 908) Trichoderma polysporum (IMI 206039)	FU	A 4	01/01/2009	31/12/2018	2008/113
Verticillium albo-atrum (WCS850) (formerly	FU	A 4	01/01/2009	31/12/2018	2008/113
Verticillium dahliae)	10	114	01/01/2007	51/12/2010	2000/115

# Table 11: Active substances suitable for biological control listed on Annex I of 91/414/EEC (EU<br/>Pesticide Database) - Status on 21 April 2009

# Table 11 (continued)

Other Natural					
Abamectin (aka avermectin)	AC, IN	A 3	01/01/2009	31/12/2018	2008/107
Acetic acid	HB	A 4	01/09/2009	31/08/2018	2008/127
Aluminium silicate (aka kaolin)	RE	A 4	01/09/2009	31/08/2019	2008/127
Blood meal	RE	A 4	01/09/2009	31/08/2019	2008/127
Carbon dioxide	IN, RO	A 4	01/09/2009	31/08/2019	2008/127
Fat distilation residues	RE	A 4	01/09/2009	31/08/2019	2008/127
Ferric phosphate	MO	С	01/11/2001	31/10/2011	01/87/EC
Kieselguhr (diatomaceous earth)	IN	A 4	01/09/2009	31/08/2019	2008/127
Milbemectin	IN, AC	С	01/12/2005	30/11/2015	05/58/EC
Quartz sand	RE	A 4	01/09/2009	31/08/2019	2008/127
Spinosad	IN	С	01/02/2007	31/01/2017	07/6/EC
Other Natural, produced by synthesis				I	
Benzoic acid	BA, FU, OT	С	01/06/2004	31/05/2014	04/30/EC
Potassium hydrogen carbonate	FU	A 4	01/09/2009	31/08/2019	2008/127
Urea	IN	A 4	01/09/2009	31/08/2019	2008/127
Other Natural, fatty acid					
Capric acid (CAS 334-48-5)	IN, AC, HB, PG	A 4	01/09/2009	31/08/2019	2008/127
Caprylic acid (CAS 124-07-2)	IN, AC, HB, PG	A 4	01/09/2009	31/08/2019	2008/127
Fatty acids C7 to C20	IN, AC, HB, PG	A 4	01/09/2009	31/08/2019	2008/127
Fatty acids C7-C18 and C18 unsaturated potassium	IN, AC, HB, PG	A 4	01/09/2009	31/08/2019	2008/127
salts (CAS 67701-09-1)					
Fatty acids C8-C10 methyl esters (CAS 85566-26-3)	IN, AC, HB, PG	A 4	01/09/2009	31/08/2019	2008/127
Lauric acid (CAS 143-07-7)	IN, AC, HB, PG	A 4	01/09/2009	31/08/2019	2008/127
Methyl decanoate (CAS 110-42-9)	IN, AC, HB, PG	A 4	01/09/2009	31/08/2019	2008/127
Methyl octaonate (CAS 111-11-5)	IN, AC, HB, PG	A 4	01/09/2009	31/08/2019	2008/127
Oleic acid (CAS 112-80-1)	IN, AC, HB, PG	A 4	01/09/2009	31/08/2019	2008/127
Pelargonic acid (CAS 112-05-0)	IN, AC, HB, PG	A 4	01/09/2009	31/08/2019	2008/127
Other Natural, repellent					•
Calcium carbonate	RE	A 4	01/09/2009	31/08/2019	2008/127
Limestone	RE	A 4	01/09/2009	31/08/2019	2008/127
Methyl nonyl ketone	RE	A 4	01/09/2009	31/08/2019	2008/127
Sodium aluminium silicate	RE	A 4	01/09/2009	31/08/2019	2008/127
Repellents by smell/Fish oil	RE	A 4	01/09/2009	31/08/2019	2008/127
Repellents by smell/Sheep fat	RE	A 4	01/09/2009	31/08/2019	2008/127
Semiochemical					
(Z)-13-Hexadecen-11yn-1-yl acetate	AT	A 4	01/09/2009	31/08/2019	2008/127
(Z,Z,Z,Z)-7,13,16,19-Docosatetraen-1-yl isobutyrate	AT	A 4	01/09/2009	31/08/2019	2008/127
Ammonium acetate	AT	A 4	01/01/2009	31/12/2018	2008/127
Hydrolysed proteins	IN	A 4	01/09/2009	31/08/2019	2008/127
Putrescine (1,4-Diaminobutane)	AT	A 4	01/09/2009	31/08/2019	2008/127
Trimethylamine hydrochloride	AT	A 4	01/09/2009	31/08/2019	2008/127
Straight Chain Lepidoptera Pheromones	AT	A 4	01/09/2009	31/08/2019	<u>2008/127</u>

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#### Table 11 (continued)

Semiochemical / SCLP					
(2E, 13Z)-Octadecadien-1-yl acetate	AT	A 4	01/09/2009	31/08/2019	2008/127
(7E, 9E)-Dodecadien 1-yl acetate	AT	A 4	01/09/2009	31/08/2019	2008/127
(7E, 9Z)-Dodecadien 1-yl acetate	AT	A 4	01/09/2009	31/08/2019	2008/127
(7Z, 11E)-Hexadecadien-1-yl acetate	AT	A 4	01/09/2009	31/08/2019	2008/127
(7Z, 11Z)-Hexadecdien-1-yl acetate	AT	A 4	01/09/2009	31/08/2019	2008/127
(9Z, 12E)-Tetradecadien-1-yl acetate	AT	A 4	01/09/2009	31/08/2019	2008/127
(E)-11-Tetradecen-1-yl acetate	AT	A 4	01/09/2009	31/08/2019	2008/127
(E)-5-Decen-1-ol	AT	A 4	01/09/2009	31/08/2019	2008/127
(E)-5-Decen-1-yl-acetate	AT	A 4	01/09/2009	31/08/2019	2008/127
(E)-8-Dodecen-1-yl acetate	AT	A 4	01/09/2009	31/08/2019	2008/127
(E,E)-8,10-Dodecadien-1-ol	AT	A 4	01/09/2009	31/08/2019	2008/127
(E/Z)-8-Dodecen-1-yl acetate	AT	A 4	01/09/2009	31/08/2019	2008/127
(Z)-11-Hexadecen-1-ol	AT	A 4	01/09/2009	31/08/2019	2008/127
(Z)-11-Hexadecen-1-yl acetate	AT	A 4	01/09/2009	31/08/2019	2008/127
(Z)-11-Hexadecenal	AT	A 4	01/09/2009	31/08/2019	2008/127
(Z)-11-Tetradecen-1-yl acetate	AT	A 4	01/09/2009	31/08/2019	2008/127
(Z)-13-Octadecenal	AT	A 4	01/09/2009	31/08/2019	2008/127
(Z)-7-Tetradecenal	AT	A 4	01/09/2009	31/08/2019	2008/127
(Z)-8-Dodecen-1-ol	AT	A 4	01/09/2009	31/08/2019	2008/127
(Z)-8-Dodecen-1-yl acetate	AT	A 4	01/09/2009	31/08/2019	2008/127
(Z)-9-Dodecen-1-yl acetate	AT	A 4	01/09/2009	31/08/2019	2008/127
(Z)-9-Hexadecenal	AT	A 4	01/09/2009	31/08/2019	2008/127
(Z)-9-Tetradecen-1-yl acetate	AT	A 4	01/09/2009	31/08/2019	2008/127
Dodecyl acetate	AT	A 4	01/09/2009	31/08/2019	2008/127
Tetradecan-1-ol	AT	A 4	01/09/2009	31/08/2019	2008/127

<sup>1</sup> AC=acaricide, AT= attractant, BA=bactericide, EL=elicitor, FU=fungicide, HB=herbicide, IN=insecticida, MO=molluscicide, NE=nematicide, PA=Plant Activator, PG=Plant Growth, RE=repellent, RO=rodenticide.

<sup>2</sup> Category in [] added by author

<sup>3</sup> A: Existing active substances divided into four lists for phased evaluations; C: New active substances

#### Micro-organisms

The term micro-organism is defined in regulation (EC) No 1107/2009: 'micro-organisms' means any microbiological entity, including lower fungi and viruses, cellular or non-cellular, capable of replication or of transferring genetic material.. This definition applies to, but is not limited to, bacteria, fungi, protozoa, viruses and viroids. It does not include multicellular organisms, such as nematodes or insects.

Twenty five microbial species are included in annex I, some of which are represented by several strains. Six bacterial (sub)species (*Bacillus subtilis*, *Pseudomonas chlororaphis* and four subspecies of *Bacillus thuringiensis*) and two virus species (*Cydia pomonella* Granulose Virus and *Spodoptera exigua* NPV) are included. All B.t. subspecies and viral agents are approved for insect control. *Pseudomonas* is approved for fungicidal seed treatments and *Bacillus subtilis* can be used against plant pathogenic fungi and bacteria. Seventeen fungal agents belonging to twelve genera are listed, *Trichoderma* being represented by five species. *Beauveria bassiana*, *Lecanicillimum muscarium* and *Metarhizium anisopliae* are approved for use as insecticides, the other fungal agents for use against fungal diseases.

#### Semiochemicals (attractants)

Semiochemicals are chemical substances such as pheromones, kairomones and allomones that act to modify the behaviour of pests or their natural enemies.

In the table based on the EU Pesticides Database, Straight Chain Lepidopteran Pheromones (SCLP) are highlighted in green, non-SCLP-pheromones in light cyan and other attractants (including hydrolysed proteins) are highlighted in yellow. There is one repellent which is marked in light red.

SCLPs are included in annex I as a group but 25 compounds of this group are also listed individually. In the inclusion directive 2008/127/EC, some molecules are mentioned three times, as an individual substance, in a blend of the same type, e.g. acetates and in mixed blends, e.g. alcohols and acetates. Often single SCLP compounds show attraction to one or more moth species and typically a combination of two or more of these compounds in a precise ratio enhances the attraction and the specificity. Thus SCLPs should be considered as a whole group and it must not be concluded that each compounds stands for one species.

The SCLPs listed individually are typical examples found in the pheromone blends of moth pest species currently of economic importance. A large variety of compounds and isomers, an estimated number of about 300 identified molecules, used by Lepidopterans are not listed here. They differ in carbon chain length, in the number of double bonds and/or their positions and in their chemical functional group (alcohol, acetate or aldehyde). SCLPs can be used for mass trapping, mating disruption or in attract and kill devices (A&K) or formulations. When associated with an insecticide, i.e in A&K products, attractants do not need to be included in annex I.

Two non SCLP pheromones, (Z)-13-Hexadecen-11yn-1-yl acetate and (Z,Z,Z,Z)-7,13,16,19-Docosatetraen-1-yl isobutyrate, as well as four semiochemicals other than pheromones attractive to different fly (Diptera) species are listed in the EU Pesticides Database: Ammonium acetate, hydrolysed proteins, putrescine (1,4-diaminobutane) and Trimethylamine hydrochloride.

#### Other Plant Protection substances of natural origin

This group has been created for the purpose of the survey. It includes mineral substances as well as substances produced by or derived from animals or from micro-organisms. Thus very diversified substances and products like limestone powder, kaolin as well as diatomaceous earth (Kieselguhr), fatty acids and their derivates (e.g. soaps) can be found in this group. Not all substances of this group do meet the expectation of low non-target toxicity and low environmental impact.

Some active substances included in annex I are produced by micro-organisms. Spinosad which is produced by the bacterium *Saccharopolyspora spinosa* finds its place here; it is accepted for organic farming. Milbemectin is a mixture of natural compounds (milbemycins) isolated from fermentation broth of the fungus *Streptomyces hygroscopicus* subsp. *aureolacrimosus*. The substance is active against insects of different families and a large range of mites. Abamectin contains avermeetins which are biosynthesised by *Streptomyces avermitilis*. The substance shows very high toxicity in Mammals and in aquatic organisms. Milbemectin and abamectin are not authorised in organic crop protection.

Potassium hydrogen carbonate is a slightly basic substance used for its fungicidal properties. The US FDA considers this substance as GRAS (Generally Recognised as Safe). Six natural substances are specifically marked in the EU List, they are used as animal repellents: three are minerals (Calcium carbonate, limestone, sodium aluminium silicate), two are of animal origin (fish oil and sheep fat) while methyl nonyle ketone is either produced by synthesis or extracted from plant oils (rue). The latter repellent acts by its strong odour. It is naturally present in some edible crops and spices.

#### Limit cases and exclusions

With regards to their (eco)toxicological profile and environmental impact neither sulphur and its derivates (iron sulphate) nor cupric compounds i.e. Bordeaux mixture, copper hydroxide, copper oxichloride and cuprous oxide are considered here as typical biological substances although they might be accepted in organic agriculture.

Tall oils (crude or pitch) are a by-product in the Kraft process used in the paper industry. Thus they are substances resulting from a chemical process and are classified as chemicals here. Calcium carbide is produced from lime and coke in electric arc furnaces. It is fitted among chemicals but is used as a repellent like some other minerals. 1-Methyl-cyclopropene is an inhibitor of the effects of

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the phytohormone ethylene and is mainly used to conserve cut flowers. It is placed among the chemicals.

#### Uses of biocontrol products in five European countries

#### Registered plant protection substances

In each country all BCAs authorised for uses in seven crops or cropping groups were identified. Lists of representative products (generally up to two) were created and sorted according to uses in crops: pomefruit (apples and pears), vine, cereals, rape, maize, potatoes and tomatoes (greenhouse and field), the latter was extended to other vegetables where deemed of interest.

In **France** twelve different microbial BCA species (or sub-species in the case of *Bacillus thuringiensis*) are authorised among which two species, *Beauveria tenella* and *Candida oleophila* are not yet included in 91/414 Annex I. Only four botanical active substances are authorised, including rotenon (EU non-inclusion decision in April 2008 but temporary authorisation in FR) and pyrethrum which were excluded from our survey. Fenugreek extracts benefited from a specific French approach to plant extracts under former national rules, while EU approval was given in 2010, after the survey. Laminarin is included in Annex I. Five Straight Chain Lepidopteran Pheromones (SCLP) blends or associations (one just specifying minor components used for the single target codling moth) are registered for mating disruption in orchards or vineyard.

In **Germany** nine microbial BCAs are authorised in Plant Protection Products (all included). Only four botanical substances are listed for plant protection, two of which are included in Annex I (pyrethrins and rape seed oil), two are not (azadirachtin and lecithin). Three different SCLP associations are authorised for mating disruption against Codling Moth or Vine Moths.

For Germany only fully registered BC products according to the rules of the PPP directive were included in the survey. As a consequence, plant strengtheners authorised according to the Federal Plant Protection Act §§ 31ff were excluded. Plant Strengtheners can avoid the EU procedures and requiremments for plant protection products but they must not claim specific protective properties either.

In **Spain** ten microbial BCAs are authorised, all of which are included in EU Annex 1. Only three botanical substances could be identified: Pyrethrins and rotenon which are excluded from the survey and Azadirachtin (Neem extract) which was re-included in EU Annex I in 2011. The plant growth regulators gibberellinic acid/gibberellin are not explored in the survey. Only four SCLP associations are authorised for mating disruption in vine and orchards including two for oriental fruit moth and peach twig borer typical for peach orchards.

In **Switzerland** twelve different microbial BCA species (or sub-species in the case of *Bacillus thuringiensis*) are authorised, among which is one species not included in 91/414 Annex I: *Beauveria brognartii*. Eleven botanicals are approved, among which the insecticides Pyrethrum (included in EU Annex I) and rotenon (rejected from Annex I) have been excluded from the survey because of their toxicological profile. The plant growth regulators gibberellic acid and gibberellin were also excluded from the survey. Five substances not included in EU Annex I are authorised: Azadirachtin (Neem extract), fennel oil, lecithin, mustard powder and Quassia extract. An impressive number of semiochemicals, eleven different SCLP associations are authorized for mating disruption allowing the control a large variety of moths in orchards (including one association of 8 compounds against five different species) and vineyards. This can be related to the facilitated approval of pheromone products in Switzerland.

In the **UK** eight microbial BCAs are approved but only a single botanical (Laminarin, EU approved) and a single pheromone blend (for codling moth). No biological plant protection products are available for use in grapevine, rape, maize or potatoes. With regard to the global availability of biological control products in the different crops, pomefruit, vegetables and vine are generally in a better position than arable crops in the countries included in the survey. In the UK e.g. only

laminarin is available on wheat and cereals, and no biological plant protection products are registered for rape, maize or potatoes.

None of the EU Member States covered in the present survey shows such a variety of BCAs as Switzerland where we find the largest numbers of microbials, botanicals and pheromone blends authorised in the crops subject of the inquiry. Only France reaches the number of twelve microbial BCAs in registered products. The privileged situation in the Helvetic Confederation can be explained by the flexible regulatory approach of the competent authorities in the past, until the progressive implementation of EU directive 91/414/EEC and the related framework, as well as the sustained support by experts in confederal agronomic institutes.

#### Invertebrate biocontrol agents

Invertebrate biocontrol agents (BCAs) used in the five European countries of this survey are listed in Appendix 11.

In **France** invertebrate BCAs cannot be registered and they do not yet need to be formally declared, but a law passed on 12<sup>th</sup> July 2010 created the basis to establish rules governing the introduction into the environment of non-indigeneous macro-organisms useful to plants. Procedures and requirements for authorisations which will also cover non-indigeneous beneficial are expected to be set up for the in the coming months. The list provided in the present survey is based on the voluntary declarations to ACTA by the producers wishing to have their beneficials published in the non-official Index Phytosanitaire.

Invertabrate BCAs must be registered in **Germany**. An official list which is regularly updated is published by the Julius Kühn Institute.

In **Spain** companies which are responsible of commercialisation of IBCAs must give information to the Ministry of Agriculture to allow the inscription into a register before commercialisation (Orden APA/1470/2007). This information given is about name of commercial product, identification of the organism, the manufacturer, the company responsible for commercialisation. Another law (43/2002; 20<sup>th</sup> of November 2002) covers the introduction of exotic organisms (article 44).

In **Switzerland** invertebrate BCAs must be formally approved by the BLW (Bundesamt für Landwirtschaft) and they are listed together with the plant protection products.

In the **United Kingdom** no authorisation is required to release indigenous beneficials but the import (and release) of non indigenous species must be approved by the Advisory Committee for the Release of Exotics (ACRE acting under DEFRA).

# **Chapter 5**

# Identified difficulties and conditions for field success of biocontrol. 1. Regulatory aspects

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## **Objectives**

The objective of the work was to identify typical hurdles for the placing of biological plant protection products on the market experienced by biocontrol industry or evaluators in the recent past under the European directive 91/414/EEC. In parallel, we examined the new regulation (No 1107/2009/EC of the European Parliament and of the Council of 21 October 2009) concerning the placing of plant protection products on the market and repealing Council Directives 79/117/EEC and 91/414/EEC and the new directive (N° 2009/128/EC of the European Parliament and of the Council of 21 October 2009) establishing a framework for Community action to achieve the sustainable use of pesticides. These two texts were examined for provisions creating new opportunities for the approval biocontrol agents, their placing on the market and use. In fine, it was the intent to establish a dialogue with EU regulators and evaluators in European institutions, i.e. in the European Commission and in the European Food Safety Agency (EFSA) and to seek solutions in common for the problems encountered.

#### Working method

An *ad hoc* group of representatives from the biocontrol industry and INRA called "Regulatory Review Team" was set up. Two full-day working sessions were organised in which regulatory experts identified difficulties and questions but also described positive experience and perspectives.

The work of the Regulatory Review Team active under Reasearch Activity RA4.3 of the ENDURE network was then summarised and reported in a meeting of a delegation of ENDURE partners (IBMA, INRA and ACTA) with representatives of the European Commission (DG SANCO, DG Agriculture, DG Research) and the EFSA in Brussels (24 September 2009).

#### Results

A PowerPoint presentation entitled "*Gaps - Problems - Opportunities for BCAs in E.U. Regulation - From Past to Future*" was prepared for the ENDURE – Commission meeting, with inputs on general regulatory issues, micro-organisms, straight chain lepidopteran pheromones and botanicals. In this document, two key issues related to directive 2001/36/EC annex II B which fixed requirements for **microbial active substances** were highlighted. Readers may note that since 14<sup>th</sup> June 2011 Regulation (EU) No 544/2011 implements these data requirements unchanged to reg.

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(EC) No 1109/2009. Tests suggested by evaluation experts and intended to establish the genetic stability of a strain do not reflect practical conditions, while in the case of potential microbial contaminants no European reference list is available. The incidence of many pathogens can be excluded by production methods or the geographic location of production sites. Tolerance limits for contamination levels could take into consideration thresholds used in food industry, application levels for the microbial product and naturally occurring background levels. The two issues presented here but also other examples put forward to the Regulatory Review Team lead to the statement that "not all the studies or tests that can be performed for microbials will necessarily yield relevant data".

The most important experience with semiochemicals was made during the on-going reassessment of **Straight Chain Lepidopteran Pheromones** (SCLPs), which were supported by an IBMA Task Force. Regulators and evaluators were flexible in accepting a single common dossier for all compounds notified but although an OECD guidance document recommends data waiving for numerous SCLP requirements, the Rapporteur Member State insisted that all existing data and study reports on all compounds be submitted on the grounds that the requirements of the directive are superior to the guidance document recommendations. So far, the re-assessment procedure resulted in the inclusion with postponed peer review of SCLPs as a group, but 25 substances are also listed individually. New substances can be included in a simplified procedure provided that the applicant has access to the existing dossier. Remaining questions include what industry input will be required during the peer review by EFSA, the E.U. status of a revised OECD guidance document for semiochemicals other than SCLPs, the decision if MRLs are required for sprayable SCLP formulations, and equivalence criteria for SCLP substances. It was also noted that under the Biocidal Product Directive, rules and fees applied to SCLPs created an economic hurdle which resulted in the submission of a dossier for only one compound.

**Extracts from plants** - as long as not purified - consist of mixtures of molecules while data requirements of directive 91/414/EEC maintained under new regulation (EC) No 1107/2009 are basically designed for defined single substances. Thus those requirements often do not fit for mixtures of several substances. It must be decided if the most "active" substance, the one with the highest content in the extract or the whole extract shall be used in studies required for different sections of a dossier i.e. for data on physical-chemical properties, metabolism, toxicology, residues, environmental fate and behaviour, and which data shall be used in risk assessment. While the whole extract can be recommended for use in toxicity studies, it is not convenient for residue, metabolism or environmental studies because in practice it is generally not possible to determine the fate of all compounds contained in an extract. Questions asked by evaluators from several Member States after the issuing of a draft assessment report for Neem extract and its lead substance Azadirachtin A illustrate the difficulties experienced by an applicant in the evaluation process for a botanical.

**Regulation** (EC) No 1107/2009 concerning the placing of plant protection products on the market provides for a specific status for "low risk active substances" (article 22). Many biocontrol substances can be expected to qualify for this new category but one exclusion criterion, the half-life in soil, may cause problems for microbial active substances unless it is clearly limited to chemicals. A full set of data is required to gain the status of low risk active substance but products containing them exclusively and without co-formulants of concern will benefit from reduced dossier requirements and time lines for approval. Micro-organisms, plant extracts or other natural substances may also meet the criteria for "Basic substances" provided for in article 23 but the discussion in the ENDURE-Commission meeting made it clear that this category is without interest for manufacturers who intend to market their substances for plant protection. It was noted that the new regulation does not provide for generic waivers i.e. for justifications of non submission of data or exemptions from requirements for groups of substances or products.

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In the **sustainable use directive 2009/128/EC** a number of provisions in favour of biological pest control measures or non-chemical methods have been identified. The new regulation also mentions in recital 35 that priority should be given to "non-chemical and natural alternatives wherever possible" but since the definition of non-chemical methods refers to "physical, mechanical or biological pest control" and does not specifically mention microbials, semiochemicals, botanicals or other natural substances with non-toxic mode of action it must be clarified how those groups are covered by the definition.

# Conclusion

In the meeting between the ENDURE delegation and representatives of the European Commission, the need for discussions between regulators, evaluators and industry about requirements especially those relevant for microbial and botanical substances was recognised. Article 77 of the new plant protection product regulation authorises the Commission to "*adopt or amend technical and other guidance documents e.g. explanatory notes or guidance documents on the content of the application concerning micro-organisms, pheromones and biological products.*" Thus at least part of the problems experienced by applicants can be addressed in guidance documents. Industry representatives and companies directly concerned by evaluations or reviews of biocontrol agents should enter into discussions with evaluators (EFSA or Competent Authorities in Member States) without forgetting the leading role of the Commission. Industry should fix priorities, prepare rationales and make substantiated proposals dealing with data requirements considered inappropriate, unnecessary or unrealistic.

# **Chapter 6**

# **Identified difficulties and conditions for field success of biocontrol. 2. Technical aspects: factors of efficacy**

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#### **Quality of the BCAs formulations**

Numerous investigations on the development of biopesticides have been initiated as legislation and government policy have demanded less reliance on chemical pesticides and greater adoption of IPM. In Europe, some countries have set goals of reducing pesticide use by 50%. Successes have been achieved through better timing of applications, so that lower dosages are effective and substituting less hazardous and more active materials, to reduce the number of applications.

Biopesticides are distinguished from conventional chemical pesticides as many are very selective and are non-toxic towards non-target organisms. While biopesticides are likely to be less harmful to the environment than the conventional ones, care needs to be taken that wastage is minimised, by selecting the most appropriate droplet spectrum. A disadvantage of biological agents relative to chemicals, is that many are not sufficiently persistent and are relatively slow acting; therefore, research has been directed at extending the period of activity. However, some such agents may persist in the field or the forest for many months, and a risk–benefit analysis should be performed to establish their environmental acceptability.

Transition from the optimised conditions of a laboratory experiment to the harsh conditions experienced in the field has so far proved more difficult for application of biopesticides in contrast to chemicals. This has undoubtedly been due to lack of investment in the development of effective formulations and delivery systems, in order to commercialise more potential biopesticides. The relatively small effort invested in target-specific sprayers, compared with the investment in laboratory studies, has led to unbalanced development, and exemplifies the need for closer integration between formulation and engineering research. The challenge is to get effective formulations so that biological control agents can be easily applied by farmers.

# A good example, the case of *Trichoderma*: direct and indirect mode of action against plant pathogens

*Trichoderma* species have long been recognized as biological control agents (BCAs) for the control of plant disease and for their ability to increase plant growth and development. They are widely used in agriculture, and some of the most useful strains demonstrate a property known as 'rhizosphere competence', the ability to colonize and grow in association with plant roots (Harman 2000). Much of the known biology and many of the uses of these fungi have been documented recently (Harman *et al.* 2004a; Kubicek *et al.* 1998; Perello *et al.* 2009). The taxonomy of this fungal genus is continually being revised, and many new species are being described (Komon-Zelazowska *et al.* 2007; Kubicek *et al.* 2008; Overton *et al.* 2006; Samuels 2006; Samuels and

Ismaiel 2009). The mechanisms that *Trichoderma* uses to antagonize phytopathogenic fungi include competition, colonization, antibiosis and direct mycoparasitism (Harman 2006, 2011; Howell 2003). This antagonistic potential serves as the basis for effective biological control applications of different *Trichoderma* strains as an alternative method to chemicals for the control of a wide spectrum of plant pathogens (Harman *et al.* 1991; Lorito *et al.* 2010).

The colonization of the root system by rhizosphere competent strains of *Trichoderma* results in increased development of root and/or aerial systems and crop yields (Bae *et al.* 2011; Chacon *et al.* 2007; Kubicek *et al.* 1998; Yedidia *et al.* 2003). *Trichoderma* has also been described as being involved in other biological activities such as the induction of plant systemic resistance (Shoresh *et al.* 2010; Tucci *et al.* 2011) and antagonistic effects on plant pathogenic nematodes (Jegathambigai *et al.* 2008; Sharon *et al.* 2001).Some strains of *Trichoderma* have also been noted to be aggressive biodegraders in their saprophytic phases, in addition to acting as competitors to fungal pathogens, particularly when nutrients are a limiting factor in the environment (Worasatit *et al.* 1994). These facts strongly suggest that in the plant root environment *Trichoderma* actively interacts with the components in the soil community, the plant, bacteria, fungi, other organisms, such as nematodes or insects, that share the same ecological niche (Lorito *et al.* 2010).

*Trichoderma* spp. are important participants in the nutrient cycle. They aid in the decomposition of organic matter and make available to the plant many elements normally inaccessible. Yedidia *et al.* (2001) noted that the presence of the fungus increased the uptake and concentration of a variety of nutrients (copper, phosphorus, iron, manganese and sodium) in the roots of plants grown in hydroponic culture, even under axenic conditions. These increased concentrations indicated an improvement in plant active-uptake mechanisms. Corn that developed from seeds treated with *T. harzianum* strain T-22 produced higher yields, even when a fertilizer containing 40% less nitrogen was applied, than the plants developed from seed that was not treated with T-22 (Harman 2000). This ability to enhance production with less nitrate fertilizer, provides the opportunity to potentially reduce nitrate pollution of ground and surface water, a serious adverse consequence of large-scale maize culture. In addition to effects on the increase of nutrient uptake and the efficiency of nitrogen use, the beneficial fungi can also solubilize various nutrients in the soil, that would be otherwise unavailable for uptake by the plant (Altomare *et al.* 1999b).

The cross-talk that occurs between the fungal BCA and the plant is important both for identification of each component to one another and for obtaining beneficial effects. Somehow, the plant is able to sense, possibly by detection of the released fungal compounds, that *Trichoderma* is not a hostile presence, therefore the plant defence system is not activated as it is when there is pest attack and the BCA is recognized as a plant symbiont rather than a plant pathogen (Woo and Lorito, 2006). Molecules produced by *Trichoderma* and/or its metabolic activity also have potential for promoting plant growth (Chacón *et al.*, 2007; Vinale *et al.* 2008a; 2008b; Yedidia *et al.* 1999). Applications of *T. harzianum* to seed or the plant resulted in improved germination, increased plant size, augmented leaf area and weight, greater yields (Altomare *et al.* 1999a; Harman *et al.* 2004c, b; Inbar and Chet 1995; Tucci *et al.* 2011; Vinale *et al.* 2008a).

Numerous studies indicated that metabolic changes occur in the root during colonization by *Trichoderma* spp., such as the activation of pathogenesis-related proteins (PR-proteins), which induce in the plant an increased resistance to subsequent attack by numerous microbial pathogens (Table 12)

species	s (11ai illa	in <i>et al.</i> , 2004	fa).		-
Species and strain	Plant	Pathogens	Evidence or effects	Time after application	Efficacy
<i>T. virens</i> G-6, G-6-5 and G-11	Cotton	Rhizoctonia solani	Protection of plants; induction of fungitoxic terpenoid phytoalexins	4 days	78% reduction in disease; ability to induce phytoalexins required for maximum biocontrol activity
T. harzianum T-39	Bean	Colletotrichum lindemuthianum; Botrytis cinerea	Protection of leaves when T-39 was present only on roots	10 days	42% reduction in lesion area; number of spreading lesions reduced
T. harzianum T-39	Tomato, pepper, tobacco, lettuce, bean	B. cinerea	Protection of leaves when T-39 was present only on roots	7 days	25–100% reduction in grey-mould symptoms
T. asperellum T-203	Cucumber	Pseudomonas syringae pv. lachrymans	Protection of leaves when T-203 was present only on roots; production of antifungal compounds in leaves	5 days	Up to 80% reduction in disease on leaves; 100-fold reduction in level of pathogenic bacterial cells in leaves
T. harzianum T-22; T. atroviride P1	Bean	B. cinerea and Xanthomonas campestris pv. phaseoli	Protection of leaves when T-22 or P1 was present only on roots; production of antifungal compounds in leaves	7–10 days	69% reduction in grey-mould ( <i>B. cinerea</i> ) symptoms with T22; lower level of control with P1. 54% reduction in bacterial disease symptoms.
T. harzianum T-1 & T22; T. virens T3	Cucumber	Green-mottle mosaic virus	Protection of leaves when Trichoderma strains were present only on roots	7 days	Disease-induced reduction in growth eliminated
T. harzianum T-22	Tomato	Alternaria solani	Protection of leaves when T-22 was present only on roots	3 months	Up to 80% reduction in early blight symptoms from natural field infection
T. harzianum T-22	Maize	Colletotrichum graminicola	Protection of leaves when Trichoderma strains were present only on roots	14 days	44% reduction of lesion size on wounded leaves; no disease on non-wounded leaves
Trichoderma GT3-2	Cucumber	C. orbiculare, P. syringae pv. lachrymans	Protection of leaves when <i>Trichoderma</i> strains were present only on roots; induction of lignification and superoxide generation	,	59% and 52% protection from disease caused by <i>C. orbiculare</i> or <i>P. syringae</i> , respectively
T. harzianum	Pepper	Phytophthora capsici	Protection of stems when Trichoderma strains were present only on roots; enhanced production of the phytoalexin capsidiol	9 days	~40% reduction in lesion length
T. harzianum NF-9	Rice	Magnaporthe grisea; Xanthomonas oryzae pv. oryzae	Protection of leaves when NF-9 was present only on roots	14 days	34–50% reduction in disease

Table 12:	Evidence for,	and	effectiveness	of,	induced	resistance	in	plants	by	Trichoderma
spec	cies (Harman et	al., 2	2004a).		_			_		

The induction of systemic resistance (ISR) observed *in planta* determines an improved control of different classes of pathogens (mainly fungi and bacteria), which are spatially and temporally distant from the *Trichoderma* inoculation site. This phenomenon has been observed in many plant species, both dicotyledons (tomato, pepper, tobacco, cotton, bean, cucumber) and monocotyledions (corn, rice). For example, *Trichoderma* induces resistance towards *Botrytis* cinerea in tomato, tobacco, lettuce, pepper and bean plants, with a symptom reduction ranging from 25 to 100% (Tucci *et al.* 2011). Moreover, *Trichoderma* determined an overall increased production of defence-related plant enzymes, including various peroxidases, chitinases,  $\beta$ -1,3-glucanases, and the lipoxygenase-pathway hydroperoxide lyase (Harman *et al.* 2004c; Howell *et al.* 2000; Yedidia *et al.* 1999) of *T. harzianum* strain T-39, the active ingredient of the commercial product TricodexTM.

Thus far, *Trichoderma* is able not only to produce toxic compounds with a direct antimicrobial activity against pathogens, but also to generate fungal substances that are able to stimulate the plant to produce its own defence metabolites. In fact, the ability of *T. virens* to induce phytoalexin accumulation and localized resistance in cotton has already been discussed (Hanson and Howell 2004). In cucumber, root colonization by strain T-203 of *T. asperellum* caused an increase in phenolic glucoside levels in the leaves; the aglycones, which are phenolic glucosides with the carbohydrate moieties removed, are strongly inhibitory to a range of bacteria and fungi (Yedidia *et al.* 2003).

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A fundamental part of the *Trichoderma* antifungal capability consists in the production and secretion of a great variety of extracellular cell wall degrading enzymes (CWDEs), including endochitinases,  $\beta$ -N-acetylhexosaminidase (N-acetyl- $\beta$ -D-glucosaminidase), chitin-1,4- $\beta$ -chitobio-sidases, proteases, endo- and exo- $\beta$ -1,3-glucanases, endo  $\beta$ -1,6-glucanases, lipases, xylanases, mananases, pectinases, pectin lyases, amylases, phospholipases, RNAses, DNAses, etc. (Benitez *et al.* 2004; Lorito *et al.* 1998). The chitinolytic and glucanolytic enzymes are especially valuable for their CWDE activity on fungal plant pathogens, hydrolyzing polymers not present in plant tissues (Woo *et al.* 1999). Each of these classes of enzymes contains diverse sets of proteins with distinct enzymatic activities. Some have been purified, characterized and their encoding genes cloned (Ait-Lahsen *et al.* 2001; de la Cruz *et al.* 1992; 1995a; 1995b; Garcia *et al.* 1994; Limon *et al.* 1995; Lora *et al.* 1995; Lorito *et al.* 1993, 1994b; Montero *et al.* 2007; Peterbauer *et al.* 1996; Suarez *et al.* 2004; Viterbo *et al.* 2001, 2002). Once purified, many *Trichoderma* enzymes have shown to have strong antifungal activity against a wide variety of phytopathogens, and they are capable of hydrolyzing not only the tender young hyphal tips of the target fungal host, but they are also able to degrade the hard, resistant conservation structures such as sclerozi.

*Trichoderma* spp. have been widely studied, and are presently marketed as biopesticides, biofertilizers and soil amendments, due to their ability to protect plants, enhance vegetative growth and contain pathogen populations under numerous agricultural conditions (Harman 2000, 2004; Vinale *et al.* 2008b). The commercial success of products containing these fungal antagonists can be attributed to the large volume of viable propagules that can be produced rapidly and readily on numerous substrates at a low cost in diverse fermentation systems. The living microorganisms, conserved as spores, can be incorporated into various formulations, liquid, granules or powder etc., and stored for months without losing their efficacy (Jin *et al.* 1996). To date more than 50 different *Trichoderma*-based preparations are commercialized and used to protect or increase the productivity of numerous horticultural and ornamental crops (Table 13; Lorito *et al.* 2006).

# The case *Trichoderma*: mode of application, persistence on the target and new formulations.

Effectiveness under controlled conditions (even under field conditions) does not necessarily guarantee that the organism will perform successfully; proper formulation is a prime condition for meeting market requirements. For instance an efficient biocontrol agent of soilborne and airborne pathogens must first and foremost protect the young seedling against detrimental attack by infective inoculum. Therefore some factors may be considered:

(a) soil ecosystem factors such as moisture, pH, structure, and temperature;

(b) root colonization capacity;

(c) reasonable shelf life;

(d) efficiency of application of the biocontrol agent in terms of its specific habitat and target (Spiegel and Chet 1998)

Commercial	Biocontrol	Product	Formulation,	Uses - Location,	Uses, Pathogens	Manufacturer/Supplier, Country, Internet Reference
Product	Organism(s)	Туре	Application	Crops	controlled	
Ago Biocontrol <i>Trichoderma</i> 50	T. harzianum	Biological fungicide	n/a	Flowers, vegetables, fruits, other crops	Fusarium, Rhizoctonia, Alternaria, Rosellinia, Botrytis, Sclerotium, Phytophthora spp	Ago Biocontrol, Colombia (http://www.sipweb.org/directorymcp/fungi.html)
<u>Antagon</u>	<i>Trichoderma</i> spp.	Biological fungicide	powder	Horticulture (commercial), parks, recreational areas, sports fields	damping-off diseases	De Ceuster Meststoffen N.V. (DCM), Belgium (http://www.agroBiologicals.com/products/P1609.htm)
<u>Binab T</u>	T. harzianum, T. polysporum	Biological fungicide	Pellets, wettable powder or granules; spray, drench, mixed in soil	Wood products; ornamental, shade, forest trees; greenhouse, nursery, field; cut flowers, potted plants, vegetables, mushrooms, flower bulbs	Wood rots causing internal decay, or originating from pruning wounds; Didymella, Chondrostereum, Heterobasidion, Botrytis, Verticillium, Pythium, Fusarium, Phytophthora, Rhizoctonia	BINAB Bio-Innovation AB, Sweden (http://www.algonet.se/~binab/index2.html); Henry Doubleday Research Association, United Kingdom; Svenska Predator AB, Sweden; E.R. Butts International, Inc., USA
BioFit	T. viride	Biological fungicide	Seed treatment, root/tuber dip, drench; Used alone or in combination with chemicals.	Gram, pepper, groundnut, wheat, potato, ginger, turmeric, peas, matki, mung, urid, tomato, bhindi, onion, other vegetables, grapes.	Pythium, Rhizoctonia, Fusarium, Sclerotium, other root rots; for Botrytis in combination with chemicals	Ajay Bio-tech (India) Ltd., India (http://www.ajaybio.com)
<u>Bio-Fungus</u> (formerly <u>Anti-Fungus),</u> <u>Supresivit</u>	Trichoderma spp.	Biological fungicide	granular, wettable powder, sticks, crumbles; soil incorporation; spray or injection	Flowers, strawberries, trees, vegetables	Sclerotinia, Phytophthora, Rhizoctonia solani, Pythium spp., Fusarium, Verticillium	BioPlant, Denmark (www.bioplant.dk); De Ceuster Meststoffen N.V. (DCM), Belgium

 Table 13:
 Trichoderma-based preparations commercialized for biological control of plant diseases.

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Table 13 (continued): Trichoderma-based preparations commercialized for biological control of plant diseases.

Commercial	Biocontrol	Product	Formulation,	Uses - Location,	Uses, Pathogens controlled	Manufacturer/Supplier, Country, Internet Reference
Product	Organism(s)	Туре	Application	Crops		
Combat	T. harzianum, T. virens (=T. lignorum G. virens), Bacillus subtilis	Biological fungicide	Talc; seed treatment, broadcast, root dip, drench, foliar spray	Grapes, cotton, pulses, tea, potato, tomato, oil seeds, tobacco, spices, cereals, vegetables, horticultural crops	Downy mildew, powdery mildew, die back, <i>Verticillium, Fusarium</i> , Panama wilt; pod, seedling, late blight; root, collar, stem, red, soft, clump, dry, bean, fruit, pod rot; black leg, damping off, abnormal leaf fall, black thread, canker	BioAg Corporation USA (http://www.bioag.com/products.html)
<u>Harzian 20</u> (under	T. harzianum	Biological fungicide	n/a	orchard crops, vineyards	Armillaria spp., Pythium spp., Sclerotinia spp.	Natural Plant Protection (NPP), France (http://www.agroBiologicals.com/products/P1362.htm)
<u>development)</u>						
<u>PlantShield</u>	T. harzianum	Biological fungicide	Granules, wettable powder; soil drench, foliar spray	Greenhouse, flowers, ornamentals, herbs, nursery, vegetable crops; hydroponic, orchard trees	Pythium, Fusarium, Rhizoctonia, Cylindrocladium, Thielaviopsis; suppresses Botrytis	BioWorks, Inc., USA (http://www.bioworksbiocontrol.com)
Primastop	G. catenulatum	Biological fungicide	Powder; drench, spray, irrigation	ornamental, vegetable, tree crops	pathogens causing seed, root, stem rot, wilt disease	Kemira Agro Oy, Finland (http://growhow.kemira- agro.com); AgBio Development Inc.USA
<u>Root Pro,</u> <u>RootProtato</u>	T. harzianum, T. cornedia	Biological fungicide	Powder; spores mixed with growing media	Seedling, rooting stage in nursery; Horticulture - flowers, vegetables, potatoes	Rhizoctonia solani, Pythium spp., Fusarium spp., Sclerotium rolfsii	Mycontrol Ltd., Israel; Efal Agri, Israel ( <u>http://www.efal.com/main.htm</u> , http://www.agroBiologicals.com/company/C1096.htm)

Many preparations have been developed to ensure a good shelf life of the product based on *Trichoderma*. Some of that formulation are stable in terms of pH, that remains constant and low (5.5) during the entire growth period, thus preventing bacterial contamination. Moreover the shelf life of the fungus at 25 °C is 1 year and from 1 to 2 years, the number of colonies-forming-units (CFUs) decreases by one order of magnitude. Many of that formulation have been proven successful in several experiments in the greenhouse and field. The rapid mass production of promising antagonists in the form of spores, mycelia or mixtures of both, has been achieved by liquid-fermentation technology: mass production of biomasses of *T. hamatum*, *T. harzianum*, and *T. viride* was reached by utilizing commercially available, inexpensive ingredients such as molasses, brewer's yeast, cotton seed flour, or corn-steeped liquor.. Other techniques have been employed to improve the delivery of the biocontrol agents. A lignite-stillage (a by-product of sorghum fermentation) carrier system was tested for applying a *T. harzianum* preparation to the soil. Encapsulation of the biocontrol agent in an alginate-clay matrix, using Pyrax as the clay material, improved yield and propagule viability over time.

Pelletized formulations of wheat bran or kaolin clay in an alginate gel containing conidia, chlamydospores or fermentex biomass of several *Trichoderma* isolates revealed increased viability of stored pellets, and the number of CFUs formed after adding these pellets to the soil was comparable to that formed from freshly prepared pellets. These growth media and delivery systems for formulations of biocontrol fungi show promise because they are able to introduce high levels  $(10^{6}-10^{10} \text{ CFU/g})$  of fungi into soils not steamed, fumigated, or treated with other biocides.

To enhance biocontrol efficacy, appropriate introduction of the antagonist into the microenvironment appears to be crucial: formulations have been applied to seedlings prior to planting or to seeds in furrows. Economic considerations have forced biotechnologists to improve the application techniques: seed-coating, a technique involving minimal amounts of inoculum was developed.

Increased biocontrol activity may be achieved by combining two types (or more, if possible) of biocontrol agents, for example combining Trichoderma with a bacterium, or another beneficial fungus. The combined activity of the antifungal compounds produced by both microorganisms could expand the spectrum of pathogens controlled. In fact, in field trials combining T. koningii with certain fluorescent pseudomonads, greater suppression of take-all disease and increased wheat yield were achieved relative to plants treated with T. koningii alone (Duffy et al. 1996). Delivery systems must ensure that biocontrol agents will grow well and achieve their purpose. It is generally recognized that delivery and application processes must be developed on a crop by crop and application by application basis. No general solutions exist, and so biocontrol systems must be developed for each crop. It is very important to use the organism properly and to have appropriate expectations. Any biocontrol organism will be unable to protect seeds as well as chemical fungicides. However, it colonizes roots, increases root mass and health, and consequently frequently provides yield increases, which chemical fungicides applied at reasonable rates cannot do. An effective method of use is to use the biocontrol fungus in conjunction with chemical fungicides. The chemicals provide good short-term seed protection, and the biocontrol fungus provides long-term root protection. As a consequence, yields frequently are increased over use of the chemical alone.

Some experiences evidence that *Trichoderma* spp. is also highly effective when applied to blossoms or fruits for control of *B. cinerea*. Even low levels of the organism applied to strawberry blossoms by bee delivery or by sprays of liquid formulations are effective. For maximum control of the *Botrytis* bunch rot of grape, this initial application needs to be augmented by sprays as fruits mature, and addition of iprodione as a tank mix to this late application appears to have synergistic activity over either the biocontrol agent or the chemical fungicide alone.

**Novel applications of** *Trichoderma* **spp.** *Trichoderma* **spp.** produce a variety of lytic enzymes that have a high diversity of structural and kinetic properties, thus increasing the probability of this fungus to counteract the inhibitory mechanisms used by neighbouring microorganisms. Further,

*Trichoderma* hydrolytic enzymes have been demonstrated to be synergistic, showing an augmented antifungal activity when combined with themselves, other microbial enzymes, PR proteins of plants and some xenobiotic compounds (Fogliano *et al.* 2002; Lorito *et al.* 1994a, 1994b, 1994c, 1996, 1998; Schirmbock *et al.* 1994; Woo *et al.* 2002). In fact, the inhibitory effect of chemical fungicides for the control of the foliar pathogen *B. cinerea* was substantially improved by the addition of minute quantities (10-20 ppm) of *Trichoderma* CWDEs to the treatment mixture (Lorito *et al.* 1994b).

Extensive testing of *T. harzianum* strain T22 conducted for the registration of this biocontrol agent in the USA by the Environmental Protection Agency (EPA) has found that the CWDEs do not have a toxic effect on humans and animals (ED50 and LD50), and that they do not leave residues, but degrade innocuously in the environment. Therefore, these *Trichoderma* hydrolytic enzymes present a novel product for plant disease control based on natural mycoparasitic compounds used by the antagonistic fungi. Single or mixed combinations of CWDEs with elevated antifungal effects, obtained from fermentation in inducing conditions, over-expression of the encoding genes in strains of *Trichoderma*, or heterologous expression of the encoding genes in other microbes are possible alternatives for pathogen control. These natural substances originating from the BCA are an improvement over the use of the living microorganism in the production of commercial formulations because they are easily characterized, resist desiccation, are stable at temperatures up to 60° C, and are active over a wide range of pH and temperatures in the agricultural environment.

The important factors to consider in a commercial bio-formulation are product stability, the capacity to produce consistent results by preserving the characteristics producing the biological effects; the storability of the material, the ability to be conserved in unspecialized conditions similar to those of chemical pesticides; and a reasonable shelf-life or time that the product can be stored and used without compromising the efficacy (Agosin and Aguilera, 1998; Agosin et al., 1997; Powell and Jutsum, 1993). When a formulation contains the living microorganism component, the treatment must consist of stabilizing the viability of the BCA. For liquid formulations this can be achieved by maintaining the product in refrigeration (<10° C) or by freezing in the presence of cryoprotectant substances. However, conservation of a commercial product in these conditions is not economic for maintaining low temperatures or efficient because the liquid is both bulky and heavy, plus it is difficult to sustain these conditions in storage and transportation. In comparison, it is preferable to obtain formulations that contain a dehydrated product, stored as a powder, granule, talc, etc. Some works (Ruocco et al. unp) demonstrated that lyophilisation did not reduce chitinolytic activity and spore vitality when the fermented cultures were treated with compounds that protect the osmotic integrity of the living material such as glycerol. Generally, lyophilisation is the method that best maintains viability, but its cost is very high. At the industrial level and in order to obtain a low-cost product, the methods preferred is spray- or fluidized bed- drying. Many products are obtained by spray-drying, but this method produces a high loss of viability in some microorganisms (observed also in this formulation), due to the thermal treatment.

In spite of the relatively abundant number of patents filed for microbial pesticides, the number of commercial applications has not been as dramatic as expected (Montesinos, 2003). In Europe, the limiting factor for registration, apart from the cost, is undoubtedly the slow process of decision-taking. As an example, the first application for patenting a biopesticide, Paecilomyces fumosoroseus, was submitted to the European Union in 1994 and approved only in 2001. In most cases, excessive specificity is a problem difficult to solve because it is intrinsic to the biological control system. In fact, success depends on three living systems: the pathogen or pest, the BCA and the host plant. Biosafety and environmental concerns are also major limiting factors for microbial pesticide prospects. Furthermore, the registration procedure to approve a biopesticide formulation on the market has not been altered to consider the biological aspects of the product, criteria which are different than those considered for the testing of chemical based products.

## Persistence, physiological stresses, timing and coverage of others biological agents

Other references have been screened for biocontrol agents considering the analysis of:

- persistence on the target,
- resistance to physiological stresses,
- timing and coverage.

*Cladosporium cladosporioides*. The antagonist has been effective in reducing sporulation of *Venturia inaequalis* under orchard conditions. Furthermore, the results of the pre-screening indicate that it is cold and drought tolerant and results of experiments on spore production in solid state fermentation show that mass production is economically feasible. These results have been obtained in a stepwise selection approach (Köhl, 2009, Köhl *et al.*, 2009).

Ulocladium atrum and Gliocladium roseum. Köhl et al., 1998 described the effect of treatments with conidial suspensions of Ulocladium atrum and Gliocladium roseum on leaf rot of cyclamen caused by *Botrytis cinerea* was investigated under commercial greenhouse conditions. Spraying U. atrum (1  $\times$  106 conidia per ml) or G. roseum (2  $\times$  106 conidia per ml and 1  $\times$  107 conidia per ml) at intervals of 2 to 3 weeks during the production period and spraying U. atrum ( $1 \times 106$  conidia per ml) at intervals of 4 to 6 weeks resulted in a significant reduction of natural infections of petioles by B. cinerea. U. atrum or G. roseum (1  $\times$  107 conidia per ml) was as effective as the standard fungicide program. B. cinerea colonized senesced leaves within the plant canopy and infected adjacent petioles and leaves later. The antagonists colonized senesced leaves and reduced B. cinerea development on these leaves. Thus, the inoculum potential on petioles adjacent to necrotic leaf tissues was reduced. The fate of U. atrum conidia on surfaces of green cyclamen leaves during a 70day period after application was studied. The number of conidia per square centimetre of leaf surface remained relatively constant during the entire experiment. Sixty percent of the conidia sampled during the experiments retained the ability to germinate. When green leaves were removed from the plants to induce senescence and subsequently were incubated in a moist chamber, U. atrum colonized the dead leaves. Senesced leaves also were colonized by other naturally occurring fungi including B. cinerea. On leaves treated with U. atrum from all sampling dates, sporulation of B. cinerea was significantly less as compared with the untreated control. Our results indicate that early applications of *U. atrum* before canopy closure may be sufficient to achieve commercially satisfactory control of Botrytis leaf rot in cyclamen.

Kessel *et al.*, 2005 developed a spatially explicit model describing saprophytic colonization of dead cyclamen leaf tissue by the plant-pathogenic fungus *Botrytis cinerea* and the saprophytic fungal antagonist *Ulocladium atrum*. Both fungi explore the leaf and utilize the resources it provides. Leaf tissue is represented by a two-dimensional grid of square grid cells. Fungal competition within grid cells is modelled using Lotka-Volterra equations. Spatial expansion into neighbouring grid cells is assumed proportional to the mycelial density gradient between donor and receptor cell. Established fungal biomass is immobile. Radial growth rates of *B. cinerea* and *U. atrum* in dead cyclamen leaf tissue were measured to determine parameters describing the spatial dynamics of the fungi. At temperatures from 5 to  $25^{\circ}$ C, B. cinerea colonies expanded twice as rapidly as *U. atrum* colonies. In practical biological control, the slower colonization of space by *U. atrum* thus needs to be compensated by a sufficiently dense and even distribution of conidia on the leaf. Simulation results confirm the importance of spatial expansion to the outcome of the competitive interaction between *B. cinerea* and *U. atrum* at leaf scale. A sensitivity analysis further emphasized the importance of a uniform high density cover of vital *U. atrum* conidia on target leaves.

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# **Chapter 7**

# Identified difficulties and conditions for field success of biocontrol. 3. Economic aspects: cost analysis

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The industrial and commercial development of biological control agents, although needed as an alternative to chemical pesticides in both organic farming and IPM systems is facing different constraints which are particularly difficult to overcome due to the size of the involved companies and the early development stage of the market. These constraints can be classified within four

- categories:size of the targeted market
  - cost of production
  - costs of registration
  - business profitability

In this paper, in order to be more specific, we shall consider the situation regarding microbial biocontrol agents (MBCAs), using the real case of a well defined product that we cannot mention here due to proprietary rights.

#### Size of the targeted markets

In most of the situations MBCAs are being developed with rather small, if not niche markets. The total value of MBCAs sold worldwide amounted in 2008 to 620 Mio Euro (122 Mio Euro in Europe) including products with insecticidal or fungicidal effects. This value can be compared with the sales of chemical insecticides and fungicides amounting to a total of 21 000 Mio Euros.

MBCAs, with the exception of Bt products which can be used in larger crops such as grapes, forestry or even cereals, are presently still used in speciality crops, greenhouses and covered crops. The size of these crops is not growing anymore or at a very reduced rate. The only optimistic perspective is the intention to develop organic faster farming (objective 20% of the production area in France in 2030) where MBCAs can find a good market.

Additionally the potential market is widely fragmented within a long list of crops such as carrots, petersillium, onions, etc, usually referred to as "Minor crops". These markets are so small that even large chemical companies refrain from the investments that would cover the needs and the manufacturers of MBCAs, due to the specificity of their products, are obliged to invest and cover costs where scale economy can never be reached.

#### **Cost of production**

Contrary to the synthesis of chemicals, producing MBCAs requires a complicated and extremely expensive process of production which can be divided into four phases: fermentation, extraction,

purification, formulation and packaging. All these phases are difficult and require relatively heavy costs.

**Fermentation.** This first step has to be undertaken either with solid or with liquid phase technology. Although the liquid phase fermentation is usually simple and cost effective, the process is more risky because the produced spores are more fragile. In the contrary using solid fermentation substrates will produce stronger, but it becomes more difficult to increase the production volume.

**Extraction**. Here again, there is a very strong difference between the MBCAs produced in liquid or in solid fermenters. In a liquid, the extraction will be rather easy by filtration, but the product will need to be dried, which is a very long, energy-demanding and expensive process. From a solid fermentation process, the extraction will be mechanical. Such a process is rather harmful for the spores: It is again energy demanding and it is extremely difficult to extract more that 60% of the spores from a substrate. In such a case the productivity becomes rather poor.

**Purification**. This step is very important to ensure the stability of the MBCAs produced. The industrially produced MBCAs always contain impurities which, although biologically inactive, may become critical over time, potentially creating risks of degradation, inactivation etc. In all situations the purification step requires a high level of sophistication and expensive processes.

**Formulation and packaging**. Formulation and packaging of MBCAs, due to their living state (and the requirement that they remain alive for satisfactory effectiveness of the product), constitute an extremely difficult step and in any case more expensive than the equivalent process for chemicals. The choice of co-formulants, adjuvants and packaging material must secure the quality of the MBCAs and its vitality. This is again a source of problems and heavy costs.

Additionally to all the above mentioned hurtles, it has to be secured that no contamination will occur, during the fermentation process naturally, but also during the extraction, the purification, the formulation and the packaging. All the safety measures are very expensive to carry out, but they are necessary in order to ensure the quality of the product brought to the market. As a consequence of all these extra expenses and technical difficulties the MBCAs used for this analysis were more than 2.5 times more expensive to produce than an equivalent chemical pesticide (Table 14).

	Typical Insecticide	MBCA	Comments
Sales value	100	100	
Type of production cost			
Raw materials	%* 8	29	40% lost material for MBCA by solid fermentation process
Packaging	1	2	
Energy and miscellaneous	1	2	
Manpower	5	9	
Consumables	2	3	
Amortisation	4	11	
TOTAL	21	56	

 Table 14:
 Compared structure of the production costs for a microbial biocontrol agent (MBCA) and a chemical insecticide (source IBMA).

\* costs are expressed as percent of the sales value of the commercial product

#### **Cost of registration**

It has been already mentioned that biological control agents suffer from a highly unfavourable situation compared to chemical pesticides. The regulations for registration have initially been set up to reduce the risks attached to molecules and the regulator is trying to extrapolate these requirements for the registration of living organisms.

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The estimated cost for registering a microbial biocontrol agent is currently lower than that for a chemical pesticide (Table 15). However, the size of this investment is still very high for a company in comparison with the market potential (Table 16). This evaluation indicates that the introduction on the market of a MBCA is about 4 times less effective than its chemical equivalent.

Table 15:	Compared potential costs of registration for a microbial biocontrol agent (MBCA) and a
	chemical pesticide (source IBMA)

Area	Study Type	Cost for Chemical (€)	Cost for MBCA (€)
	Acute studies (6 tests)	140 000	140 000
Toxicity of the	Sub-acute (rat study)	140 000	120 000
active substance	Mutagenicity	40 000	may be waived
	Toxicity on cultured cells	10 000	not required
Toxicity of the	Acute studies	140 000	140 000
formulation	Toxicity on cultured cells	10 000	not required
Environmental fate	Soil, water, air	200 000	70 000
Biology	Mode of action etc	150 000	*50 000
Ecotoxicology of	Birds, fish, bees, algae, daphnia, earthworm	60 000	40 000
active substance	Beneficials	20 000	may be waived
Ecotoxicology of	Birds, fish, bees, algae, daphnia, earthworm	60 000	40 000
formulation	Beneficials	20 000	
Residues	8 trials / crop	80 000	may be waived
Kesiuues	Development of analytical methods	100 000	**variable
Formulation	Physical properties, shelf life, etc.	200 000	220 000
Efficacy	8 field trials	40 000	40 000
TOTAL		1 410 000	860 000

\* cost of strain identification

\*\* e.g. development of strain-specific markers

Table 16:	Compared estimated market potential for a microbial biocontrol agent (MBCA) and for a
	chemical pesticide (source: IBMA)

Year	Estimated sales value ( Mio€)		
1 ear	Chemical pesticide	MBCA	
1	0.1	0.05	
2	1.2	0.15	
3	6.0	0.90	
4	15.0	1.50	
5	35.0	3.50	
Total early sales	57.3	6.10	
Plateau sales	120.0	15.00	
Registration costs	1.410	0.860	
Ratio registration/ early sales	2.4 %	14.0 %	
Ratio registration/ Plateau sales	1.1 %	5.7 %	

## **Business profitability**

Comparing estimated production and other costs, relative to the sales value at plateau level, points out large differences between chemical pesticides and microbial biocontrol agents (Table 17). The gap between the two in terms of estimated profit is nearly 10-fold in favour of the chemical industry.

%	Chemical pesticide	MBCA
Sales value at plateau level	100	100
Costs of production	21	56
Gross margin	79	44
Cost of sales	21	15
Cost of research	8	12
Cost of administration	4	3
<b>Earnings</b> before investments taxes and amortisation (EBITA)	46	14
<b>Profit</b> after taxes, provisions and amortisation	18	2

 Table 17:
 Compared margin structure estimates for the production and sales of a microbial biocontrol agent (MBCA) and a chemical pesticide (source IBMA)

\* costs and margins are expressed as percent of the sales value of the commercial product

## **Conclusion and outlook for industry**

These data show clearly that the profitability of a biocontrol business is much less attractive than that of chemical pesticides and may explain why the large chemical companies decided in the 90's to retreat from this business. Although these companies show presently some new signs of interest, they seem to remain basically reluctant to re-enter despite the new attractiveness of a fast growing biocontrol market. Contrary to European and US-based companies, several Japanese firms, such as Sumitomo chemicals or Mitsui appear to have invested for a potential long term return. Taking advantage of the divestment by the chemical majors, they have been able to acquire a good business basis at very attractive conditions. This should enable them to consider optimistically the future development of the biocontrol industry and its positive trend.

The smaller companies which have invested in this business and try to overcome their financial problems have only two alternatives:

- Either develop, often at a loss, into larger markets (grapevine, field crops etc), if they can. In order to sustain these efforts, they will need a strong support from venture capital companies;
- or enter into venture agreements with other manufacturers/suppliers, in order to build up a product portfolio which will make them successful in the future.

### **Chapter 8**

## **Identified difficulties and conditions for field success of biocontrol. 4. Socio-economic aspects: market analysis and outlook**

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With estimated sales amounting to only 200 Mio€ in Europe in 2008, the market for biological control agents appears to be extremely small compared with the 7 000 Mio€ turnover achieved with chemical pesticides. However, very important efforts have been undertaken for the development of biocontrol agents. The OECD estimated that 5 000 Mio\$ have been spent worldwide in public research for biocontrol during the last 40 years. This amounts to a yearly average of 500Mio\$, not far from the 600 Mio\$ spent yearly in research by the agrochemical industry, but with a comparatively poor result!

In the Conference on biological control organised in 2003 by IBMA in Béziers, France, the major stakeholders (farmers, retailers, distributors, regulators etc.) have provided a list of gaps considered to play a role in preventing wide adoption of biocontrol products. This list was meant to cover all potential explanations, but provided neither figures nor priority ranking, making it difficult to prioritize actions for improvement. It was however a general opinion that the complicated and costly system of registration was the major reason of the problem. As a result, important efforts have been undertaken to convince the regulators to adopt more facilitating procedures for the registration of biologicals. These efforts were not without effect and the newly adopted "Pesticides package" makes it easier, under certain conditions, to register biologicals. In the meantime, several EU member states have adopted easier registrations tracks, such as the Biopesticides Scheme in the UK, for example.

In reality, the unique assumption that the current regulations in Europe significantly hamper the development and the use of biologicals does not seem to be proven by the facts. During a very long period, the biologicals were not subject to registration and very few products were brought successfully to the market. At the same time countries such as the USA, New Zealand or Japan have adopted very liberal registration procedures, but the sales of biologicals remain marginal.

In the frame of ENDURE, it has been therefore decided to get a detailed and quantified idea on the gaps which, in Europe, restrain the adoption of biologicals, especially at the users and commercial levels. In order to achieve this objective, a Pan-Europa survey was undertaken from 2007 until 2008, with the assistance of the public opinion organisation Agridata.

#### Methodological approach: survey of European farmers

Since no validated data were available about the real market and the use of biological control agents in Europe, it has been necessary to build up a form of electronic map of the European agriculture and of the distribution of the potential users. A survey was carried out to evaluate the size of the biocontrol market in Europe and to identify key factors that could influence its future evolution. This study included four main steps:

- Localisation of the main crops and cropping systems. Using the data from EUROSTAT and national statistics a model of European agriculture was constructed.
- Randomised sampling of farmers and retailers.
   The model was used for the selection of 12 production systems (Table 18) located on 25 sites in 9 countries (Table 19) where 2000 farmers and 21 retailers were identified.
- The selected sample was contacted by phone directly and a questionnaire (Table 20) was sent to those who agreed to participate in the survey. A total of 675 full responses were obtained and analysed.
- Complementary survey. In order to validate the process, more specific data was collected i

In order to validate the process, more specific data was collected in a survey concerning the biological control of wood diseases of grapevine in France

Table 18:	Production systems selected for a survey of factors influencing biocontrol use in Europe
	(source IBMA)

Type of cropping system	Geographical sub-categories
Large arable crops	North, South
Multicropping	
<ul> <li>arable crops dominant</li> </ul>	Mountains, North, South
animal production dominant	Mountains, North South
Fruit production	
orchards	
• grapes	
Tomato production	
• protected	
• field	

 Table 19:
 Geographical distribution of sampling sites for a survey of factors influencing biocontrol use in Europe (source IBMA)

Country	number of sampling site
Austria	2
Denmark	1
Germany	4
Greece	2
France	4
Italy	4
Poland	3
Spain	3
United Kingdom	2

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Table 20:	Structure of the	questionnaire	used	in a	survey	of	European	farmers	and	retailers	of
	biological contro	l products									

Categories of questions	Nbr of Questions
Geographical identification	5
System of production concerned	12
Ownership and social related aspects	5
Crop protection issues / pest occurrence, etc	18
Economy of the farm, actual costs, revenues etc	12
Expectations for future, cropping systems, investments, etc	9
Relations with input suppliers	18
Relations with advisors	18
Relations with authorities	18
Relation with the food chain (coops, supermarkets etc.)	18
Relations with the consumers	18
Relations with the public	18
Expectations about innovations, role of science	12
Open comments	2

#### Survey Results: The estimated market of biocontrol in Europe

The questionnaire made it possible to estimate the total biological market in ha and in value (Figure 11) and its partition among different crops (Figure 12).

These data confirm that in 2008, the main use of biologicals was in protected crops, followed by grapevine and fruit production. Nearly 40% of the estimated biocontrol market consisted in sales of beneficial insects, compared to 25% for microorganisms and 21% for semiochemicals (Figure 11).



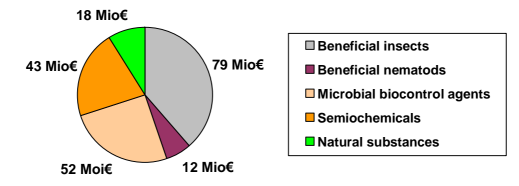
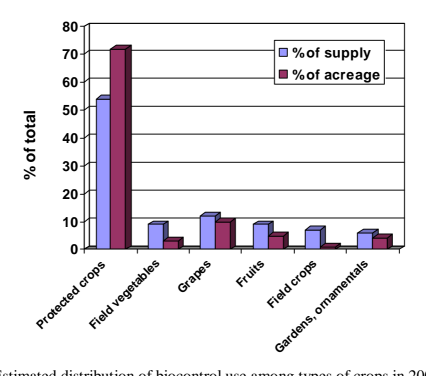
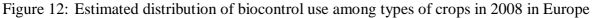


Figure 11: Estimated sales of biocontrol products in Europe in 2008 (in Million €). The estimates were obtained by extrapolating use patterns in a representative sample of EU farmers.





#### **Survey results: Factors of development of biocontrol**

The exploitation of the questionnaires was somewhat difficult due to the large variety of farmers and situations. Additionally, several open ended questions were introduced to collect opinions on possible additional gaps and opportunities which were not mentioned in the form.

**Qualitative analysis.** In a first step, the analysis of the responses led to the identification of 12 factors deemed to have a significant influence on the future development of biological control

Nine factors with a positive influence:

- o Ability of manufacturers to invest in R&D
- Financial strength of manufacturers
- Direct involvement of leading distributors
- o Pull from the fresh food wholesalers and from the food industry
- Demand from consumers and NGOs
- o Incentives given to growers
- o Education of advisors and growers
- Availability of Decision Support Systems (DSS)
- o Regulatory obstacles to chemical pesticides

Three factors with a negative influence:

- o Regulations not adapted to Biological control
- o Discovery of novel effective and safe chemicals
- Development of new resistant crops

Quantitative analysis. In a second step, a quantitative analysis was conducted to estimate the influence of the identified factors. For this, 320 contacts (50% of the sample) were requested to

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indicate which of the 12 factors they considered as important in terms of their potential impact on the evolution of future use of biological control agents. For those factors selected as important, the respondents were asked to weigh the expected impact positively or negatively on a scale from 0 to 20.

The data were used to compute for each of the 12 factors:

- a) an **Influence Index**, calculated as the percentage of respondents who selected the factor as important
- b) a **Weight Index**, calculated as the average of the weights attributed to the factor by those respondents who selected it as important
- c) a **Growth Index**, combining the two other indices according to the following formula:

GI = (Influence Index)\*(Weight Index)/10

This index represents the overall estimate of the influence of a factor on the future use of biological control agents by European farmers.

The scores computed for each of the 12 factors are presented in Table 21. Among the factors deemed to carry the most impact on future use of biological control by European farmers the action by far the most cited was the establishment of incentives for farmers (factor D).

	Factors	Influence Index (%)*	Weight Index* (scale from -20 to +20)	Growth Index*	Rank of positive influence
А	Registration for biological control products remains as present	12	- 15	- 18.0	
В	Involvement of distribution	65	8	52.0	4
С	Size / strength of the manufacturers	55	12	66.0	3
D	Incentives to growers	87	18	156.6	1
E	Education of advisors and growers	27	8	21.6	5
F	Decision Support Systems available	12	7	7.2	9
G	Pull from wholesalers and food industry	43	16	66.8	2
Н	Stringent registration of chemicals	16	14	22.4	6
Ι	New safe chemical pesticides	42	- 12	- 3.0	
J	Progress in R&D of Biocontrol	8	14	11.2	8
K	New resistant varieties	16	- 4	- 6.4	
L	Pull from Consumers	67	2	13,4	7

Table 21: Impact of twelve factors on the future use of biocontrol agents by European farmers according to a survey of 320 farmers

\* see main text above for the specific definition of the indices

The second most important factors based on the Growth Index (G, C and B in Table 21) were linked to the influence of key economic actors (the wholesalers, the food industry, the distributors and manufacturers of biocontrol products). The factors with the lowest scores were those related to scientific innovation (factors K, I, J). Interestingly, both factors linked to regulatory aspects (factors H and A) also had a relatively low Growth Index. The registration requirements are obviously more a concern for the industry than for the users of the plant protection products. Surprisingly, the efficacy and the price of the biologicals, usually considered as two critical factors, were not mentioned as real constraints. This may be due to two reasons:

(1) It is anticipated that only "effective" solutions will be registered in the EU, showing the high confidence of the farmers and the retailers in the registration systems

(2) The selling price of the new solutions (biological control products) will necessarily cope with the current price levels. Too highly priced, the new solutions will simply be ignored.

#### Conclusions

The gaps and the opportunities for the development of biological crop protection products are extremely relative to people concerned. While the industry, due to the heavy factor time/cost to the market, considers the regulation requirements as a major obstacle, the users and the retailers are much more influenced by the pull and push actions exercised at the market level. Somewhat disappointing is the relative low concern about the technical progress offered by the biological solutions.

## **Conclusions and perspectives**

# Perspectives for future research-and-development projects on biological control of plant pests and diseases

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The review of published scientific literature on the biological control of selected pests and diseases has lead to the identification of clear knowledge gaps highlighted in previous chapters. Further bottlenecks were revealed by seeking the possible reasons for the striking discrepancy between the rich inventory of potential biocontrol agents described by scientists and a very small number of commercial products on the market.

To complement these analyses, the participants of Research Activity 4.3 of the European Network ENDURE organized consultations of experts (scientists, extension specialists and representatives of the Biocontrol industry) at the occasion of scientific meetings of three Working Groups of IOBC-wprs.

- Working Group "Integrated Control of Plant Pathogens": meeting on "Molecular Tools for Understanding and Improving Biocontrol" in Interlaken (Switzerland) September 9-12, 2008. (attended by P.C. Nicot and B. Blum – discussion session about the outlook on biocontrol against plant diseases)
- Working Group "Multitrophic Interactions in Soil" meeting in Uppsala (Sweden), 10-13 June 2009. (attended by C. Alabouvette roundtable about the outlook on biocontrol of soilborne pests and diseases)
- Working Group "Insect Pathogens and Insect Parasitic Nematodes": meeting on "Future Research and Development in the Use of Microbial Agents and Nematodes for Biological Insect Control" in Pamplona (Spain), 22-25 June, 2009 (attended by C. Alabouvette his plenary presentations about the outlook on biocontrol of diseases and pests has been published<sup>\*</sup>).

These consultations were further complemented by discussions at the occasion of various meetings of participants of Research Activity 4.3 to identify the most prominent issues that could be tackled by future research and development activities. The key elements are organised below in three categories, based on their relevance to the concern of the research community, development or industry.

#### **Research issues**

Five key issues have been identified in term of research needs:

**Devise better strategies for the screening of biocontrol agents**. The demand for new biocontrol agents is already high. It is expected to increase sharply in the EU, with the ongoing

<sup>&</sup>lt;sup>\*</sup> Alabouvette, C, Cordier, C. 2009 Biological control of plant diseases: Future research goals to make it successful. IOBC/WPRS Bulletin 45:3-5.

reduction of available chemical pesticides and the need for new non-chemical plant protection tools to comply with Directive 2009/128/EC. Current methods need to be improved both in terms of logistics (high throughput to allow rapid mass screening of large numbers of candidates) and in terms of the pertinence of criteria for efficacy, production and commercialization. This topic has been tackled within Research Activity 4.3 of the European Network ENDURE for microbial biocontrol agents against diseases (Deliverable DR4.9) and the results have been published (Köhl *et al.*, 2011<sup>\*</sup>).

**Improve knowledge on efficacy-related issues.** The criteria traditionally used to asses the efficacy of biological control methods may be misleading because contrarily to conventional pesticides, biocontrol does not intend to eradicate pests and diseases but, rather, to install a biological balance which will enable the plants to grow more healthily. However the consistency of field efficacy remains one of the constraints for the large scale use of biological control of plant diseases. Despite much recent progress, research efforts are still necessary for (1) a better understanding of key parameters of field efficacy in relation to the type of biocontrol agent and their modes of action and (2) implementing the most promising methods for efficacy improvement. Promising avenues of research are to be sought both in terms of exploiting the biological properties of the biocontrol agents and enhancing their effectiveness through formulation of the products. Results obtained on these topics should provide key information both for the design of optimised production and application strategies, but also for improving the screening process of future biocontrol agents as mentioned in the point above.

**Promote multidisciplinary approaches to integrate better biocontrol with IPM and other production issues.** Based on passed published experience, it is clear that levels of protection provided by a single biocontrol agent alone will seldom be sufficient, especially when faced with field conditions unfavourable to their effectiveness or with very high inoculum pressures of a pest or plant pathogen. More emphasis will need to be placed on the compatibility of biocontrol agents with the implementation of IPM, preferably in a <u>systemic approach</u> of integrated production. Among the many possible interactions to be considered, compatibility and combined used of biocontrol and plants genetically modified for improved resistance to pest or plant diseases should not be overlooked.

**Develop adapted delivery technologies.** Much progress has been made in packaging technology and delivery for macrobial biocontrol agents (e.g. beneficial arthropods). In contrast, treatments with microbial biocontrol agents (against pests or diseases) still rely on sprayers developed for the application of pesticides. Research is needed to provide growers with low pressure spraying equipment to preserve the viability of the microbials. Technological improvements are also needed for optimal coverage of the target plant surfaces to be protected by the biocontrol agents.

**Safeguard the durability of biocontrol.** Certain pests and pathogens are known for their capacity to develop resistance to chemical pesticides or to overcome varietal resistance. The durability of biological control has often been assumed to be higher than that of chemical control, but several examples of resistance of pests have already been reported. Much less is known about plant pathogens, probably in part because biological control against diseases is still very rare. Significant research efforts are needed to anticipate the potential hurtles in this domain and integrate durability concerns both in the screening of new biocontrol agents and in the careful management of their use once they become commercially available.

<sup>&</sup>lt;sup>\*</sup>Köhl, J., Postma, J., Nicot, P., Ruocco, M., Blum, B. 2011. Stepwise screening of microorganisms for commercial use in biological control of plant pathogenic fungi and bacteria. Biological Control 57, 1-12.

#### **Issues for development**

Three key issues have been identified in terms of development. They are directly related to improving the efficacy of crop protection but also to acceptability of biocontrol by farmers.

**Training of advisers and farmers**. Compared to chemical control, the implementation of biological control presents an additional level of technical complexity when the "active substance" is a living organism or microorganism, whose liveliness and development on the target crop underpins the effectiveness of the protection. In many situations, achievement of successful biocontrol of pests has been linked to an active role of advisers in accompanying the farmers, at least during their initial phase of adoption and implementation. The success of large scale use of biological control in the future will require stepping up the technical training of farmers and of advisors. Such action will also positively influence the adoption issues mentioned below.

**Development and dissemination of Decision Support Systems (DSS).** Growers routinely make decisions that take into account multiple constraints (both technical and economic) of their activity. However, the complexity of biocontrol and its necessary integration in a systems approach of crop protection and crop production make DSS more and more indispensible, including in their function as easily consultable repositories of knowledge on available choices.

**Establishment of demonstration schemes and development of farmers' networks.** This action is needed to stimulate the dissemination of information to and among farmers, but also to facilitate exchange between the end users of biocontrol and the other actors of research, development and commercialization of the products. Breaking up regional and national barriers and including a European dimension to such networks is desirable for optimal efficacy of multisite experimental trials.

#### **Industrial issues**

**Quality control.** Ongoing efforts by the manufacturers of biological control agents to guarantee the quality of their products need to be stepped up. The definition of tests and their routine implementation is crucial to ensure reliable effectiveness and maintain confidence of farmers for biocontrol. Whenever possible, such tests should include not only an evaluation of viability of the biocontrol agent but also an evaluation of physiological parameters related to its efficacy, based on knowledge of its modes of action.

**Improve distribution systems.** Distibution systems need to be improved to safeguard the quality of the products and provide technical advice for the users. In many cases, the distribution of biocontrol products is common with that of chemical pesticides. One possible avenue of progress would be to improve awareness on the specificities of handling biocontrol products, especially those containing living organisms or micro-organisms. Another would be the development of sizeable distributions networks focused on biocontrol, which could be brought together by groups of (currently often small) producers of biocontrol products.

## Appendices

#### For Chapter 1

- Appendix 1. Inventory of biocontrol agents described in primary literature (1998-2008) for successful effect against *Botrytis* sp. in laboratory experiments and field trials with selected crops
- Appendix 2. Inventory of biocontrol agents described in primary literature (1998-2008) for successful effect against powdery mildew in laboratory experiments and field trials on selected crops.
- Appendix 3. Inventory of biocontrol agents described in primary literature (1973-2008) for successful effect against the rust pathogens in laboratory experiments and field trials on selected crops
- Appendix 4. Inventory of biocontrol agents described in primary literature (1973-2008) for successful effect against the downy mildew / late blight pathogens in laboratory experiments and field trials on selected crops
- Appendix 5. Inventory of biocontrol agents described in primary literature (1973-2008) for successful effect against *Monilinia* in laboratory experiments and field trials on selected crops
- Appendix 6. Primary literature (2007-2009) on biological control against Fusarium oxysporum

#### For Chapter 2

- Appendix 7. Number of references retrieved by using the CAB Abstracts database in order to review scientific literatures on augmentative biological control in selected crops for Chapter 2.
- Appendix 8. Collection of data on augmentative biological control of pests in grapevine. Each table refers to a group of biocontrol agents.

#### For Chapter 3

Appendix 9. References on classical biological control against insect pests (cited in Chapter 3)

#### For Chapter 4

- Appendix 10. Substances included in the "EU Pesticides Database" as of April 21 2009
- Appendix 11. Invertebrate beneficials available as biological control agents against invertebrate pests in five European countries.

#### **Tomato + Cucumber + Pepper** (target pathogen = B. cinerea) **Success in laboratory conditions** (*in vitro* and/or *in planta* in controlled conditions) Success in field trials Bacteria Bacteria Bacillus amyloliquefaciens BL3, pepper (Park et al., 1999) Bacillus antagonists (Tsomlexoglou et al., 2000) (Enva et al., 2007) (Tsomlexoglou et al., 2001) (Tsomlexoglou et Bacillus licheniformis > FG (Lee *et al.*, 2006) al., 2002) Bacillus subtilis strain QST 713 (Serenade ASO) (Ingram and Meister, 2006), Bacillus circulans (Wang et al., 2008b) Quadra 136, preventive (Utkhede and Mathur, 2006) Bacillus subtilis (Wang et al., 2008b) (Sadfi-Zouaoui et al., 2007a) (Gu et al., 2008) (Sadfi-Zouaoui et al., 2007b) Brevibacillus brevis (Seddon et al., 2000) (McHugh et al., 2002) (Schmitt et al., Bacillus licheniformis (Lee et al., 2006) (Sadfi-Zouaoui et al., 2007a) 2001) Brevibacillus brevis (White et al., 2001) (Seddon and Schmitt, 1999) (Seddon et al., 2000) (Allan et al., 2003) Brevibacillus brevis WT + Milsana / cucumber (Konstantinidou-Doltsinis et al., Cupriavidus campinensis / cuc, tom (Schoonbeek et al., 2007) Halomonas subglaciescola, Halobacillus litoralis, Marinococcus halophilus, Salinococcus roseus, Halovibrio 2002)variabilis, Halobacillus halophilus, Halobacillus trueperi (Sadfi-Zouaoui et al., 2008) Paenibacillus polymyxa BL4, pepper (Park et al., 1999) Pseudomonas putida Cha94, pepper (Park et al., 1999) Halomonas sp. K2-5 (Sadfi-Zouaoui et al., 2007b) Streptomyces (Mycostop(R), (Lahdenpera and Korteniemi, 2008) actinomyces Micromonospora coerulea (Kim et al., 1999) (Yao et al., 2007), strains III-61 and A-21 (Pan et al., 2005) Pantoea (Enya et al., 2007) Bakflor (consortium of valuable bacterial physiological groups) (Kornilov et al., Pseudomonas aeruginosa (Hernandez-Rodriguez et al., 2004) 7NSK2 (Audenaert et al., 2002) 2007)Pseudomonas fluorescens (Yildiz et al., 2007) (Hernandez-Rodriguez et al., 2004) Burkholderia cepacia (Hernandez-Rodriguez et al., 2004) Serratia plymuthica HRO-C48 (Ma et al., 2007), IC1270 (Meziane et al., 2006), IC14 / cucumber (Kamensky et Fungi + veasts: Clonostachys rosea (ADJ 710 OMRI), (Shipp et al., 2008) al., 2002, Kamensky et al., 2003) Gliocladium sp. (Georgieva, 2004) Streptomyces ahygroscopicus var. wuviensis (Sun et al., 2004) Μ Gliocladium catenulatum Prestop(R), preventive (Utkhede and Mathur, 2006) Streptomyces lydicus/ cucumber (Farrag, 2003) (Utkhede and Mathur, 2002) (Lahdenpera and Korteniemi, 2008) Fungi + veasts: Gliocladium viride (Lisboa et al., 2007) Aureobasidium pullulans (Dik et al., 1999) (Dik and Elad, 1999) Microdochium dimerum (Nicot et al., 2003) (Trottin-Caudal et al., 2001) Beauveria sp. (Diaz et al., 2007) Rhodosporidium diobovatum S33 preventive (Utkhede and Mathur, 2006) Candida guilliermondii strains 101 and US 7 (Saligkarias et al., 2002) curative (Utkhede and Mathur, 2002), /cucumber (Utkhede and Bogdanoff, Candida oleophila strain I-182 (Saligkarias et al., 2002) Candida pelliculosa (Bello et al., 2008) 2003) Clonostachys rosea (Nobre et al., 2005) (Sutton et al., 2002) (Yohalem, 2001) Trichoderma sp. (Georgieva, 2004) Trichoderma harzianum (Lisboa et al., 2007), T39 (Trichodex) tomato (Apablaza Cryptococcus laurentii (Xi and Tian, 2005) and Jalil R, 1998) (Moreno Velandia et al., 2007), tomato + cucumber Cryptococcus albidus (Dik et al., 1999) (Dik and Elad, 1999) (Elad, 2000b) (Dik and Wubben, 2001) / cucumber (Elad, 2000a), TM / Gliocladium (Hmouni et al., 2005) (Hmouni et al., 2006, Hmouni et al., 1999) pepper (Park et al., 1999), RootShield curative (Utkhede and Mathur, 2002). Gliocadium viride (Bocchese et al., 2007) (Lisboa et al., 2007) T22 PlantShield(R) curative (Utkhede and Mathur, 2006) Microdochium dimerum (Bardin et al., 2008) (Bardin et al., 2004b) (Bardin et al., 2004a) (Decognet and Nicot, 1999) (Decognet et al., 1999) (Trottin-Caudal et al., 2001) (Nicot et al., 2002) Variable little or no effect once in the field (good in lab): Pichia guilliermondii (Zhao et al., 2008) Brevibacillus brevis WT / cucumber (Konstantinidou-Doltsinis et al., 2002) Rhodosporidium diobovatum (S33), (Utkhede et al., 2001) Gliocladium catenulatum (Prestop). (Ingram and Meister, 2006) Rhodotorula glutinis Y-44 (Kalogiannis et al., 2006) Trichoderma (tomato + pepper) (Salas Brenes and Sanchez Garita, 2006) Rhodotorula rubra (Bello et al., 2008) Trichoderma harzianum T39 Trichodex with BOTMAN (Moyano et al., 2003) Trichoderma (Hmouni et al., 2005) (Hmouni et al., 1999) Trichoderma harzianum (Hmouni et al., 2006) (Fiume et al., 2008) (Barakat and Al-Masri, 2005) (Lisboa et al.,

## Appendix 1. Inventory of biocontrol agents (M: microbials; B: botanicals; O: others) described in primary literature (1998-2008) for successful effect against Botrytis sp. in laboratory experiments and field trials with selected crops

## Appendix 1

		<ul> <li>2007) T115 (Meyer et al., 2001) Trichodex T39 (Elad et al., 1998) (Yohalem et al., 1998) (Meyer et al., 1998) (Jalil R et al., 1997) (Dik et al., 1999) (Dik and Elad, 1999), RootShield (Utkhede et al., 2001), Th-B /pepper (Li et al., 2004), Rifai (Gromovikh et al., 1998)</li> <li>Trichoderma taxi ZJUF0986 (Wang et al., 2008a)</li> <li>Trichosporon pullulans (Cook, 2002)</li> <li>Ulocladium atrum (Nicot et al., 2002) (Fruit and Nicot, 1999) (Yohalem, 2001) / cucumber (Yohalem, 1997)</li> <li>Ustilago maydis (Teichmann et al., 2007)</li> <li><u>Oomvcetes</u></li> <li>Pythium oligandrum (Floch et al., 2001) (Wang et al., 2007a)</li> <li>Little or no effect once in the field (good in lab):</li> <li>Trichoderma spp. commercial preparations/ cucumber (Yohalem, 1997)</li> </ul>
в	<ul> <li>Milsana + Brevibacillus brevis WT / cucumber (Konstantinidou-Doltsinis <i>et al.</i>, 2002)</li> <li>Variable little or no effect once in the field: Reynoutria sachalinensis extract (Milsana); (Ingram and Meister, 2006)</li> </ul>	volatile substances produced by grape cv. Isabella (Vitis labrusca) (postharvest) (Kulakiotu <i>et al.</i> , 2004) (Kulakiotu and Sfakiotakis, 2003)
0	calcium foliar fertilizers (CaH2O2, CaSO4, Ca(NO3)2, CaCl2 and CaO), (Mizrakci and Yildiz, 2002)	<ul> <li>Compost water extracts prepared from animal sources (horse, sheep, and cattle) and a plant source (olive), (Hmouni <i>et al.</i>, 2006)</li> <li>Adipic acid monoethyl ester (Vicedo <i>et al.</i>, 2005)</li> <li>Calcium foliar fertilizers (CaH2O2, CaSO4, Ca(NO3)2, CaCl2 and CaO), (Mizrakci and Yildiz, 2002)</li> <li>Chitosan Elexa (Acar <i>et al.</i>, 2008)</li> <li>Benzothiadiazole (BTH) (Hernandez-Rodriguez <i>et al.</i>, 2004)</li> <li>Variable little or no effect :</li> <li>Vital pasta, Vital gel and Elot-Vis (Gielen <i>et al.</i>, 2004)</li> </ul>

Gr	<b>apes</b> (target pathogen = $B$ . <i>cinerea</i> )	
	Success in field trials	Success in laboratory conditions (in vitro and/or in planta in controlled conditions)
	BacteriaAcinetobacter Iwoffii PTA-113, (Magnin-Robert et al., 2007)Pseudomonas fluorescens PTA-CT2, (Magnin-Robert et al., 2007)Pantoea agglomerans PTA-AF1 (Magnin-Robert et al., 2007)Bacillus (isolate UYBC38) (Rabosto et al., 2006)Bacillus subtilis strain QST 713 (serenade) (Benuzzi et al., 2002)	Bacteria         Bacillus sp., (Paul et al., 1998) (Krol, 1998) (Trotel-Aziz et al., 2003), isolate UYBC38 (Rabosto et al., 2006)         Cupriavidus campinensis (Schoonbeek et al., 2007)         Pseudomonas sp. (Trotel-Aziz et al., 2003), strain PsJN (Barka et al., 2002)         Pseudomonas fluorescens (Krol, 1998)         Pantoea (Trotel-Aziz et al., 2003)         Fungi + yeasts:         Alternaria spp., (Walter et al., 2006)         Aureobasidium pullulans, L47 postharvest (Lima et al., 1997), LS-30 postharvest (Castoria et al., 2001)
М	<ul> <li>Fungi + veasts: Acremonium cephalosporium, strain B11 (Zahavi <i>et al.</i>, 2000) Candida guilliermondii, strain A42 (Zahavi <i>et al.</i>, 2000) Chaetomium cochlioides (Lennartz <i>et al.</i>, 1998) Gliocladium (Cherif and Boubaker, 1998) Gliocladium roseum (Holz and Volkmann, 2002) Hanseniaspora uvarum (isolate UYNS13) (Rabosto <i>et al.</i>, 2006) Trichoderma (Cherif and Boubaker, 1998) Trichoderma harzianum (Holz and Volkmann, 2002), Rootshield(R) (Marco and Osti, 2007) Rifai, 1295-22, (Harman <i>et al.</i>, 1996), Trichodex 25 WP (Turcanu, 1997)</li> <li>Trichoderma virens 31 (Harman <i>et al.</i>, 1996)</li> <li>Trichosporon pullulans (Holz and Volkmann, 2002)</li> <li>Ulocladium atrum, low disease pressure (Metz <i>et al.</i>, 2002) (Roudet and Dubos, 2001) (Schoene <i>et al.</i>, 1999) (Holz and Volkmann, 2002) (Lennartz <i>et al.</i>, 1998) (Schoene and Köhl, 1999), isolate 385 (Schoene <i>et al.</i>, 2000)</li> <li>Ulocladium oudemansii + 5-chlorosalicylic acid in combination (Reglinski <i>et al.</i>, 2005)</li> <li>Variable little or no effect once in the field: Trichoderma harzianum partial effect (Monchiero <i>et al.</i>, 2005)</li> <li>Ulocladium oudemansii partial effect (Monchiero <i>et al.</i>, 2005)</li> <li>Ulocladium atrum, high disease pressure (Metz <i>et al.</i>, 2002) (Roudet and Dubos, 2001)</li> </ul>	<ul> <li>Candida oleophila (Lima et al., 1997), postharvest (El-Neshawy and El-Morsy, 2003)</li> <li>Coniothyrium (Sesan et al., 2002)</li> <li>Debaryomyces hansenii (Santos et al., 2004)</li> <li>Epicoccum spp (Sesan et al., 2002) (Walter et al., 2006) (Fowler et al., 1999)</li> <li>Gliocladium, (Sesan et al., 2002)</li> <li>Hanseniaspora uvarum (isolate UYNS13) (Rabosto et al., 2006)</li> <li>Kloeckera spp. (Cirvilleri et al., 1999)</li> <li>Metschnikowia fructicola, postharvest (Karabulut et al., 2003), postharvest (Kurtzman and Droby, 2001)</li> <li>Muscodor albus, postharvest (Gabler et al., 2006)</li> <li>Pichia anomala (strain FY-102) (Masin et al., 2000) (Santos et al., 2004)</li> <li>Pichia membranaefaciens (Masih and Paul, 2002) (Masih et al., 2001) (Santos and Marquina, 2004) (Santos et al., 2004)</li> <li>Scytalidium, (Fowler et al., 1999)</li> <li>Trichoderma spp. (Walter et al., 2006) (Fowler et al., 1999)</li> <li>Trichoderma viride, (Sesan et al., 2002)</li> <li>Trichothecium, (Sesan et al., 2002)</li> <li>Trichothecium, (Sesan et al., 2002)</li> <li>Trichothecium, (Sesan et al., 2002)</li> <li>Ulodadium spp (Walter et al., 2006) (Fowler et al., 1999)</li> <li>Ulocladium atrum isolate 385 (Schoene et al., 1999)</li> <li>Ulocladium atrum isolate 385 (Schoene et al., 2000)</li> <li>Verticillium, (Sesan et al., 2002)</li> <li>Oomvectes</li> <li>Pythium paroecandrum (Abdelghani et al., 2004)</li> <li>Pythium periplocum (Paul, 1999b)</li> </ul>
в	<ul> <li>Croplife (citrus and coconut extract) + Plantfood (foliar fertilizer), moderate to good control (Schilder <i>et al.</i>, 2002)</li> <li>Milsana (giant knotweed [Fallopia sp.] extract), moderate control (Schilder <i>et al.</i>, 2002)</li> </ul>	volatile substances produced by grape cv. Isabella (Vitis labrusca) (postharvest) (Kulakiotu <i>et al.</i> , 2004) (Kulakiotu and Sfakiotakis, 2003)
0	Chitosan (Amborabe et al., 2004)	

Success in field trials		Success in laboratory conditions (in vitro and/or in planta in controlled conditions)
		Bacteria
		Bacillus sp. (isolate 17141) (Helbig et al., 1998)
		Bacillus pumilus (Essghaier et al., 2007), NCIMB 13374 (Swadling and Jeffries, 1998)
		Bacillus subtilis, (Essghaier et al., 2007) (Sardi et al., 2008) (Helbig and Bochow, 2001) (Marquenie et al., 1999)
Bacteria		(Zhao et al., 2007) (Abada et al., 2002) (Gengotti et al., 2000)
Paenibacillus polymyxa 18191	(Helbig, 2001b)	Bacillus marismortui, (Essghaier et al., 2007)
Pseudomonas fluorescens (Aba	da <i>et al.</i> , 2002)	Bacillus licheniformis, (Essghaier et al., 2007)
		Bacillus thuringiensis (Bacikol) (Kandybin, 2003)
Fungi + yeasts:		Virgibacillus marismortui, (Essghaier et al., 2007)
Aureobasidium pullulans (Stror		Enterobacteriaceae (10B1, 5B4) (Guinebretiere et al., 2000)
Candida fructus, (El-Neshawy a		Halomonas sp. (Essghaier et al., 2007)
C. glabrata, (El-Neshawy and S		Pantoea agglomerans strain EPS125, postharvest (Bonaterra et al., 2004)
C. oleophila (El-Neshawy and S		Pseudomonas fluorescens (Abada et al., 2002), NCIMB 13373 (Swadling and Jeffries, 1998)
Cryptococcus albidus (Helbig, 2		Pseudomonas cepacia (Marquenie et al., 1999)
Epicoccum nigrum, (Stromeng		Pseudomonas chlororaphis isolate I-112 (Gulati <i>et al.</i> , 1999)
Metschnikowia fructicola (=FG		Pseudomonas syringae but phytotox (Pellegrini et al., 2007)
	mycoides mixture (Guetsky et al., 2001)	<u>Fungi + yeasts</u> :
(Guetsky et al., 2002)		Aureo basidium pullulans (Adikaram et al., 2002)
Rhodotorula glutinis (Helbig, 2		Candida reukaufii, (Guinebretiere et al., 2000)
	a et al., 2002) (Antoniacci et al., 2000)	Candida pulcherrima, (Guinebretiere et al., 2000)
	, 1295-22 (Kovach et al., 2000), (atroviride) P1	Clonostachys rosea (Cota et al., 2008), IK726 (Mamarabadi et al., 2008)
	9 (Shafir et al., 2006), Trichodex (Freeman et al.,	Cryptococcus albidus (Helbig, 2002)
	02) (Freeman <i>et al.</i> , 2004)	Cryptococcus laurentii (Zheng et al., 2003)
Trichoderma products (BINAB)		Gliocladium virens (Tehrani and Alizadeh, 2000)
	(Boff <i>et al.</i> , 2002a) (Boff <i>et al.</i> , 2002b) (Köhl <i>et</i>	Metschnikowia fructicola (Shemer(R) postharvest (Ferrari <i>et al.</i> , 2007)
	04) (Köhl and Fokkema, 1998)	Pichia guilermondii + Bacillus mycoides mixture (Guetsky <i>et al.</i> , 2002b) (Guetsky <i>et al.</i> , 2001b) (Guetsky <i>et al.</i> ,
Variable little or no effect once		2001a) (Guetsky <i>et al.</i> , 2002a)
Bacillus subtilis (Gengotti <i>et al.</i>		Rhodotorula glutinis, postharvest (Zhang <i>et al.</i> , 2007a), (Helbig, 2001a)
Gliocladium roseum (Chaves an		Trichoderma sp (Santorum <i>et al.</i> , 2002)
	kkola <i>et al.</i> , 2003), but low disease incidence	Trichoderma harzianum (Abada <i>et al.</i> , 2002) (Tehrani and Alizadeh, 2000) (Sanz <i>et al.</i> , 2002), T39 (Bilu <i>et al.</i> , 2004) (Least of a 200
(Prokkola and Kivijarvi, 2		2004) (Levy <i>et al.</i> , 2004a) (Levy <i>et al.</i> , 2006) (Levy <i>et al.</i> , 2004b), atroviride P1 (Hjeljord, Stensvand <i>et al.</i> 2001)
	97), (Stensvand, 1998) (Hjeljord <i>et al.</i> , 2000) It low disease incidence (Prokkola and Kivijarvi,	$\frac{2001}{\text{Trickederma commutery (Commuter)}}$
(PTOKKOTA <i>et al.</i> , 2005), 00 2007)	it low disease incluence (Plokkola and Kivijarvi,	Trichoderma asperellum (Sanz <i>et al.</i> , 2005) (Sanz <i>et al.</i> , 2002) Trichoderma longibrachiatum (Sanz <i>et al.</i> , 2002)
	ride) (Hjeljord, 2002) (Hjeljord et al., 2001),	Trichoderma atroviride (Sanz <i>et al.</i> , 2002)
	ichodex 40 WP (Meszka and Bielenin, 2004)	Trichoderma koningii, (Tehrani and Alizadeh, 2000)
(Oengotti <i>et al.</i> , 2002), 11	100000 + 0  with $(101002 Ka allu Dicicilili, 2004)$	Trichoderma viride (Tehrani and Alizadeh, 2000)
		Ulocladium atrum (Boff, 2001) (Berto <i>et al.</i> , 2001) (Boff <i>et al.</i> , 2001)
		Verticillium lecanii (Koike <i>et al.</i> , 2004)
		Variable little or no effect once in the field:
		Pichia guilermondii (Wszelaki and Mitcham, 2003)

I	<b>Variab</b> B Biosept	nger (harpin), (Meszka and Bielenin, 2004) <b>ble little or no effect once in the field:</b> ot 33 SL (grapefruit extract) (Meszka and Bielenin, 2004) red, garlic, and compost extracts (Prokkola <i>et al.</i> , 2003), but low disease	
(	ir sodium Variabl Biochic silicon	incidence (Prokkola and Kivijarvi, 2007) n bicarbonate (Funaro, 1997) <b>ble little or no effect once in the field:</b> icol 020 PC (chitosan) (Meszka and Bielenin, 2004) n (Prokkola <i>et al.</i> , 2003), but low disease incidence (Prokkola and Kivijarvi, 2007)	Natural volatile compounds : benzaldehyde, methyl benzoate, methyl salicylate, 2-nonanone, 2-hexenal diethyl acetal, hexanol, and E-2-hexen-1-ol (Archbold <i>et al.</i> , 1997)

<b>Field vegetables (lettuce, onion, cabbage, melon)</b> (target pathogen = <i>B. cinered</i>	a)
Success in field trials	Success in laboratory conditions (in vitro and/or in planta in controlled conditions)
<ul> <li>M Microsphaeropsis ochracea / onion (Carisse <i>et al.</i>, 2006) Ulocladium atrum 385, onion (Köhl and Fokkema, 1998) (Köhl <i>et al.</i>, 1999)</li> </ul>	<ul> <li>Bacteria Bacillus subtilis / lettuce (Fiddaman et al., 2000), L-form / Chinese cabbage (Walker et al., 2002), / melon (Wang et al., 2008c)</li> <li>Brevibacillus brevis / lettuce (McHugh and Seddon, 2001)</li> <li>Bacillus amyloliquefaciens/ melon (Wang et al., 2008c)</li> <li>Pseudomonas spy. (LC8, PF13, PF14, PF15), / lettuce (Card et al., 2002)</li> <li>Pseudomonas syringae pv. phaseolicola / Chinese cabbage (Daulagala and Allan, 2003)</li> <li>Fungus + yeast:</li> <li>Clonostachys rosea / onion (Nielsen et al., 2000) (Yohalem et al., 2004)</li> <li>Coniothyrium minitans / lettuce (Fiume and Fiume, 2005)</li> <li>Epicoccum sp. (E21) / lettuce (Card et al., 2002)</li> <li>Gliocladium virens [Trichoderma virens], / lettuce (Lolas et al., 2005)</li> <li>Penicillium griseofulvum, / onion (Tylkowska and Szopinska, 1998)</li> <li>Pichia onychis /onion postharvest (German Garcia et al., 2001) (Cotes, 2001)</li> <li>Ulocladium sp. (U13), /lettuce (Card et al., 2002)</li> <li>Ulocladium atrum / onion (Köhl et al., 2003), 385 and 302 / onion (Nielsen et al., 2000) (Yohalem et al., 2004)</li> <li>Trichoderma harzianum, / onion (Tylkowska and Szopinska, 1998), T39 / lettuce (Meyer et al., 1998) (Lolas et al., 2005), Supresivit' / cress (Borregaard, 2000)</li> <li>Trichoderma koningii / onion (Tylkowska and Szopinska, 1998)</li> <li>T. viride / onion (Tylkowska and Szopinska, 1998)</li> </ul>
B	
0	

Success in field trials	Success in laboratory conditions (in vitro and/or in planta in controlled conditions)
<ul> <li>Bacteria Pantoea agglomerans (CPA-2) (Nunes <i>et al.</i>, 2002b) (Nunes <i>et al.</i>, 2001b) Pseudomonas syringae, MA-4, MB-4, MD-3b and NSA-6 (=FG) (Zhou <i>et al.</i>, 2001)</li> <li>Fungi + veasts: Aureobasidium pullulans, Rhodotorula glutinis and Bacillus subtilis in combination (=FG) (Leibinger <i>et al.</i>, 1997)</li> <li>Candida saitoana (El-Ghaouth <i>et al.</i>, 2001a), with chitosan (Bio-Coat) or lyric enzyme (Biocure) (El-Ghaouth <i>et al.</i>, 2001b)</li> <li>Candida sake strain CPA-1 combined with diphenylamine (Zanella <i>et al.</i>, 2003), CPA- 1 + ammonium molybdate /pear (Nunes <i>et al.</i>, 2002a)</li> <li>Metschnikowia pulcherrima (Migheli <i>et al.</i>, 1997)</li> <li>Pichia anomala strain K beta -1,3-glucans and calcium chloride (Jijakli <i>et al.</i>, 2002)</li> </ul>	<ul> <li>Bacteria         <ul> <li>Bacillus licheniformis (EN74-1) (Jamalizadeh et al., 2008)</li> <li>Bacillus subtilis (Ongena et al., 2005), GA1 (Toure et al., 2004), Rizo-N (El-Sheikh Aly et al., 2000)</li> <li>Bacillus amyloliquefaciens 2TOE. /pears (Mari et al., 1996)</li> <li>Bacillus subtilis (Ongena et al., 2005), GA1 (Toure et al., 2001a)</li> <li>Bacillus of (Floros et al., 1998)</li> </ul> <ul> <li>Partoca agglomerans (Sobiczewski and Bryk, 1999) (Nunes et al., 2001a)</li> <li>Pseudomonas syring Strain ESC-11 BioSave (Janisiewicz and Jeffers, 1997), / pear (Sugar and Benbow, 2002) (Benhow and Sugar, 1997), MA-4 (Zhou et al., 2002), CPA5 (Nunes et al., 2007)</li> </ul> </li> <li>Pseudomonas fluorescens (Mikani et al., 2007) (Mikani et al., 2008)</li> <li>Pseudomonas viridiflava (Bryk et al., 1999)</li> <li>Rahnella aquatilis (Calvo et al., 2007)</li> </ul> <li><b>Fungi + veasts:</b> <ul> <li>Aureobasidium pullulans (Achbani et al., 2005) (Lima et al., 2005) (Schena et al., 1999), LS-30 (Lima et al., 2005a)</li> <li>Candida butyri JCM 1501, (Wagner et al., 2006)</li> <li>Candida nelibiosica 2515 (Wagner et al., 2006)</li> <li>Candida nelibiosica 2515 (Wagner et al., 2006)</li> <li>Candida nelibiosica 2515 (Wagner et al., 2007), peach (Karabulut and Baykal, 2004) (Baji and Jijakli, 2007)</li> <li>(Jijakli et al., 2004) (Lahlai et al., 2007), peach (Karabulut and Baykal, 2004)</li> <li>Candida sationan (El-Ghaouth et al., 2001c) (El-Ghaouth et al., 2000a) (El-Ghaouth et al., 2001b)</li> <li>Candida sake (Vinas et al., 1998) (Nunes et al., 2002) (Giraud and Crouzet, 2004) (Cook, 2002b), CPA-1 + Pantoea agglomerans (Nunes et al., 2002)</li> <li>Candida pulcherrima (Cook, 2002b)</li> <li>Cryptococcus laurentii (Benhow and Sugar, 1997) (Zhang et al., 20</li></ul></li>

	Kloeckera apiculata / peach (Karabulut and Baykal, 2003) (Karabulut et al., 2005)
	Metschnikowia pulcherrima (Spadaro et al., 2002) (Piano et al., 1998) (Spadaro et al., 2004), MACH1 (Duraisamy
	<i>et al.</i> , 2008)
	Metschnikowia fructicola (Karabulut <i>et al.</i> , 2005)
	Muscodor albus (Mercier and Jimenez, 2004) (Ramin et al., 2008) (Schotsmans et al., 2008)
	Penicillium spp. (El-Sheikh Aly et al., 2000)
	Pichia stipitis CBS 5773 (Wagner <i>et al.</i> , 2006)
	Pichia anomala strain K (Grevesse <i>et al.</i> , 2003) (Jijakli, 2000) (Friel and Jijakli, 2007) (Friel <i>et al.</i> , 2007) (Jijakli
	and Lepoivre, 1998) (Lahlali et al., 2007)
	Pichia guilliermondii (29-A), (Lima <i>et al.</i> , 1999)
	Rhodotorula glutinis (Sugar and Benbow, 2002) (Benhow and Sugar, 1997) (Lima et al., 2005) (Lima et al., 1998)
	(Sansone et al., 2005), LS-11 (Lima et al., 1999) (Lima et al., 2003),
	Rhodosporidium toruloides NRRL Y1091, (Filonow et al., 1996)
	Sporobolomyces roseus FS-43-238 (Filonow et al., 1996) (Filonow, 1998)
	Saccharomyces cerevisiae, (Faten, 2005)
	Trichoderma harzianum Plant-guard (El-Sheikh Aly et al., 2000), Rifai (Batta, 2004)
	Trichoderma Viride (El-Sheikh Aly et al., 2000),
	Trichosporon sp., (Fan <i>et al.</i> , 2001b) (Tian <i>et al.</i> , 2002)
	Trichosporon pullulans (Cook, 2002b)
	R. glutinis SL 1 + C. laurentii SL 62 mixture (Calvo et al., 2003)
	Variable little or no effect :
	Candida oleophila (Aspire), (Colgan, 1997)
В	volatile substances produced by grape cv. Isabella (Vitis labrusca) (Kulakiotu and Sfakiotakis, 2003b) (Kulakiotu
D	<i>et al.</i> , 2004a)
	Chitosan, (Faten, 2005)
0	Calcium (Chardonnet et al., 2000) (Holmes et al., 1998)
J	Phosphonate (Holmes et al., 1998)
	sodium bicarbonate (Karabulut <i>et al.</i> , 2005)

Flo	Flowers (target pathogen = B. cinerea)				
	Success in field trials	Success in laboratory conditions ( <i>in vitro</i> and/or <i>in planta</i> in controlled conditions)			
М	Bacteria         Bacillus amyloliquefaciens B190 / lily (Chiou and Wu, 2003)         Bacillus cereus / lily (Liu et al., 2008)         Bacillus amyloliquefaciens / lily (Chiou and Wu, 2001)         Burkholderia gladioli, / lily (Chiou and Wu, 2001)         Pseudomonas putida / lily (Liu et al., 2008)         Fungi + veasts:         Clonostachys rosea /rose (Morandi et al., 2003)         Ulocladium atrum / cyclamen (Köhl et al., 2000) (Köhl et al., 1998)         Variable little or no effect :         Trichoderma harzianum / cyclamen (Minuto et al., 2002) (Minuto et al., 2004)	Bacteria         Bacillus amyloliquefaciens / lily (Chiou and Wu, 2001)         Bacillus subtilis / rose buds (Tatagiba et al., 1998)         Burkholderia gladioli, / lily (Chiou and Wu, 2001)         Pseudomonas sp. 677 /geraldton waxflower (Beasley et al., 2001)         Serratia marcescens strain B2 / cyclamen (Someya et al., 2001)         Serratia marcescens strain B2 / cyclamen (Someya et al., 2001)         Fungi + yeasts         Cladosporium oxysporum, / rose debris + buds (Tatagiba et al., 1998)         Cladosporium cladosporioides / rose buds (Tatagiba et al., 1998)         Cladosporium cladosporioides / rose buds (Tatagiba et al., 1998)         Clonostachys rosea /rose (Morandi et al., 1999) (Morandi et al., 2006) (Morandi et al., 2001) (Morandi et al., 2007) (Morandi et al., 2008) (Morandi et al., 2006) (Morandi et al., 2004) (Yohalem, 2000)         Epicoccum sp. / Geraldton waxflower (Beasley et al., 2001)         Fusarium sp., / Geraldton waxflower (Beasley et al., 2001)         Glicoladium roseum FR136 / rose debris (Tatagiba et al., 1998)         Rhizoctonia (BNR), / geranium (Buck and Jeffers, 2004) (Buck, 2004)         Rhodotorula glutinis PM4 / geranium (Buck, 2004)         Trichoderma harzianum (Trichodex) / Geraldton waxflower (Beasley et al., 2005)         Trichoderma harzianum (Trichodex) / Geraldton waxflower (Beasley et al., 2005)         Trichoderma harzianum (Trichodex) / Geraldton waxflower (Beasley et al., 2005)         Trichoderma inhamatum / r			
		Variable little or no effect : Trichoderma hamatum 382 in compost / begonia (Horst <i>et al.</i> , 2005) Trichoderma harzianum preparations (Yohalem, 2000) (Trichodex and Supresivit) (Yohalem, 2004)			
B		grapefruit [Citrus paradisi] extract / lily, peony and tulip (Orlikowski <i>et al.</i> , 2002), / tulips, Gerbera jamesonii and carnations (Orlikowski and Skrzyoczak, 2003), Biosept 33 SL / tulip (Orlikowski and Skrzypczak, 2001) chitosan / tulips, Gerbera jamesonii and carnations (Orlikowski and Skrzypczak, 2003)			
0					

Mis	Miscellaneous crops (target pathogen = B. cinerea)					
	Success in field trials	Success in laboratory conditions (in vitro and/or in planta in controlled conditions)				
М	Bacteria         Streptomyces griseoviridis (Mycostop) / Pinus sylvestris (Capieau et al., 2001) (Capieau et al., 2004)         Fungi + yeasts         Gliocladium sp (GlioMix) / Pinus sylvestris (Capieau et al., 2001) (Capieau et al., 2004)         Gliocladium roseum / Eucalyptus nurseries (Stowasser and Ferreira, 1997)         Trichoderma harzianum and T. polysporum (Binab TF.WP), / Pinus sylvestris (Capieau et al., 2001) (Capieau et al., 2004)         Trichoderma viride (Trichosemin 25 PTS (25% Tv), / sunflower (Eva, 2003)         Variable little or no effect :         Penicillium sp. / Eucalyptus nurseries (Stowasser and Ferreira, 1997)         Trichoderma harzianum , Trichoderma viride / Eucalyptus nurseries (Stowasser and Ferreira, 1997)	Bacteria         Bacillus spp./ Ginseng (Kim et al., 1997) (Chung et al., 1998)         Bacillus subtilis Cot1 and CL27 / Astilbe hybrida, Aster hybrida, Daphne blayana, Photinia fraseri (Li et al., 1998)         Bacillus amyloliquefaciens / oilseed rape (Danielsson et al., 2007)         Bacillus licheniformis / Perilla (Son et al., 2002)         B. megaterium / Perilla (Son et al., 2002)         Cupriavidus campinensis / Arabidopsis thaliana (Schoonbeek et al., 2007)         Erwinia / Ginseng (Kim et al., 1997)         Pseudomonas fluorescens / castor crop (Raoof et al., 2003), WCS374r / Eucalyptus (Ran et al., 2005)         Pseudomonas putida WCS358r / Eucalyptus (Ran et al., 2005)         Streptomyces griseoviridis (Mycostop) / Pinus sylvestris (Capieau et al., 2001) (Capieau et al., 2004) <b>Fungi + veasts</b> Clonostachys (A-10) / Pinus radiate, Eucalyptus globulus (Molina Mercader et al., 2006)         Cylindrocladium spp. / Eucalyptus (Fortes et al., 2007)         Gliocladium spp. / Eucalyptus (Fortes et al., 2007)         Gliocladium roseum / Picea mariana (Zhang et al., 1996)         Trichoderma harzianum / Arabidopsis thaliana (Korolev and Elad, 2004) / castor crop (Tirupathi et al., 2006)         (Raoof et al., 2003) (Bhattiprolu and Bhattiprolu, 2006), / hazelnut (Machowicz-Stefaniak et al., 2004)         Trichoderma iride / castor crop (Tirupathi et al., 2006) (Raoof et al., 2003) (Bhattiprolu, 2006), T         13-82 (Trichodermin-BL) / flax (Pristchepa et al., 2006), / ha				
В		<i>al.</i> , 2004) Mature leaf extract of Lantana camera / castor crop (Bhattiprolu and Bhattiprolu, 2006)				
Б О		Cryptogein, elicitor secreted by Phytophthora cryptogea / tobacco (Blancard <i>et al.</i> , 1998)				
U	1	Cryptogeni, enclor secrete by Englophinora cryptogea / tobacco (Biancard et al., 1998)				

**Successful inhibition** *in vitro* (target pathogen = *B. cinerea*)

	Bacteria
	Alcaligenes faecalis (Honda et al., 1999)
	Azotobacter (Khan et al., 2006)
	Bacillus sp mutant strain (Bernal et al., 2002)
	Bacillus amyloliquefaciens CCMI 1051 (Caldeira et al., 2007), BL-3 (Lee et al., 2001)
	Bacillus brevis [Brevibacillus brevis](Gu et al., 2001) (Edwards and Seddon, 2001)
	Bacillus cereus (Guven et al., 2008) (Huang and Chen, 2004)
	Bacillus circulans (Paul et al., 1997)
	Bacillus licheniformis W10 (Ji et al., 2007) (Gu et al., 2001)
	Bacillus subtilis (Gu et al., 2001) (Chen et al., 2008) (Chen et al., 2004b) (Zhao et al., 2003) (Chen et al., 2004a) (Zakharchenko et al., 2007) (Gu et al., 2004) (Novikova et al., 2003) (Hsieh et al., 2004a) (Zakharchenko et al., 2007) (Gu et al., 2004) (Novikova et al., 2008) (Hsieh et al., 2008) (Chen
	2003) (Feng et al., 2003) (Liu et al., 2007b)
	Bacillus thuringiensis CMB26 (Kim et al., 2004)
	Paenibacillus polymyxa BL-4 (Lee <i>et al.</i> , 2001)
	Photorhabdus luminescens ATCC 29999 (Hsieh et al., 2004)
	Plutella xylostella (Indiragandhi et al., 2008)
	Pseudomonas (Lian et al., 2007) (Cornea et al., 2007) (Kim et al., 2000) (Woo et al., 2002) (Bryk et al., 2004)
	Pseudomonas aeruginosa PUPa3 (Kumar et al., 2005)
	Pseudomonas antimicrobica (Walker et al., 2001)
	Pseudomonas corrugata strain P94 (Guo et al., 2007)
	Pseudomonas fluorescens (Nian et al., 2007) (Khan and Almas, 2002)
3.4	Pseudomonas putida (Cornea et al., 2007), Cha 94 (Lee et al., 2001)
Μ	Pseudomonas syringae pv. syringae strain B359 (Fogliano et al., 2002)
	Lysobacter capsici sp. Nov (Park et al., 2008)
	Serratia plymuthica C48 (Frankowski et al., 2001a) (Frankowski et al., 2001b)
	Streptomyces + actinomycetes (Tian et al., 2004b) (Nadkarni et al., 1998) (Liang et al., 2007, Yan et al., 2004) (Han et al., 2004) (Liang et al., 2007) (Long et al., 2005) (Stoppacher et al., 2007) (Kim
	<i>et al.</i> , 2007b)
	Streptomyces ahygroscopicus (Sun et al., 2003) (Yang et al., 2007) (Zhao et al., 1998)
	Streptomyces luteogriseus ECO 00001 (Li et al., 2008)
	Streptomyces rimosus subsp. daheishanensis strain MY02 (Liu et al., 2004)
	Streptomyces roseoflavus strain LS-A24 (Park et al., 2006)
	Tripterygiun wilfordii (Shentu et al., 2006)
	Xenorhabdus sp. strain CB43 (Xiao et al., 2005)
	Xenorhabdus nematophilus YL001 (Liu et al., 2006)
	marine bacteria (Nie et al., 2007)
	Fungi + yeasts
	Acremonium strictum (Kim et al., 2002)
	Aspergillus fumigatus and A. terreus (El-Zayat, 2008)
	Aspergillus clavatonanicus (Zhang et al., 2008)
	Cryptococcus laurentii (isolate LS-28) (Castoria et al., 1997)
	Fusarium lateritium extracts (Anitha, 2006)
	Fusarium semitectum (Altomare et al., 2000)
<u>L</u>	

Lecanicillium muscarium (Fenice and Gooday, 2006)			
	Muscodor albus (Mercier and Jimenez, 2007)		
	Rhodotorula (Calvente et al., 2001)		
	Rhodotorula glutinis (Castoria et al., 1997)		
	Trichoderma (Pezet et al., 1999) (Chen et al., 2005) (Liu et al., 2007a)		
	Trichoderma viride (Machowicz-Stefaniak, 1998) T15 and T17 (Silva-Ribeiro et al., 2001)		
	Trichoderma atroviride (Navazio et al., 2007) (Klemsdal et al., 2006) GMO (Brunner et al., 2005)		
	Trichoderma harzianum (Dana et al., 2001) (Ding et al., 2002) (Limon et al., 2004) (Mach et al., 1999) T5A, T1 and T1A (Silva-Ribeiro et al., 2001) (Lee et al., 2001), T-33 (Witkowska and Maj, 2002)		
	Trichoderma hamatum C-1 (Witkowska and Maj, 2002)		
	Trichoderma reesei [T. longibractiatum] M7-1 (Witkowska and Maj, 2002)		
	Oomycetes		
	Pythium bifurcatum (Paul, 2003)		
	Pythium citrinum (Paul, 2004)		
	Pythium contiguanum (Paul, 2000)		
	Pythium radiosum (Paul, 1999a)		
	Antifungal metabolites of endophytic fungus, A10 (Qian et al., 2006)		
	antimicrobial peptide Ar-AMP from Amaranthus retroflexus L. (Lipkin, Anisimova et al. 2005)		
	basic haem-peroxidase (WP1) from wheat (Triticum aestivum) kernels (Caruso, Chilosi et al. 2001)		
в	Extracts from Bazzania trilobata, Diplophyllum albicans, Sphagnum quinquefarium, Dicranodontium denudatum and Hylocomium splendens (Tadesse, Steiner et al. 2003)		
D	Extracts of Sophora flavescens (Zheng et al., 2000) (Zheng et al., 1999)		
	Irpex lacteus (Fr.) Fr., Trametes versicolor (L.:Fr.) Pilat, and Chondrostereum purpureum (Pers.:Fr.) Pouzar (White and Traquair, 2006)		
	Pyrrolnitrin, produced by several bacteria (Okada et al., 2005)		
	Ten sesquiterpenes and six diterpenes from Pilgerodendron uviferum wood and bark (Solis <i>et al.</i> , 2004)		
	chlorine dioxide (Zoffoli et al., 2005)		
0	earthworm (Eisenia fetida) polysaccharides (Wang et al., 2007b)		
	chitosan derivatives (Rabea et al., 2003)		

#### References on biocontrol against Botrytis

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#### Appendix 1

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#### Appendix 1

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### Appendix 1

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# Appendix 2. Inventory of biocontrol agents (M: microbials; B: botanicals; O: others) described in primary literature (1998-2008) for successful effect against powdery mildew in laboratory experiments and field trials on selected crops.

Powdery	vdery mildew on cereals		
	Success in field trials	Success in laboratory conditions (in vitro and/or in planta in controlled conditions)	
	Bacteria Pseudomonas aureofaciens ; Bacillus subtilis ; P. fluorescens (Sanin et al., 2008)	General paper: Crop protection: management strategies (Prasad, 2005)	
М	<u>Fungi + yeasts:</u>	Bacteria         Rhizobacteria (Yigit, 2004)         Bacteria, (Azarang, 2004)         Fungi + yeasts:         Acremonium alternatum (Kasselaki, 2006a, b)         Alternaria alternata, Aspergillus niger, Bipolaris spicifera, Cladosporium cladosporioides, Curvularia lunata,         Fusarium acuminatum F. semitectum, Penicillium rubrum, (Simian, 2008)         BCAs mix (David, 2007)         Fungi (Azarang, 2004)         Fusarium oxysporum f. sp. radicis-lycopersici (Nelson, 2005)         Paecilomyces farinosus (Szentivanyi, 2006)         Verticillium lecanii (Koike, 2004)	
В		Bryophyte extracts (Tadesse, 2003)	
0		Aromatic substances (Koitabashi, 2002) Mycelial extracts (Haugaard, 2002) PAF from Penicillium chrysogenum (Barna, 2008) Secondary metabolic products of strain A19 of actinomycetes (Shen <i>et al.</i> , 2008) Verlamelin (Kim, 2002)	

Powdery	ry mildew on pome/stone fruits		
	Success in field trials	Success in laboratory conditions (in vitro and/or in planta in controlled conditions)	
	Bacteria	General paper:	
	Fungi + yeasts:	Bacteria	
	yeast (Y16) (Alaphilippe, 2007)		
М		Fungi + yeasts:	
		Ampelomyces quisqualis (Harvey, 2006)	
		Ampelomyces quisqualis (Sonali, 2005)	
		yeast (Y16) (Alaphilippe, 2008)	
В			
0			

Powdery	Powdery mildew on grapes		
	Success in field trials	Success in laboratory conditions (in vitro and/or in planta in controlled conditions)	
М	Bacteria Bacillus subtilis (Crisp, 2006)Photosynthetic bacteria (Robotic, 2002)Fungi + veasts: Ampelomyces hyperparasites (Fuzi, 2003) Ampelomyces quisqualis (Angeli, 2006a, b, c, 2007a, b)Ampelomyces quisqualis (Hoffmann, 2007) Ampelomyces quisqualis 94013 (Lee, 2004)BCAs (Amaro, 2003)BCAs (Ari, 2004)BCAs (Kaine, 2003)BCAs (Linder et al., 2006)BCAs (Zulini, 2004)Pseudozyma flocculosa (Schmitt, 2001)Yeast (Robotic, 2002)	General paper:         Bacteria         Brevibacillus brevis (Schmitt, 2001, 2002)         PGPR (Konstantinidou-Doltsinis, 2007)         Pseudomonas syringes pv. Syringe (Kassemeyer, 1998)         Serenade (Bacillus subtilis)(Schilder, 2002)         Fungi + yeasts:         Ampelomyces quisqualis (Angeli, 2006a, b, c, 2007a, b)         Ampelomyces quisqualis 94013 (Lee, 2004)         Ampelomyces quisqualis AQ10, (Schweigkofler, 2006)         BCA mix (David, 2007)         BCAs (Kaine, 2003)         BCAs {Amaro, 2003 #177         Pseudozyma flocculosa (Schmitt, 2001)         Pseudozyma flocculosa (SporodexReg. L) (Konstantinidou-Doltsinis, 2007)         Tilletiopsis spp (Haggag, 2007)	
В	Milsana (VP99) (Schmitt, 2001, 2002)	Milsana (VP99) (Konstantinidou-Doltsinis, 2001)	
0	fresh or dried milk (10%),pinolene 1%, calcium chloride (2%), tripotassium phosphate (1%) and a mixture of mineral oil (1%),sodium bicarbonate/sodium silicate (0.5%) (Casulli, 2002) mycophagous mite (Melidossian, 2005)	Milk, whey, whey protein, <i>Bacillus subtilis</i> , yeast extract medium (Crisp, 2006) Mycophagous mite (Melidossian, 2005) <i>Orthotydeus lambi</i> mites (English-Loeb, 1999, 2006, 2007)	

Powdery	mildew on strawberry pathogen: Podosphaera aphanis f.sp.	fragariae; Sphaerotheca macularis f.sp. fragariae
	Success in field trials	Success in laboratory conditions (in vitro and/or in planta in controlled conditions)
	Bacteria	Bacteria
		B. subtilis QST (Fiamingo, 2007a)
		Bacillus subtilis (Amsalem, 2004)
	Fungi + yeasts:	Bacillus subtilis (Pertot, 2004) (Pertot, 2008)
		Pseudomonas reactans (Fiamingo, 2007b)
М		<u>Fungi + yeasts:</u>
		Ampelomyces quisqualis, Trichoderma harzianum T39, Bacillus sp. F77, Cladosporium tenuissimum (Amsalem,
		2004)
		BCAs mix (David, 2007)
		T. harzianum T39 (Fiamingo, 2007a)
		Trichoderma harzianum Rifai strain T-22 (Picton, 2003)
		Trichoderma harzianum T39 (Pertot, 2004) (Pertot, 2008)

B	
0	

Powdery	Powdery mildew on tomato, pathogen: Leveillula taurica, Oidium neolycopersici, Oidium lycopersicum, Oidium spp.	
	Success in field trials	Success in laboratory conditions (in vitro and/or in planta in controlled conditions)
М	Bacteria Pseudomonas fluorescens (Shashi, 2007) Fungi + veasts: Trichoderma harzianum (Shashi, 2007)	General paper:         Bacteria         Bacillus brevis (Seddon, 1999)         Bcillus subtilis (Jacob, 2007)         Rhizobacteria B101R, B212R, and A068R, (Silva, 2004)         Serenade ; Pseudomonas strains (Laethauwer, 2006)         Fungi + veasts:         Acremonium alternatum (Kasselaki, 2006a, b)         Lecanicillium lecanii (Mycotal) (Bardin, 2004)         Lecanicillium muscarium (Bardin, 2008)         Sporothrix flocculosa (Jarvis, 2007)         Trichoderma spp. (Moreno-Velandia, 2007) (Velandia, 2007)
В		Milsana (Seddon, 1999) MilsanaReg. (VP 1999)(Malathrakis, 2002) Milsana (Trottin-Caudal, 2003) Malsana (Bardin, 2004) (Bardin, 2008) Milsana ; (Laethauwer, 2006)
0		

Powdery	dery mildew on pepper, pathogen: Podosphaera leucotricha		
	Success in field trials	ccess in field trials Success in laboratory conditions ( <i>in vitro</i> and/or <i>in planta</i> in controlled conditions)	
	Bacteria	General paper:	
	<u>Fungi + yeasts:</u>	<u>Bacteria</u>	
М		$\frac{\text{Fungi} + \text{yeasts:}}{1 + 200}$	
		AQ10 (Ampelomyces quisqualis) (Tsror, 2004)	
		Trichoderma harzianum (Gupta, 2005)	
		Trichoderma harzianum T39; Ampelomyces quisqualis (Brand, 2002)	
		Verticillium lecanii, Tilletiopsis minor (Haggag, 2008)	
В		Milsana (Haggag, 2008)	
0		Water extract of cattle manure compost, grape marc compost, , Kaligrin and Rifol (Tsror, 2004)	

	Success in field trials	Success in laboratory conditions (in vitro and/or in planta in controlled conditions)
М	Bacteria         Bacillus brevis (Schmitt, 1999)         Bacillus isolates (Koumaki, 2001)         Brevibacillus brevis (Abd-El-Moneim, 2004)         Fungi + veasts:         Acremonium alternatum (Kasselaki, 2006a)         Ampelomyces quisqualis (Kristkova, 2003)         Ampelomyces quisqualis isolate M-10 (Benuzzi, 2006)         Ampelomyces quisqualis, Verticillium lecanii, Sporothrix         flocculosa (Dik, 1998)         Cryptococcus laurentii and Aureobasidium pullulans (Lima, 2002)         PlantShield Trichoderma harzianum (Utkhede, 2006)         Rhodotorula glutinis (Lima, 2002)         T. harzianum T39 (Levy, 2004)         Tilletiopsis washingtonensis (yeast) (El-Hafiz-Mohamed, 1999)         Verticillium lecanii; (Verhaar, 1999)	Bacceria         Bacceria         Bacillus spp (Romero, 2004a)         Bacillus subilis (Abd-El-Moneim, 2004) (Gilardi, 2008) (Keinath, 2004) (Romero, 2007b) (Romero, 2007d)         BCAs mix (David, 2007)         Brevibacillus brevis (Allan, 2007) (Konstantinidou-Doltsinis, 2002) (Schmitt, 2001) (White, 2001)         Enterobacter cloacae (Georgieva, 2003)         Xenorhabdus nematophilus (Shi, 2004)         Fungi + veasts:         Accremonium alternatum, Ampelomyces quisqualis , Lecanicillium lecanii (Romero, 2003)         Acremonium alternatum, Verticillium lecanii (Romero, 2007b)         Ampelomyces quisqualis (Gilardi, 2008) (Rankovic, 1998)         AQ10Reg. (Ampelomyces quisqualis) and MycotalReg. (Lecanicillium lecanii) (Romero, 2007b)         BCAs mix (David, 2007)         Acremonium alternatum and Verticillium lecanii, (Romero, 2001)         Lecanicillium longisporum (Kim, 2008)         Lecanicillium longisporum (Kim, 2008)         Meira geulakonigii (Steipherg, 2004)         Paecilomyces fumosoroseus (Kavkova, 2005)         Paacillousyma flocculosa, Ampelomyces quisqualis, Verticillium lecanii, Trichoderma harzianum (Dik, 2002)         Saccharomyces cerevisiae (El-Gamal, 2003)         Trichoderma harzianum (Abd-El-Moneim, 2004) (Elad, 2000)         Trichoderma harzianum (Abd-El-Moneim, 2004) (Elad, 2000)         Trichoderma harzianum T39; Ampelomyces quisqualis AQ110 (Elad, 1998)
В		Milsana (VP99) (Dik, 2002) (Schmitt, 2001) (White, 2001) Milsana (VP99) from Fallopia sachalinensis (Konstantinidou-Doltsinis, 2001)
0	fresh or dried milk (10%), pinolene 1%, calcium chloride (2%), tripotassium phosphate (1%) and a mixture of mineral oil (1%),sodium bicarbonate/sodium silicate (0.5%) (Casulli, 2002)	Fresh or dried milk (Casulli, 2002) gramicidin S; (Schmitt, 1999) lactoperoxidase system (Ravensberg, 2007) lipopeptide antibiotic neopeptins from Streptomyces sp. (Kim, 2007) lipopeptides (iturin and fengycin families of Bacillus subtilis) (Romero, 2007c) Lipopeptides of antagonistic strains of Bacillus subtilis (Romero, 2007a) oil formulations (Verhaar, 1999) Psyllobora bisoctonotata (Soylu, 2002) undiluted homogenised milk (Utkhede, 2006)

Powdery	wdery mildew on various crops, pathogen: Oidium spp. Sphaerotheca spp., Erysiphe spp		
	Success in field trials	Success in laboratory conditions (in vitro and/or in planta in controlled conditions)	
М	Bacteria         Bacillus subtilis (Nofal, 2006)         Fungi + veasts:         Verticillium lecanIi, Tilletiopsis minor (Nofal, 2006)	Bacteria         Pseudomonas fluorescens (Vimala, 2006)         P. fluorescens (Hooda, 2006)         Fungi + veasts:         Acremonium spp., Ampelomyces spp., Penicillium spp., Cladosporium spp., Trichoderma spp., Bacillus spp., Pseudomonas spp., Bradyrhizobium spp., Brachybacterium spp., Curtobacterium spp., Cryptocoocus spp., Rhodosporidium spp (Mmbaga, 2008)         Ampelomyces mycoparasites (Kiss, 2004)         BCAs (Dhananjoy, 2008)         BCAs (Eken, 2005)         BCAs(Casey, 2007)         Cladosporium cladosporioides, Cladosporium oxysporum, Drechslera hawaiensis,T richoderma viride (Sankar, 2007b)         Cladosporium oxysporum (Sankar, 2007a)         Gliocladium roseum (Lahoz, 2004)         Kyu-W63 (Koitabashi, 2005)         Trichoderma viride, T. harzianum, Pseudomonas fluorescens, mixture of T. harzianum P. fluorescens (Hooda, 2006)	
В			
0		Exudates from sclerotia of two <i>Sclerotium rolfsii</i> isolates (Pandey, 2007) <i>Mycophagous Ladybird</i> (Sutherland, 2005) <i>Phyllactinia corylea</i> (Krishnakumar, 2004) <i>Psyllobora bisoctonotata</i> (Muls.) (Soylu, 2002) <i>Psyllobora vigintimaculata</i> , (Sutherland, 2008; Sutherland, 2006)	

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Success in field trials		Success in laboratory conditions (in vitro and/or in planta in controlled
		conditions)
		<b>Bean</b> – target pathogen = <i>Uromyces appendiculatus</i>
		Pantoea agglomerans B1 (Yuen et al., 2001)
		Stenotrophomonas maltophilia C3 (Yuen et al., 2001)
		Cladosporium tenuissimum (Assante et al., 2004)
		<b>Groundnut</b> – target pathogen = <i>Puccinia arachidis</i>
		Bacillus subtilis AF 1 (Manjula et al., 2004)
		Pseudomonas fluorescens strain Pf1 (Meena et al., 2000) (Meena et al., 2002)
<b>Bean</b> – target pathogen = $U_{i}$	romvces appendiculatus	Acremonium obclavatum (Gowdu and Balasubramanian, 1993)
Bacillus subtilis (Baker et al., 1		Fusarium chlamydosporum (Mathivanan and Murugesan, 2000) (Mathivanan e al., 1998)
<b>Groundnut</b> – target pathog		<b>Soybean</b> – target pathogen = <i>Phakopsora pachyrhizi</i>
Pseudomonas fluorescens strain	n Pf1 (Meena et al., 2002)	Verticillium psalliotae, Verticillium lecanii (Saksirirat and Hoppe, 1990)
		(Saksirirat and Hoppe, 1991)
		Wheat, Oat $-$ target pathogens $=$ <i>Puccinia recondite</i> , <i>P. coronata</i>
		Pseudomonas putida strain BK8661 (Flaishman et al., 1996)
		Chaetomium globosum strain F0142 (Park et al., 2005b)
		Verticillium chlamydosporium (Leinhos and Buchenauer, 1992)
		endophytic fungi (Dingle and McGee, 2003)
		Fusaric acid from Fusarium oxysporum EF119 (Son et al., 2008)
В		
		<b>Bean</b> – target pathogen = <i>Uromyces appendiculatus</i>
·		2,6-dichloro-isonicotinic acid (CGA 41396) (Dann and Deverall, 1995)

# Appendix 3. Inventory of biocontrol agents (M: microbials; B: botanicals; O: others) described in primary literature (1973-2008) for successful effect against the rust pathogens in laboratory experiments and field trials on selected crops

	Success in field trials	Success in laboratory conditions (in vitro and/or in planta in controlled
		conditions)
		Chrysanthemum
		Verticillium lecanii (Whipps, 1993)
	<b>Coffee</b> – target pathogens = <i>Hemileia vastatrix</i>	Coffee – target pathogen = Hemileia vastatrix Bacillus lentimorbus (Shiomi et al., 2006) Bacillus cereus (Shiomi et al., 2006) Bacillus (Haddad et al., 2004) Cedecea davisae (Silva et al., 2008) Pseudomonas (Haddad et al., 2004) Acremonium (Haddad et al., 2004) Aspergillus (Haddad et al., 2004) Cladosporium (Haddad et al., 2004) Fusarium (Haddad et al., 2004) Penicillium (Haddad et al., 2004)
Μ	Bacillus sp. (Haddad et al., 2006) Pseudomonas sp. (Maffia et al., 2005), variable effect (Haddad et al., 2006)	<ul> <li>Geranium – target pathogen = Puccinia pelargonii-zonalis</li> <li>Bacillus subtilis (Rytter et al., 1989)</li> <li>Safflower – target pathogen = Puccinia carthami</li> <li>Trichoderma viride and T. harzianum, Bacillus subtilis, B. cereus, B. thuringiensis</li> <li>Pseudomonas fluorescens added alone and in combination (Tosi and Zazzerini, 1994)</li> </ul>
		<b>Poplar</b> – target pathogen = <i>Melampsora ciliata</i> Alternaria alternata and Cladosporium oxysporum (Sharma <i>et al.</i> , 2002)
		<b>Pine</b> – target pathogens = <i>Cronartium</i> and <i>Peridermium</i> Cladosporium tenuissimum (Moricca <i>et al.</i> , 2001) Scytalidium uredinicola (Moltzan <i>et al.</i> , 2001) Plant-growth-promoting rhizobacteria (Enebak and Carey, 2004)
В		
0	<b>Coffee</b> – target pathogens = <i>Hemileia vastatrix</i> acibenzolar-S-methyl (ASM) (Patricio <i>et al.</i> , 2008)	

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# Appendix 4. Inventory of biocontrol agents (M: microbials; B: botanicals; O: others) described in primary literature (1973-2008) for successful effect against the downy mildew / late blight pathogens in laboratory experiments and field trials on selected crops

	Success in field trials	<b>Success in laboratory conditions</b> ( <i>in vitro</i> and/or <i>in planta</i> in controlled conditions)
М	Bacillus subtilis (Basu et al., 2001)         Bacillus sp. isolate PB2 (Atia, 2005) effect < fungicides	<ul> <li>Serenade (<i>Bacillus subtilis</i> strain QST 713) (Stephan <i>et al.</i>, 2005) (Olanya and Larkin, 2006)</li> <li><i>Bacillus subtilis</i> B5 (Ajay and Sunaina, 2005)</li> <li><i>Bacillus, Pseudomonas, Rahnella</i>, and <i>Serratia</i> (Daayf <i>et al.</i>, 2003)</li> <li><i>Enterobacter cloacae</i> (Slininger <i>et al.</i>, 2007)</li> <li><i>Pseudomonas fluorescens</i> (Slininger <i>et al.</i>, 2007)</li> <li><i>Xenorhabdus bovienii</i> (Eibel <i>et al.</i>, 2004)</li> <li><i>Penicillium aurantiogriseum</i> (Jindal <i>et al.</i>, 2004)</li> <li><i>Trichoderma viride</i> (Hemant <i>et al.</i>, 2004)</li> <li><i>Trichoderma viride</i> (Hemant <i>et al.</i>, 2004)</li> <li><i>Penicillium, Rhizoctonia</i> and <i>Trichoderma</i> spp (Phukan and Baruah, 1991)</li> <li>various microorganisms (Stephan and Koch, 2002)</li> </ul>
В	carvone (Quintanilla, 2002)	carvone, thymol, pinochamphone, plumbagin (Quintanilla, 2002) extracts of <i>Rheum rhabarbarum</i> and <i>Solidago canadensis</i> (Stephan <i>et al.</i> , 2005) oregano extract (Olanya and Larkin, 2006) Elot-Vis (Stephan <i>et al.</i> , 2005) patatin J from potato tuber (Sharma <i>et al.</i> , 2004)
0	culture filtrates from Streptomyces padanus (Huang <i>et al.</i> , 2007) <b>negative effect:</b> salicylic acid (Quintanilla, 2002)	chitosan ElexaTM (Acar <i>et al.</i> , 2008) cyclic lipopeptides from Pseudomonas: massetolide A (Tran Thi Thu, 2007) extracts from <i>Pseudomonas fluorescens</i> (Martinez and Osorio, 2007)

	Success in field trials	Success in laboratory conditions (in vitro and/or in planta in controlled
		conditions)
М	Bacillus cereus (Silva et al., 2004) Burkholderia (Lozoya-Saldana et al., 2006), Pseudomonas (Lozoya-Saldana et al., 2006), Streptomyces (Lozoya-Saldana et al., 2006)	<ul> <li>Bacillus pumilus (Yan et al., 2002)</li> <li>Cellulomonas flavigena (Lourenco Junior et al., 2006)</li> <li>Pseudomonas fluorescens (Yan et al., 2002) (Ha et al., 2007) (Tran Thi Thu, 2007)</li> <li>Streptomyces sp. AMG-P1 (Lee et al., 2005)</li> <li>Aspergillus sp., (Lourenco Junior et al., 2006)</li> <li>Candida sp. (Lourenco Junior et al., 2006)</li> <li>Cryptococcus sp. (Lourenco Junior et al., 2006)</li> <li>Fusarium oxysporum (Kim et al., 2007a)</li> <li>Penicillium sp. (Perez Mancia and Sanchez Garita, 2000)</li> <li>Trichoderma harzianum T39 (Ferrari et al., 2007)</li> </ul>
В	Nochi leaf extract (Vanitha and Ramachandram, 1999)	capsidiol (El-Wazeri and El-Sayed, 1977) Elot-vis (Ferrari <i>et al.</i> , 2007)
0	compost extracts (Zaller, 2006)	acibenzolar-S-methyl (Becktell <i>et al.</i> , 2005) beta -amino butyric acid (Yan <i>et al.</i> , 2002) Bion (benzothiadiazole) (Surviliene <i>et al.</i> , 2003) bikaverin and fusaric acid (Son <i>et al.</i> , 2008) cellulose (Perez Mancia and Sanchez Garita, 2000) chaetoviridin A (Park <i>et al.</i> , 2005a) chitosan ElexaTM (Acar <i>et al.</i> , 2008) Chitoplant (Ferrari <i>et al.</i> , 2007) extracts from actinomycete isolates (Mutitu <i>et al.</i> , 2008) extracts from <i>Bazzania trilobata</i> and <i>Diplophyllum albicans</i> (Tadesse <i>et al.</i> , 2003) extract from <i>Gibberella zeae</i> (Kim <i>et al.</i> , 1995) phosphate (Becktell <i>et al.</i> , 2005)

	Success in field trials	<b>Success in laboratory conditions</b> ( <i>in vitro</i> and/or <i>in planta</i> in controlled conditions)
М	Bacillus brevis (Schmitt <i>et al.</i> , 2002) Bacillus subtilis (Serenade) (Schilder <i>et al.</i> , 2002) Pseudomonas fluorescens (Rizoplan) (Kilimnik and Samoilov, 2000) (Rajeswari <i>et al.</i> , 2008) Fusarium proliferatum (Falk <i>et al.</i> , 1996) Trichoderma harzianum T39 (Vecchione <i>et al.</i> , 2007) <b>little or no effect once in the field:</b> Bacillus licheniformis (Cravero <i>et al.</i> , 2000) Biorange (Bacillus subtilis, Candida oleophila, Pseudomonas spp. and Streptomyces spp.) (Spera <i>et al.</i> , 2003)	Alternaria alternata (Musetti <i>et al.</i> , 2004) Fusarium proliferatum (Bakshi <i>et al.</i> , 2001)
В	Croplife (citrus and coconut extract) (Schilder <i>et al.</i> , 2002) Plantfood (foliar fertilizer) (Schilder <i>et al.</i> , 2002) Milsana (giant knotweed extract) (Schilder <i>et al.</i> , 2002) (Schmitt <i>et al.</i> , 2002) neem (Rajeswari <i>et al.</i> , 2008)	neem (Achimu and Schlosser, 1992) extract of giant knotweed (Schmitt, 1996)
0	acylbenzolar-s methyl (Dagostin <i>et al.</i> , 2006) chitosan (Elexa) (Schilder <i>et al.</i> , 2002) Mycosin (Angeli <i>et al.</i> , 2006)	Alternaria alternata extracts (Musetti <i>et al.</i> , 2006) EXP1, copper gluconate, salt of fatty acid, plant based alcohol extract (Dagostin <i>al.</i> , 2006)

**Pearl millet** *Pennisetum glaucum* (target pathogen = *Sclerospora graminicola*)

	Success in field trials	<b>Success in laboratory conditions</b> ( <i>in vitro</i> and/or <i>in planta</i> in controlled conditions)
М	<ul> <li>Bacillus pumilus strain INR7, strain SE34 (Raj et al., 2003)</li> <li>Bacillus subtilis (Raj et al., 2003) (Raj et al., 2005)</li> <li>Pseudomonas fluorescens (Umesha et al., 1998) (Latake and Kolase, 2007)</li> <li>Gliocladium virens (Arun et al., 2004) (Raj et al., 2005)</li> <li>Trichoderma harzianum (Raj et al., 2005) (Latake and Kolase, 2007)</li> <li>Trichoderma lignorum (Raj et al., 2005)</li> </ul>	Pseudomonas fluorescens (Raj <i>et al.</i> , 2004) Aspergillus flavus, Trichoderma harzianum and T. viride (Surender <i>et al.</i> , 2005)
В		
0	milk (cow) (Arun et al., 2004)	

# Other Vegetables and fruits

	Success in field trials	<b>Success in laboratory conditions</b> ( <i>in vitro</i> and/or <i>in planta</i> in controlled conditions)
Cauliflo	wer and other crucifers (target pathogen = Peronospora parasitica)	
М		Pseudomonas sp. XBC-PS (Li <i>et al.</i> , 2007) Trichoderma harzianum (Pratibha <i>et al.</i> , 2004)
В		
0	Bion (Pratibha <i>et al.</i> , 2004) phosphonate (Kofoet and Fischer, 2007)	Bion (Gawande and Sharma, 2003)
Lettuce	(Bremia lactucae)	
Μ		
В		
0	phosphonate (Kofoet and Fischer, 2007) Trichodermin (Borovko, 2005) Pimonex, Timorex and also Alkalin potassium+silicon (Robak and Ostrowska, 2006)	
Melon /	cucumber (target pathogen = Pseudoperonospora cubensis)	
М		actinomycete (Shu and An, 2004) Bacillus strains, Z-X-3 and Z-X-10 (Li <i>et al.</i> , 2003)
В		
0	phosphonate (Kofoet and Fischer, 2007)	attenuated cucumber mosaic cucumovirus (Qin <i>et al.</i> , 1992) chitosan ElexaTM (Acar <i>et al.</i> , 2008) compost extracts (Winterscheidt <i>et al.</i> , 1990)

#### Miscellaneous

М	Azotobacter slight effect against Peronospora arborescens on opium poppy (Chakrabarti and Yadav, 1991)	Cladosporium chlorocephalum against Peronospora arborescens (Chaurasia and Dayal, 1985) (Nalini and Rai, 1988)
В		
0	phosphonate against Peronospora destructor on Allium (Kofoet and Fischer, 2007)	DL- beta -amino-n-butyric acid (BABA) against Plasmopara helianthi (Tosi and Zazzerini, 2000)

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# Appendix 5. Inventory of biocontrol agents (M: microbials; B: botanicals; O: others) described in primary literature (1973-2008) for successful effect against *Monilinia* in laboratory experiments and field trials on selected crops

Apple (tar	Apple (target pathogens = Monilinia fructigena; M. laxa)	
	Success in field trials	Success in laboratory conditions (in vitro and/or in planta in controlled
		conditions)
		Aureobasidium pullulans, Epicoccum purpurascens, Sordaria fimicola and
		Trichoderma polysporum (Falconi and Mendgen, 1994)
М		Metschnikowia pulcherrima and (Spadaro et al., 2002), (Migheli et al., 1997)
IVI		Pseudomonas syringae (Migheli et al., 1997)
		(M laxa)
		Pantoea agglomerans strain EPS125 (Bonaterra et al., 2004)

Apricot (	Apricot (target pathogen = Monilinia laxa)	
	Success in field trials	<b>Success in laboratory conditions</b> ( <i>in vitro</i> and/or <i>in planta</i> in controlled conditions)
м	<b>bacteria</b> Burkholderia gladii OSU 7 (Altindag <i>et al.</i> , 2006) (Esitken <i>et al.</i> , 2005) Bacillus OSU-142 and Pseudomonas BA-8 (Esitken <i>et al.</i> , 2005)	<b>bacteria</b> Pantoea agglomerans strain EPS125 (Bonaterra <i>et al.</i> , 2003) ( <i>M fructicola</i> ) Bacillus subtillis strain B3 (Pusey and Wilson, 1984)
		<b>fungi, yeasts</b> Metschnikowia pulcherrima (Grebenisan <i>et al.</i> , 2006) (Grebenisan <i>et al.</i> , 2008)

Plum (targ	Plum (target pathogen = Monilinia laxa)	
	Success in field trials	Success in laboratory conditions (in vitro and/or in planta in controlled
		conditions)
		bacteria
		Pantoea agglomerans strain EPS125 (Bonaterra et al., 2004)
М		Epicoccum nigrum (Larena et al., 2001)
141		Penicillium frequentans (Cal et al., 2002)
		(M fructicola)
		Bacillus subtillis strain B3 (Pusey and Wilson, 1984)

remark: no B or O for any of the crops

	Success in field trials	Success in laboratory conditions (in vitro and/or in planta in controlled
		conditions)
М	<ul> <li>bacteria (<i>M laxa</i>) Serenade (Bacillus subtilis QRD137) (Haseli and Weibel, 2002),</li> <li>fungi, yeasts (<i>M fructicola</i>) Cryptococcus laurentii (Tian <i>et al.</i>, 2004a) Epicoccum purpurascens (E. nigrum) and Gliocladium roseum (Wittig <i>et al.</i>, 1997) (<i>M laxa</i>) Aureobasidium pullulans isolates 533 and 547 (Schena <i>et al.</i>, 2003)</li> </ul>	<ul> <li>bacteria <ul> <li>(M fructicola)</li> <li>Bacillus subtilis (15 isolates) (Utkhede and Sholberg, 1986)</li> <li>Burkholderia cepacia, Bacillus subtilis (Fan et al., 2001)</li> <li>(M laxa)</li> <li>Risoplan (Pseudomonas fluorescens), Gaupsin (Pseudomonas aureofaciens = P. chlororaphis) (Shevchuk, 2006)</li> <li>Pantoea agglomerans strain EPS125 (Bonaterra et al., 2004)</li> </ul> </li> <li>fungi, yeasts <ul> <li>(M fructicola)</li> <li>Candida guilliermondii, Kloeckera apiculata, Debaryomyces hansenii (Fan et al., 2001)</li> <li>Cryptococcus infirmo-miniatus (Spotts et al., 2002)</li> <li>Cryptococcus laurentii (Wang and Tian, 2007) (Qin and Tian, 2005) (Qin et al., 2006)</li> <li>(M laxa + M fructigena)</li> <li>Trichodex (Trichoderma harzianum) (Cardei, 2001)</li> </ul> </li> </ul>
В	( <i>M laxa</i> ) Trilogy (azadirachtin-free Neemoil) (Haseli and Weibel, 2002)	
0	( <i>M laxa</i> ) lime sulphur (calcium polysulfide) (Haseli and Weibel, 2002)	

Blueberry	<b>Blueberry</b> (target pathogen = <i>Monilinia vaccinii-corymbos</i> )		
	Success in field trials	Success in laboratory conditions (in vitro and/or in planta in controlled	
		conditions)	
М	<b>bacteria</b> Serenade (Bacillus subtilis QRD137) (Ngugi <i>et al.</i> , 2005) (Dedej <i>et al.</i> , 2004) (Scherm and Stanaland, 2001) (Schilder <i>et al.</i> , 2006)	<ul> <li>bacteria</li> <li>BlightBan (Pseudomonas fluorescens A506) (Scherm <i>et al.</i>, 2004)</li> <li>Serenade (Bacillus subtilis QRD137) (Scherm <i>et al.</i>, 2004) (Thornton <i>et al.</i>, 2008)</li> <li>Pantoea agglomerans C9-1Sv (Thornton <i>et al.</i>, 2008)</li> <li>fungi, yeasts</li> <li>Gliocladium roseum H47 (Thornton <i>et al.</i>, 2008)</li> </ul>	
В			
0			

	Success in field trials	Success in laboratory conditions ( <i>in vitro</i> and/or <i>in planta</i> in controlled
		conditions)
		bacteria
		(M fructicola)
		Rizo-N (Bacillus subtilis) (El-Sheikh Aly et al., 2000)
		Bacillus amyloliquefaciens C06 (Zhou et al., 2008)
		Bacillus subtillis (Gueldner <i>et al.</i> , 1988)
		Bacillus subtillis strain B3 (Pusey <i>et al.</i> , 1986) (Pusey <i>et al.</i> , 1988) (Pusey, 1989) (Pusey and Wilson, 1984)
		Pantoea agglomerans strain IC1270 (Ritte et al., 2002)
	bacteria	Pseudomonas syringae NSA-6 (Zhou et al., 1999)
	(M fructicola)	(M laxa)
	Pseudomonas corrugata and P. cepacia; Bacillus subtilis strain B3 (Smilanick <i>et al.</i> , 1993)	Pantoea agglomerans strain EPS125 (Bonaterra <i>et al.</i> , 2003) (Bonaterra <i>et al.</i> , 2004)
	fungi, yeasts	
Μ	Epicoccum nigrum (Mari et al., 2007)	fungi, yeasts
	(M laxa)	(M fructicola)
	Epicoccum nigrum (Cal et al., 2004) (Foschi et al., 1995) (Larena et al., 2005) (Madrigal et al.,	Candida sp(Karabulut <i>et al.</i> , 2002)
	1994) (Melgarejo <i>et al.</i> , 1986)	Cryptococcus laurentii (Yao and Tian, 2005)
	Penicillium frequentans (Cal <i>et al.</i> , 1990) (Melgarejo <i>et al.</i> , 1986) (Pascual <i>et al.</i> , 2000)	Debaryomyces hansenii (Stevens <i>et al.</i> , 1997) (Stevens <i>et al.</i> , 1998)
	Penicillium purpurogenum (Melgarejo et al., 1986)	Kloeckera apiculata yeast (Karabulut and Baykal, 2003) (McLaughlin <i>et al.</i> , 1992
		Muscodor albus (Mercier and Jimenez, 2004) (Schnabel and Mercier, 2006) Pichia membranaefaciens (Xu <i>et al.</i> , 2008)
		Trichoderma atroviride (2 isolates), T viride & Rhodotorula sp (Hong <i>et al.</i> , 1998
		Plant-guard (T. harzianum) (El-Sheikh Aly <i>et al.</i> , 2000)
		( <i>M</i> laxa)
		Penicillium purpurogenum (Foschi <i>et al.</i> , 1995) (Larena and Melgarejo, 1996)
		Penicillium frequentans (Foschi et al., 1995)
		Trichoderma koningii (Foschi et al., 1995)
B	(	
		Extract from Bacillus subtillis (McKeen <i>et al.</i> , 1986)
0	Sodium bicarbonate enhances effect of Aspire (Candida oleophila) (Droby et al., 2003)	Iturin peptides from Bacillus subtillis (Gueldner <i>et al.</i> , 1988)
2		Sodium bicarbonate (Wisniewski <i>et al.</i> , 2001); enhances effect of Aspire (Candid oleophila) (Droby <i>et al.</i> , 2003)

Suc	Successful inhibition <i>in vitro</i> (target pathogen = B. <i>cinerea</i> )		
	Bacteria		
	Pseudomonas syringae pv. morsprunorum BA35, Erwinia herbicola C9- (Voland et al., 1999)		
м	Serratia plymuthica, isolate EF-5 (Frommel et al., 1991)		
IVI	<u>Fungi + yeasts</u>		
	Penicillium frequentans (Cal and Melgarejo, 1994) (Melgarejo et al., 1985)		
	Aspergillus flavus, Epicoccum nigrum, Penicillium chrysogenum and P. purpurogenum (Melgarejo et al., 1985)		
В			
0	Thiolutin from Streptomyces luteosporeus (Deb and Dutta, 1984)		
	Deferences on biogentual against Manilia		

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# Appendix 6. Primary literature (2007-2009) on biological control against Fusarium oxysporum

Abo-Elyousr, K. A. M. and H. M. Mohamed (2009). "Biological Control of Fusarium Wilt in Tomato by Plant Growth-Promoting Yeasts and Rhizobacteria." Plant Pathology Journal 25(2): 199-204. Three plant growth-promoting yeasts and two rhizobacteria were tested for controlling tomato wilt caused by Fusarium oxysporum L sp. lycopersici under greenhouse and field conditions. Under greenhouse and field conditions, all treatments were significantly reduced disease severity of tomato wilt relative to the infected control. The highest disease reductions in pots (75.0, 67.4%) and field (52.5, 42.4%) were achieved by Azospirillum brasilense and Bacillus subtilis compared to infected control. Under field condition all treatments produced the highest tomato yield compared to the control plants inoculated with the pathogen

#### Al-Jedabi, A. A. (2009). "Biological control of Fusarium root-rot of sorghum." Research Journal of Agriculture and Biological Sciences 5(4): 465-473.

several crops including sorghum that result in low grain yield. All antagonists showed inhibition of mycelial growth of F. oxysporum and the maximum inhibition was recorded when Bacillus subtilis as biocontrol agent (67.7%). The in vitro root colonization study demonstrated that after four days of germination, the cell counts obtained from the roots have increased and the maximum count is achieved by B. subtilis (16.9\*105 cfu/cm root). The greenhouse pot experiment demonstrated that T. viride and B. subtilis resulted in more than 80% suppression of root rot. The reduction in fresh weight of roots amounted to 93.6% in the control treatment inoculated with F. oxysporum alone, whereas 71.1% reduction in fresh root weight was recorded for the treatments inoculated with only F. oxysporum decreased by 94.5% in relation to the non-inoculated control. Among the potential biological control agents in this study, B. cereus resulted in 42.3 reduction in root dry weight compared to an incidence ranging from 20 to 55% for plants treated with B. subtilis, B. lecheniformis, B. cereus, T. harzianum and T. viride. Both chlorophyll fractions increased when treated with antagonist and the maximum enhancement was recorded when Bacillus subtilis used as antagonist compared with that of control. The maximum values of the carbohydrate components were recorded when Bacillus subtilis used as antagonist relative to those of control.

Amini, J. (2009). "Induced Resistance in Tomato Plants Against Fusarium Wilt Invoked by Nonpathogenic Fusarium, Chitosan and Bion." Plant Pathology Journal 25(3): 256-262.

The potential of nonpathogenic Fusarium oxysporum strain Avr5, either alone or in combination with chitosan and Bion, for inducing defense reaction in tomato plants inoculated with E oxyysporum f. sp lycopersici, was studied in vitro and glasshouse conditions. Application Bion at concentration of 5, 50, 100 and 500 mu g/ml, and the highest concentration of chitosan reduced in vitro growth of the pathogen. Nonpathogenic F oxysporum Avr5 reduced the disease severity of Fusarium wilt of tomato in split plants, significantly. Bion and chitosan applied on tomato seedlings at concentration 100 mu g a.i./plant; 15, 10 and 5 days before inoculation of pathogen. All treatments significantly reduced disease severity of Fusarium wilt of tomato relative to the infected control. The biggest disease reduction and increasing tomato growth belong to combination of nonpathogenic Fusarium and Bion. Growth rate of shoot and root markedly inhibited in tomato plants in response to tomato Fusarium wilt as compared with healthy control. These results suggest that reduction in disease incidence and promotion in growth parameters in tomato plants inoculated with nonpathogenic Fusarium and sprayed with elicitors could be related to the synergistic and cooperative effect between them, which lead to the induction and regulation of disease resistance. Combination of elicitors and nonpathogenic Fusarium synergistically inhibit the growth of pathogen and provide the first experimental support to the hypothesis that such synergy can contribute to enhanced fungal resistance in tomato. This chemical could provide a new approach for suppression of tomato Fusarium wilt, but its practical use needs further investigation.

Anand, R., S. Kulothungan, *et al.* (2009). "Assay of chitinase and beta-1,3 glucanase in Gossypium hirsutum seedlings by Trichoderma spp. against Fusarium oxysporum." International Journal of Plant Sciences (Muzaffarnagar) 4(1): 255-258.

wilt in cotton. In this regard, the six species of Trichoderma, namely T. viride, T. virens [Gliocladium virens], T. hamatum, T. harzianum, T. koningii and T. reesi, were evaluated for its biocontrol properties and induction of defence-related enzymes, namely chitinase and beta1-3-glucanase in 30 days old cotton (G. hirsutum) seedlings. Trichoderma spp. could efficiently control the growth rate of F. oxysporum. In vitro assay of chitinase and beta-1,3-glucanase revealed the maximum production by T. harzianum (56 U/ml) and T. hamatum (80 U/ml), respectively. It also produced appreciable quantities of defence enzymes. The maximum induction of chitinase and beta1-3-glucanase in plants was found to be 80 U/ml when challenged with T. harzianum, in addition to the enhancement of defence mechanism in plants. Trichoderma spp. improved the germination rate of seedlings.

Anitha, A. and M. Rebeeth (2009). "Self-fusion of Streptomyces griseus enhances chitinase production and biocontrol activity against Fusarium oxysporum f. sp. lycopersici." Biosciences, Biotechnology Research Asia 6(1): 175-180.

Protoplasts were isolated from Streptomyces griseus (MTCC - \*4734) strain using lysing enzymes and self-fusion of Streptomyces griseus protoplasts was carried out using 50% polyethylene glycol (MW 1000, Sigma Chemicals Co., USA) in protoplast buffer. The regenerated 8 self fused Streptomyces griseus were studied detailed for chitinase production and biocontrol activity. Parent strain (PSg)

# Appendix 6

showed protein content of 2.7 mg/ml with chitinase activity of 120 IU/ml. High chitinase activity was measured in the culture filtrates of most of the self-fusants (87%) than the parent. Among the fusants, the strain SFSg 5 produced protein content of 7.8 mg/ml, maximum chitinase activity of 283.3 IU/ml with a two-fold increase as compared to the parent strain. All the self-fusants exhibited increased antagonistic activity against F. oxysporum f. sp. lycopersici than the parent. Maximum inhibition (82%, 80%) of mycelial growth of F. oxysporum was recorded with fusant of SFSg 5, SFSg 1 as against 61.1% with PSg. The result implies that, the self-fused Streptomyces griseus resulted in appreciable increase of chitinase production and biocontrol activity also the significance of the protoplast fusion technique, which could successfully be used to develop hybrid strains also for commercial formulation.

Baysal, O., M. Calskan, *et al.* (2008). "An inhibitory effect of a new Bacillus subtilis strain (EU07) against Fusarium oxysporum f. sp. Radicis-lycopersici." PMPP Physiological and Molecular Plant Pathology 73(1/3): 25-32.

destructive disease on tomato (Lycopersicon esculentum Mill.) transplant seedlings and the causal organism of crown and root rot of tomato plants growing in southern coast greenhouses of Turkey. An isolate of Bacillus subtilis (EU07) identified by the 16s RNA region code gene was selected as the best antagonist and evaluated against FORL in vitro studies. Strain EU07 at 106 CFU ml-1 was able to reduce disease incidence by 75%, when applied as an inoculant. It efficiently inhibited FORL compared to the control and QST 713 (AgraQuest, Davis, CA) whose inhibition ratio was only 52% in vivo. Random amplified polymorphic DNA analyses showed banding (genetic) differences between EU07 and QST 713 whereas there were no differences between DNAs of strains that have high homology to genes involved in the synthesis of antibiotics fengycin, bacillomycin and iturin when screened by oligonucleotide primers designed based on sequence information obtained from the NCBI database. Furthermore, one specific fragment in the EU07 genome showed the highest similarity to YrvN protein by 99% and AAA ATPase domain protein (72.2%) after amplifying oligonucleotide primers that are specific to the N-acyl-homoserine lactonase (HLS) gene as a biocontrol activity marker. These results suggested an effect of EU07 on control FORL by YrvN protein as subunit of protease enzyme. Furthermore, this fragment associated with HLS gene may be a potential molecular marker for selecting effective biological control agent belonging to Bacillus in order to control soilborne pathogens such as Fusarium, suggesting impairment in FORL invasion by signaling in the plant rhizosphere.

Bernal-Vicente, A., M. Ros, *et al.* (2009). "Increased effectiveness of the Trichoderma harzianum isolate T-78 against Fusarium wilt on melon plants under nursery conditions." Journal of the Science of Food and Agriculture 89(5): 827-833.

BACKGROUND: The use of isolates of the genus Trichoderma to control Fusarium wilt in melon plants is one of the most recent and effective alternatives to chemical treatments. In this work we have studied the immobilization of the isolate Trichoderma harzianum T-78 on different carriers as an efficient method to control vascular Fusarium wilt of melon in nurseries. Different formulations were developed: liquids (spore suspension, guar gum and carboxymethylcellulose) and solids (bentonite, vermiculite and wheat bran). RESULTS: The introduction of F. oxysporum resulted in a significant decrease in seedling fresh weight. The treatments which gave a lesser reduction in weight and showing a greater biocontrol effect were the liquid conidial suspension and the solid treatments with bentonite and superficial vermiculite. Microbiological analyses revealed that the conidial suspension and all the solid treatments, except wheat bran, significantly decreased F. oxysporum populations. Of all the treatments assayed, bentonite produced the greatest decline in the F. oxysporum population. CONCLUSIONS: The most effective treatments against Fusarium wilt on melon plants were the solid treatments bentonite and superficial vermiculite. These two treatments gave the greatest plant weight, the lowest percentage of infected plants and the greatest T. harzianum population throughout the assay. (C) 2009 Society of Chemical Industry

Boureghda, H. and Z. Bouznad (2009). "Biological control of Fusarium wilt of chickpea using isolates of Trichoderma atroviride, T. harzianum and T. longibrachiatum." Acta Phytopathologica et Entomologica Hungarica 44(1): 25-38.

The efficiency of the antagonist species Trichoderma atroviride (strains Ta.3, Ta.7 and Ta.13), T. harzianum (Th.6, Th.12, Th.15, Th.16 and Th.18) and T. longibrachiatum (TL.1, TL.2, TL.4, TL.5, TL.8, TL.9, TL.10, TL.11, TL.14 and TL17) against Fusarium wilt (caused by Fusarium oxysporum f.sp. ciceris) was compared using in vitro- and in vivo-based bioassay. A significant decrease of both growth and conidia production of the pathogen was obtained compared to the control. The highest percentages of diameter colony reduction and conidial production were obtained with Ta.13, causing 65.64% reduction in colony diameter (direct confrontation), 48.71% reduction in colony diameter (indirect confrontation), and a complete inhibition of conidial production. Once more in direct confrontation, T. atroviride overgrowth the pathogen colony and sporulate above. The seed treatment by Trichoderma spp. isolates before sowing in a soil already infested by the pathogen led to a significant decrease of disease severity compared to the untreated control. The weakest index of disease severity was obtained with Ta.13, which caused 83.92% reduction compared to the control. The most effective isolates in protecting chickpea seedlings against the disease were Ta.3, Ta.7 and Ta.13 as well as Th.16. The reduction of disease severity was associated with an increase of the vegetal growth including the stem height as well as the plant fresh and dry weights.

Casimiro Michel-Aceves, A., M. Antonio Otero-Sanchez, *et al.* (2009). "In vitro biocontrol of Fusarium subglutinans (Wollenweb. and Reinking) Nelson, Toussoun and Marasas and F. oxysporum Schlecht., causal agents of "Witches' broom" of mango (Mangifera indica L.) by Trichoderma spp." Revista Mexicana de Fitopatologia 27(1): 18-26.

The antagonistic effect of native strains of Trichoderma spp. was evaluated in vitro against Fusarium oxysporum (Fo) and Fusarium subglutinans (Fs), causal agents of mango "witches' broom". Ten strains of the antagonistic fungus were isolated, one of which was selected and identified to the species level (T. harzianum); this species showed the highest percentage of antagonism inhibiting mycelial growth of Fo by 62.9% and 42.0% of Fs. In dual Cultures between Fo and/or Fs with the selected strains of Trichoderma, the time for the first contact for Fo was between 3 and 4 days, and between 2 and 3 for Fs. The greatest intersection area (0.87 cm) was observed in T. lignorum against Fo, while the intersection area in Fs with the native strain Thzn-2 was 0.85 cm. Native strains Thzn-2 and Thzcf-12, and the commercial one showed antagonism class 2, being able to stop growth of both plant pathogens. Strain Thzn-2 is promising as an alternative for biocontrol of Fo and Fs; however, it is necessary to evaluate it under field conditions.

Chebotar, V. K., N. M. Makarova, et al. (2009). "Antifungal and phytostimulating characteristics of Bacillus subtilis Ch-13 rhizospheric strain, producer of bioprepations." Applied Biochemistry and Microbiology 45(4): 419-423.

Bacillus subtilis Ch-13 industrial strain was shown to have a wide spectrum of antagonistic activities against different species of phytopathogenic fungi and bacteria. The B. subtilis Ch-13 strain produces lytic enzymes; cyanide and other antifungal metabolites; stimulates plant growth, producing phytohormones-auxin derivatives. This strain by 2.5 times reduced the quantity of tomato plants infected with phytopathogenic fungus Fusarium oxysporum during inoculation. Fungi abundance on roots with bacterial inoculation was 6.9 times less than in the absence of inoculation. The application of detected antifungal metabolites as biochemical markers for the strain enables to control the stability of physiologic and biochemical characteristics of the producer, and ensures a rapid quality assay of biopreparations with high performance liquid chromatography (HPLC).

Chen, L. and W. Chen (2009). "Genome shuffling enhanced antagonistic activity against Fusarium oxysporum f. sp. melonis and tolerance to chemical fungicides in Bacillus subtilis BS14." Journal of Food, Agriculture & Environment 7(2): 856-860.

enhance antagonistic activity against Fusarium oxysporum f. sp. melonis (FOM) and tolerance to two chemical fungicides. Strain BS14 was identified as a strain of Bacillus subtilis by the analysis of 16S rDNA sequences. A stable recombinant F35 was obtained after three rounds of shuffling. Antagonistic activity of recombinant F35 against FOM was increased by 34.52% and 65.48% compared to that of the parent strain HN8-7 with highest activity and another parent strain utilized, BS14. The tolerance to chemical fungicides was also significantly improved (p0.05) compared to that of strain BS14. Reduction of FOM of 94% was observed by using recombinant F35, which was increased by 45% compared to that of strain BS14 (p0.05) and no significant differences (p>0.05) compared to that of thiophanate methyl (MRL). Reduction of FOM of 100% was dramatically observed by using an integrated treatment combining MRL (50% of usual dosage) with recombinant F35. Strain F35 with these improved traits would be a promising biocontrol agent in the control of FOM. Here genome shuffling was proved to be a practical methodology for strain improvement of antagonistic microorganism Bacillus subtilis BS14 for enhancing antagonistic activity against FOM and tolerance to chemical fungicides.

- Clematis, F., M. L. Gullino, *et al.* (2009). "Antagonistic activity of microorganisms isolated from recycled soilless substrates against Fusarium crow rot." Protezione delle Colture(3): 29-33. We report the results obtained in biological control trials against crown and root rot of tomato incited by Fusarium oxysporum f. sp. radicis lycopersici by using microorganisms isolated from soilless cultivation systems that showed suppressiveness against this disease. Among the tested microorganisms belonging to fluorescent bacteria (32 isolates) and to fungi belonging to Trichoderma (39 isolates) and Fusarium (38 isolated), 5 bacteria and 6 fungi showed a good activity against the pathogen. Such strains will be used in greenhouse trials, under situations closer to the field, in order to evaluate their potential to be adopted under practical conditions.
- Eden Paredes-Escalante, J., J. Armando Carrillo-Fasio, *et al.* (2009). "Antagonistic microorganismos for control of the fungal complex that cause wilt in chickpea (Cicer arietinum L.) in the state of Sinaloa, Mexico." Revista Mexicana de Fitopatologia 27(1): 27-35.

The antagonistic activity in vitro of microorganisms isolated from chickpea rhizosphere, was evaluated against Fusarium oxysporum, Sclerotium rolfsii, and Rhizoctonia solani, causal agents of chickpea wilt. The native strains with the higher percentage of pathogen mycelial growth inhibition were selected and identified as Trichoderma lignorum (CIAD 06-540903), T. harzianum (CIAD 05-550903), Bacillus subtilis (CIAD-940111), and Pseudomonas fluorescens (CIAD-990111). These strains and a commercial strain of T. harzianum (T-22) were mixed with Glomus intraradices and their effectiveness to reduce chickpea wilt was compared against a chemical treatment (PCNB) and all absolute control in the field. The seed was treated with the microorganisms before sowing and evaluations of disease severity were conducted each 15 days, while root colonization by the antagonistic microorganisms was assessed 45 days after sowing. Colonization of T, harzianum CIAD 05-550903 + G. infraradices was 33 x 10(3) ufc/g fresh root-75% and B. subtilis + G. intraradices was 1.3 x 10(8) Ufc/g fresh root-75%; while the combination P.fluorescens + G. intraradices was 1.4 x 10(7) Ufc/g fresh root-88%. These treatments also showed a reduction of disease severity in 64, 57, and 51%, respectively in comparison with the control.

- El-Khallal, S. M. (2007). "Induction and modulation of resistance in tomato plants against Fusarium wilt disease by bioagent fungi (arbuscular mycorrhiza) and/or hormonal elicitors (jasmonic acid & salicylic acid): 2 changes in the antioxidant enzymes, phenolic compounds and pathogen related-proteins." Australian Journal of Basic and Applied Sciences 1(4): 717-732.
  - Induction of plant defense against pathogen attack is regulated by a complex network of different signals. In the present study interaction between hormonal signals [jasmonic acid (JA) or salicylic acid (SA)] and bioagent [arbuscular mychorrhiza (AM) fungi] was used as new strategy to enhance tomato defense responses against wilt disease caused by Fusarium oxysporum (Fo). Thus changes in various physiological defenses including antioxidant enzymes, phenolic compounds and pathogenesis related (PR) proteins were investigated in leaves of tomato plants. Results appeared that production of reactive oxygen species (ROS), mainly H2O2 and O2 increasing the time of infection. Application with bioagent AM fungi and/or hormonal elicitors (JA & SA) markedly decreased these levels, while LOX activity greatly increased as compared with infected control. SA - treated plants had the highest MDA level but JA+AM fungi treated plants recorded the highest LOX activity. Infection by Fusarium oxysporm significantly increased activity of antioxidant enzymes (SOD, APX and CAT) in tomato leaves at different stages of growth. The highest activity was recorded in leaves of AM fungi+JA-treated plants, while treatments with SA especially when applied alone markedly decreased H2O2 scavenging enzymes (APX and CAT) and greatly increased SOD activity. Thus, imbalance between H2O2 - generation and scavenging enzymes in leaves may reflect a defense mechanism in tomato or a pathogenicity strategy of the fungus. Levels of certain phenolic acids greatly changed in tomato leaves in response to Fusarium oxysporum. AM fungi and hormonal elicitors. Benzoic and Galleic acids contents markedly decreased, however, contents of coumaric, cinnamic, chlorogenic and ferulic acids increased in leaves of all treatments. Also, activity of lignification enzymes POX, PPX and PAL significantly increased in leaves of infected tomato plants. JA-treated plants caused the highest POX and PPX activities, while SA-treated plants having the highest PAL activities. High accumulation of phenolic compounds and activity POX, PPX and PAL in these plants may reflect a component of many defense signals activated by bioagent and hormonal inducers which leading to the activation of power defense system in tomato against attack. Analysis of protein electrophoresis revealed that interaction between hormone signal (JA & SA) and bioagent AM fungi mediating the expression of the majority of different PR-proteins leading to increasing defense mechanism against Fusarium oxysporum infection. Thus, induction of protein bands of molecular weights 35, 33, 32, 31 (PR-2, beta-1, 3 glucanase), 30.5 and 27 (PR-3,-4, chitinase) in infected leaves indicated the important role which played in disease resistance. Finally, the new mechanism of the combination strategy between bioagent and hormonal signals (either synergistically) played important roles for increasing various defense systems and altering expression of defense genes which leading to different PR-proteins working together to increased resistance in tomato plants against wilt disease caused by Fusarium oxysporum. In addition, results revealed that defense mechanism in plants treated with AM fungi and JA are more effective than AM fungi plus SA-treated plants.
- Floch, G. l., J. Vallance, *et al.* (2009). "Combining the oomycete Pythium oligandrum with two other antagonistic fungi: root relationships and tomato grey mold biocontrol." Biological Control 50(3): 288-298.

To reduce Pythium oligandrum biocontrol variability and improve its efficacy, experiments were performed by combining the oomycete with two other antagonistic fungi, Fusarium dishes, Fo47 or T. harzianum hyphae destroyed P. oligandrum cells by antibiosis and mycoparasitism processes; in the rhizosphere of tomato plants (Lycopersicon esculentum), the same antagonistic features were observed. However, in the rhizosphere, hyphae are frequently separated by a certain distance; this allows the coexistence and the persistence of the three microorganisms on the root systems. When introduced in the rhizosphere, Fo47 and P. oligandrum were able to penetrate the root tissues with Fo47 limited to the epidermal and upper layers of cortical cells while P. oligandrum colonized deeper tissue at a faster rate. The two antagonists were killed in few days within roots following elicited plant-defense reactions. T. harzianum was not able to penetrate root tissues. Root colonization with either P.oligandrum alone or in combination with Fo47 and/or T. harzianum resulted in systemic plant resistance which provided plant protection against Botrytis cinerea infection of leaves. The level of control and the expression of pathogenesis-related proteins (PR-proteins) in leaves were similar whatever the antagonistic microbial treatment applied to roots.

Gay, M. I. T., Anonymous, et al. (2009). Substrates containing a Trichoderma asperellum strain for biological control of Fusarium and Rhizoctonia, Universidad de Barcelona.

The strain of Trichoderma asperellum T34(2) CECT No. 20417 is useful for preparing substrates for biological control of vascular fusariose and death of plants caused by Rhizoctonia solani. The substrates can be peats, composts (hardwood compost, pine bark compost, cork compost, sludge compost from sewage treatment plants, garden residues, etc.) or formulations based on CPV-type compost (compost+peat+vermiculite). The fact that the substrates suppress both Fusarium oxysporum f. sp. lycopersici and Rhizoctonia solani provides an advantage in comparison with other substrates known in prior art. Another advantage is that the use of methyl bromide, a highly harmful product for the environment, in the control of vascular fusariose is avoided.

#### Huang, X., J. Luo, et al. (2009). "Isolation and bioactivity of endophytic fungi in Derris hancei." Journal of South China Agricultural University 30(2): 44-47.

Derris hancei Hemsl. The antagonism of endophytic fungi against fungal pathogens was tested in vitro. Penicillium sp. Q1, Rhizoctonia sp. S1, Phomopsis sp. N2, and Corticium sp. F1 isolated from the caudex of D. hancei, and Penicillium sp. Q2 isolated from the leaf, inhibited the hyphal growth of Colletotrichum gloeosporioides Penz, Fusarium oxysporum f. niveum (E. F. Smith) Snyber et Hansen, Rhizoctonia sp. S1 against Colletotrichum orbiculare Arx, and Phomopsis sp. N2 against Colletotrichum musae (Berk1 & Curt1) Arx1 on dual culture with inhibition index II. It was reported that endophytic fungus in D.hancei could produced antibacterial substances in this paper. The culture filtrates of Penicillium sp. Q2 treated in 48 h after treatment possessed 100.00% of adjusted mortality against the 2nd larvae of Spodoptera litura by leaves disc feeding bioassays, and 75.10% against Lipaphis erysimi Kaltenbach (apterous adult) by insect-soaking method, respectively, which showed that the activity of Penicillium sp. Q2 was higher than that of other endophytic fungi.

Jadeja, K. B. and D. M. Nandoliya (2008). "Integrated management of wilt of cumin (Cuminum cyminum L.)." Journal of Spices and Aromatic Crops 17(3): 223-229.

pathogen, Fusarium oxysporum f. sp. lycopersici though the effect was well pronounced with consortium developed from Santalum album.

Four components of integrated management namely, soil solarization, crop rotation, chemicals and biocontrol agents were tested under field condition at Junagadh (Gujarat) for the management of wilt of cumin (Cuminum cyminum) caused by Fusarium oxysporum f. sp. cumini. Growing of sorghum (Sorghum bicolor) or maize (Zea mays) during kharif season did not reduce wilt incidence during the following rabi season. Soil solarization with 25 m LLDPE plastic cover for 15 days in summer proved most effective in reducing wilt incidence to 26.27% as against 44.90% in non-solarization and increasing yield to 396 kg ha-1 as against 286 kg ha-1 in non-solarized plots. Application of carbendazim granules @ 10 kg ha-1 one month after sowing or Trichoderma viride in organic carrier @ 62.5 kg ha-1 at sowing time were also effective. Integrating soil solarization followed by growing of sorghum in kharif and application of either carbendazim granules @ 10 kg ha-1 one month after sowing or application of T.viride in organic carrier @ 62.5 kg ha-1 was effective for the management of cumin wilt.

Kamilova, F., S. Validov, *et al.* (2009). Biological control of tomato foot and root rot caused by Fusarium oxysporum f.sp. radicis-lycopersici by Pseudomonas bacteria. Proceedings of the Second International Symposium on Tomato Diseases, Kusadasi, Turkey, 8-12 October 2007.

Rhizobacteria are a natural and most suitable source for the isolation of potential microbiological control agents that can protect plants from soilborne pathogens and consequently improve crop quality and yield. The beneficial effect of such bacteria on plant health depends in many cases on their ability to aggressively colonize the rhizosphere and compete with the indigenous, including pathogenic, microflora for nutrients and niches on the plant root. Bacterial strains Pseudomonas chlororaphis PCL1391 and P. fluorescens WCS365 employ antibiosis and induced systemic resistance, respectively, to control tomato foot and root rot (TFRR) caused by phytopathogenic fungus Fusarium oxysporum f.sp. radicis-lycopersici (Forl). For the selection of biocontrol bacteria acting via the mechanism "competition for nutrients and niches" we have developed an enrichment method for enhanced tomato root tip colonizers, starting from a crude mixture of rhizobacteria coated on the seed, using a sterile quartz sand/plant nutrient solution gnotobiotic system. As a result of this enrichment procedure, and subsequent tests on competitive tomato root tip colonization, the strongly competitive biocontrol strains P. fluorescens PCL1751 and P. putida PCL1760 were isolated. Both strains effectively suppress TFRR under soil and hydroponic cultivation conditions.

Kamilova, F., S. Validov, *et al.* (2009). "Biological control of tomato foot and root rot caused by Fusarium oxysporum f.sp. radicis-lycopersici by Pseudomonas bacteria." Acta Horticulturae(808): 317-320. isolation of potential microbiological control agents that can protect plants from soilborne pathogens and consequently improve crop quality and yield. The beneficial effect of such bacteria on plant health depends in many cases on their ability to aggressively colonize the rhizosphere and compete with the indigenous, including pathogenic, microflora for nutrients and niches on the plant root. Bacterial strains Pseudomonas chlororaphis PCL1391 and P. fluorescens WCS365 employ antibiosis and induced systemic resistance, respectively, to control tomato foot and root rot (TFRR) caused by phytopathogenic fungus Fusarium oxysporum f.sp. radicis-lycopersici (Forl). For the selection of biocontrol bacteria acting via the mechanism "competition for nutrients and niches" we have developed an enrichment method for enhanced tomato root tip colonizers, starting from a crude mixture of rhizobacteria coated on the seed, using a sterile quartz sand/plant nutrient solution gnotobiotic system. As a result of this enrichment procedure, and subsequent tests on competitive tomato root tip colonization, the strongly competitive biocontrol strains P. fluorescens PCL1751 and P. putida PCL1760 were isolated. Both strains effectively suppress TFRR under soil and hydroponic cultivation conditions.

Kannan, V. and R. Sureendar (2009). "Synergistic effect of beneficial rhizosphere microflora in biocontrol and plant growth promotion." Journal of Basic Microbiology 49(2): 158-164. Biological systems are getting more relevance than chemical control of plant pathogens as they are not only eco-friendly and economic in approach but are also involved in improving the soil consistency and maintenance of natural soil flora. Plant growth promoting rhizosphere microorganisms were isolated from three different tree rhizospheres using selective culture media. Five microorganisms were selected from each rhizosphere soil based on their efficiency and screened for their ability to promote plant growth as a consortium. Each of the developed consortium has a phosphate solubilizer, nitrogen fixer, growth hormone producer, heterotrophic member and an antagonist. The plant growth promoting ability of the microbial members present in the consortium was observed by estimating the IAA production level and also by the nitrogenase activity of the nitrogen fixers. The biocontrol potentiality of the consortium and the antagonist present in the consortium were checked by both dual plate assay and cross-streaking technique. Consortial treatments effected very good growth promotion in Lycopersicon esculentum Mill and the treated plants also developed resistance against wilt

Li, J., Q. Yang, *et al.* (2009). "Evaluation of biocontrol efficiency and security of A Bacillus subtilis strain B29 against cucumber Fusarium wilt in field." China Vegetables(2): 30-33. cucumerinum, was isolated from cucumber rhizosphere. After twice of 4-field-plot experiments, the control efficiencies of 100, 250 and 500 dilution times to cucumber Fusarium wilt were 70.3-88.2%, 62.3-85.9%, and 54.7-80.6%, respectively. The average efficiency of field trials with B29 was 84.9% during 2 years and the yield of cucumber increased by 12.57%. The acute toxicity of Bacillus subtilis strain B29 to big mouse through its mouth and skin was examined, and the LD50 was more than 5000 mg/kg. The application of strain B29 on cucumber, tomato, bean and seed pumpkin was safe based on the observed seedling rate, growth and development.

#### Liu, Q., J. C. Yu, et al. (2009). "Antagonism and Action Mechanism of Antifungal Metabolites from Streptomyces rimosus MY02." Journal of Phytopathology 157(5): 306-310.

The genus of Streptomyces, a saprophytic Gram-positive bacterium, has properties, which make them useful as pharmaceutical and biocontrol agents. A streptomyces strain MY02 from soil samples showed significant antagonism against 14 plant pathogenic fungi including Fusarium oxysporum f. sp. cucumarinum. Antifungal metabolite(s) SN06 from the culture of the strain MY02 were extracted with n-butanol and purified by silica gel column chromatography. The minimum concentration of SN06 inhibiting any visible fungal growth of F. oxysporum f. sp. cucumarinum is 12.5 mu g/ml by twofold serial dilutions method. The mycelia of F. oxysporum f. sp. cucumarinum treated with SN06 were observed under the normal optics microscope. The results showed that some cells of hyphae began to dilate and formed some strings of beads. The cytoplasm oozed out of the cells with the culture time and so most of the cells became empty. The hyphae broke into many segments and then collapsed after 48 h. After inoculated in potato dextrose medium for 48 h, the filtrate of mycelia treated with 1% NaCl containing 12.5 mu g/ml SN06 was scanned using ultraviolet spectrophotometer and absorption peak at 260 nm showed that the mycelia cell membrane of F. oxysporum f. sp. cucumarinum was broken and that nucleic acid oozed out of the cell.

#### Maina, M., R. Hauschild, et al. (2008). "Protection of tomato plants against fusaric acid by resistance induction." Journal of Applied Biosciences(JABs) 1: 18-31.

Objectives: The rhizobacteria Bacillus sphaericus B43, Pseudomonas fluorescens T58, and P. putida 53 are able to induce systemic resistance (ISR) against Fusarium oxysporum f.sp. lycopersici (FOL) in tomato. This study investigated if the ISR reduced the damage by the toxin fusaric acid (FA) produced by FOL. Methodology and Results: The bacteria were applied to the rhizosphere of tomato plants. Chlorophyll content and ion leakage were determined after placing the leaf discs in FA. Active oxygen species (AOS), superoxide and hydrogen peroxide levels were determined in leaves of plants injected with FA. Activities of superoxide dismutase (SOD), ascorbate (AS) and guaiacol peroxidases (GPX) involved in AOS metabolism were quantified. In untreated plants, FA led to high ion leakage and chlorophyll degradation caused by H2O2 accumulation. All the bacteria treatments decreased the chlorophyll degradation. Ion leakage was reduced by treatment with P. fluorescens T58 and B. sphaericus B43, while P. putida 53 was less effective. Treatment of plants with bacteria resulted in increased superoxide contents, but varying over time. Increased SOD and GPX activities in untreated plants were suppressed after bacteria treatment. Plants treated with P. fluorescens T58 showed only a transient increase in superoxide. P. putida 53-treated plants removed AOS, but high initial superoxide levels led to membrane damages. Treatment with B. sphaericus B43 suppressed the effects of FA, but AOS metabolism showed only slight alterations. Conclusions and potential applications of findings: ISR could also protect plant tissues from damage by pathogen toxins, which is a potential new dimension to the known mechanisms of action of biological control agents.

Martinez-Medina, A., J. A. Pascual, *et al.* (2009). "Interactions between arbuscular mycorrhizal fungi and Trichoderma harzianum and their effects on Fusarium wilt in melon plants grown in seedling nurseries." Journal of the Science of Food and Agriculture 89(11): 1843-1850.

BACKGROUND: Biological control through the use of Trichoderma spp. and arbuscular mycorrhizal fungi (AMF) could contribute to a reduction of the inputs of environmentally damaging agrochemical products. The objective of this study was to evaluate the interactions between four AMF (Glomus intraradices, Glomus mosseae, Glomus claroideum and Glomus constrictum) and Trichoderma harzianum for their effects on melon plant growth and biocontrol of Fusarium wilt in seedling nurseries. RESULTS: AMF colonisation decreased fresh plant weight, which was unaffected by the presence of T. harzianum. Dual inoculation resulted in a decrease in fresh weight compared with AMF-inoculated plants, except for G. intraradices. AMF colonisation level varied with the AM endophyte and was increased by T. harzianum, except in G. mosseae-inoculated plants. Negative effects of AMF on T. harzianum colony-forming units were found, except with G. intraradices. AMF alone were less effective than T. harzianum in suppressing disease development. Combined inoculation resulted in a general synergistic effect on disease control. CONCLUSION: Selection of the appropriate AMF species and its combination with T. harzianum were significant both in the formation and effectiveness of AM symbiosis and the reduction of Fusarium wilt incidence in melon plants. The combination of G. intraradices and T. harzianum provided better results than any other tested. (C) 2009 Society of Chemical Industry

#### Matar, S. M., S. A. El-Kazzaz, et al. (2009). "Antagonistic and inhibitory effect of Bacillus subtilis against certain plant pathogenic fungi, I." Biotechnology 8(1): 53-61.

subtilis isolates (B1 to B14), obtained from different Egyptian sites, were tested against six fungal isolates belonging to four different genera, Rhizoctonia solani, Helminthosporium spp., Alternaria spp. and Fusarium oxysporum. Cultural, morphological and physiological characteristics of these isolates were found to be identical to B. subtilis. Four B. subtilis isolates (B1, B4, B7, B8) had more antagonistic effect on all fungal isolates. Supernatant of B. subtilis isolate B7 had antagonistic effect on 6 fungal isolates but it was more effective on Helminthosporium spp., Alternaria spp. and F. oxysporum. B. subtilis as well as isolate B7 showed effectiveness in reducing disease incidence and severity levels of tomato plants when added to the F. oxysporum and R. solani-infested soil. Also, it stimulated the growth of tomato plants compared to the other. HPLC analysis of the HCl precipitate of B.subtilis isolate B7 culture supernatant revealed that an identical pattern of five peaks to that of a purified preparation of iturin A was obtained.

Matar, S. M., S. A. El-Kazzaz, *et al.* (2009). "Bioprocessing and scaling-up cultivation of Bacillus subtilis as a potential antagonist to certain plant pathogenic fungi, III." Biotechnology 8(1): 138-143. isolate G-GANA7 (GenBank accession No. EF583053), collected from Abo-Homos in Egypt, was tested against six fungal isolates belonging to four different genera, i.e. Rhizoctonia solani, Helminthosporium sp., Alternaria sp. and Fusarium oxysporum. B. subtilis isolate G-GANA7 was cultured in 3 litre bench-top New Brunswick Scientific BioFlow III bioreactor for producing the maximum yield of biomass and antifungal compound. Fed-batch processes were automated through a computer aided data bioprocessing system AFS-BioCommand multi-process management program

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to regulate the cell growth rate by controlling interactively the nutrient feed rate, temperature, pH and agitation speed based on dissolved oxygen. In batch cultivation, the process suffered from low yield of cell mass (3.2 g litre-1) and antifungal activity because of high initial glucose concentration followed by acetate formation which the causal agent for inhibition of cell growth. Constant and exponential fed-batch strategies were adopted to circumvent this potential problem. Fed-batch cultivation of B. subtilis was conducted at the specific growth rate of 0.13 and 0.1 h-1 for constant and exponential strategies, respectively. High cell density of 12.8 and 14.6 g litre-1 for both operations, with an overall biomass yield of 0.45 g g-1 was achieved. The inhibitory activity of antifungal in supernatant reached its maximum value of 2 and 2.2 cm for constant and exponential fed-batch cultivations.

- Mazurier, S., T. Corberand, *et al.* (2009). "Phenazine antibiotics produced by fluorescent pseudomonads contribute to natural soil suppressiveness to Fusarium wilt." ISME Journal 3(8): 977-991. Natural disease-suppressive soils provide an untapped resource for the discovery of novel beneficial microorganisms and traits. For most suppressive soils, however, the consortia of microorganisms and mechanisms involved in pathogen control are unknown. To date, soil suppressiveness to Fusarium wilt disease has been ascribed to carbon and iron competition between pathogenic Fusarium oxysporum and resident non-pathogenic F. oxysporum and fluorescent pseudomonads. In this study, the role of bacterial antibiosis in Fusarium wilt suppressiveness was assessed by comparing the densities, diversity and activity of fluorescent Pseudomonas species producing 2,4-diacetylphloroglucinol (DAPG) (phID+) or phenazine (phzC+) antibiotics. The frequencies of phID+ populations were similar in the suppressive and conducive soils but their genotypic diversity differed significantly. However, phID genotypes from the two soils were equally effective in suppressing Fusarium wilt, either alone or in combination with non-pathogenic F. oxysporum strain Fo47. A mutant deficient in DAPG production provided a similar level of control as its parental strain, suggesting that this antibiotic does not play a major role. In contrast, phzC+ pseudomonads were only detected in the suppressive soil. Representative phzC+ isolates of five distinct genotypes did not suppress Fusarium wilt on their own, but acted synergistically in combination with strain Fo47. This increased level of disease suppression was ascribed to phenazine production as the phenazine-deficient mutant was not effective. These results suggest, for the first time, that redox-active phenazines produced by fluorescent pseudomonads contribute to the natural soil suppressiveness to Fusarium wilt disease and may act in synergy with carbon competition by resident non-pathogenic F. oxysporum.
- Minerdi, D., S. Bossi, *et al.* (2009). "Volatile organic compounds: a potential direct long-distance mechanism for antagonistic action of Fusarium oxysporum strain MSA 35." Environmental Microbiology 11(4): 844-854.

Fusarium oxysporum MSA 35 [wild-type (WT) strain] is an antagonistic Fusarium that lives in association with a consortium of bacteria belonging to the genera Serratia, Achromobacter, Bacillus and Stenotrophomonas in an Italian soil suppressive to Fusarium wilt. Typing experiments and virulence tests provided evidence that the F. oxysporum isolate when cured of the bacterial symbionts [the cured (CU) form], is pathogenic, causing wilt symptoms identical to those caused by F. oxysporum f. sp. lactucae. Here, we demonstrate that small volatile organic compounds (VOCs) emitted from the WT strain negatively influence the mycelial growth of different formae speciales of F. oxysporum. Furthermore, these VOCs repress gene expression of two putative virulence genes in F. oxysporum lactucae strain Fuslat10, a fungus against which the WT strain MSA 35 has antagonistic activity. The VOC profile of the WT and CU fungus shows different compositions. Sesquiterpenes, mainly caryophyllene, were present in the headspace only of WT MSA 35. No sesquiterpenes were found in the volatiles of ectosymbiotic Serratia sp. strain DM1 and Achromobacter sp. strain MM1. Bacterial volatiles had no effects on the growth of the different ff. spp. of F. oxysporum examined. Hyphae grown with VOC from WT F. oxysporum f. sp. lactucae strain MSA 35 were hydrophobic whereas those grown without VOCs were not, suggesting a correlation between the presence of volatiles in the atmosphere and the phenotype of the mycelium. This is the first report of VOC production by antagonistic F. oxysporum MSA 35 and their effects on pathogenic F. oxysporum. The results obtained in this work led us to propose a new potential direct long-distance mechanism for antagonism by F. oxysporum MSA 35 mediated by VOCs. Antagonism could be the consequence of both reduction of pathogen mycelial growth and inhibition of pathogen virulence gene expression.

Nam, M. H., M. S. Park, et al. (2009). "Biological Control of Strawberry Fusarium Wilt Caused by Fusarium oxysporum f. sp fragariae Using Bacillus velezensis BS87 and RK1 Formulation." Journal of Microbiology and Biotechnology 19(5): 520-524.

Two isolates, Bacillus sp. BS87 and RK1, selected from soil in strawberry fields in Korea, showed high levels of antagonism towards Fusarium oxysporum f. sp. fragariae in vitro. The isolates were identified as B. velezensis based on the homology of their gyrA sequences to reference strains. BS87 and RK1 were evaluated for control of Fusarium wilt in strawberries in pot trials and field trials conducted in Nonsan, Korea. In the pot trials, the optimum applied concentration of BS87 and RK1 for pre-plant root-dip application to control Fusarium wilt was 10(5) and 10(6) colony-forming units (CFU)/ml, respectively. Meanwhile, in the 2003 and 2005 field trials, the biological control efficacies of formulations of RK1 were similar to that of a conventional fungicide (copper hydroxide) when compared with a non-treated control. The RK1 formulation was also more effective than BS87 in suppressing Fusarium wilt under field conditions. Therefore, the results indicated that formulations of B. velezensis BS87 and RK1 may have potential to control Fusarium wilt in strawberries.

Narayan, M., P. Tini, et al. (2009). "Biological and chemical management of tomato wilt caused by Fusarium oxysporum f.sp. lycopersici." Journal of Soils and Crops 19(1): 118-121.

Wilt of tomato is one of the most important known disease caused by Fusarium oxysporum f. sp. lycopersici. In the present study four bioagents (Trichoderma harzianum, T. viride, Bacillus subtilis and Pseudomonas fluorescens) and two fungicides (Carbendazim and Thiram) were evaluated both in vitro and in vivo conditions. In vitro evaluation, of Carbendazim (0.1%) completely inhibited the growth

of tomato wilt pathogen Fusarium oxysporum f.sp. lycopersici and was found significantly superior over the rest of fungicides. While, among the biological agents Trichoderma viride was found significantly superior to the rest in checking the growth of pathogens and showed 85.69 per cent inhibition. In vivo under field condition, seedling dip treatment of Carbendazim (1 gl-1 water) was found most significant followed by Carbendazim+ T.viride (1+100 gl-1 water) and T. viride (100 gl-1 water) significantly reduced wilt incidence by 73.91, 69.56 and 68.11 per cent respectively as against 71.88 per cent wilting in control (under epiphytotic condition i.e. wilt sick soil).

- Ortega-Morales, B. O., F. N. Ortega-Morales, *et al.* (2009). "Antagonism of Bacillus spp. Isolated from Marine Biofilms Against Terrestrial Phytopathogenic Fungi." Marine Biotechnology 11(3): 375-383. We aimed at determining the antagonistic behavior of bacteria derived from marine biofilms against terrestrial phytopathogenic fungi. Some bacteria closely related to Bacillus mojavensis (three isolates) and Bacillus firmus (one isolate) displayed antagonistic activity against Colletotrichum gloeosporioides ATCC 42374, selected as first screen organism. The four isolates were further quantitatively tested against C. gloeosporioides, Colletotrichum fragariae, and Fusarium oxysporum on two culture media, potato dextrose agar (PDA) and a marine medium-based agar [yeast extract agar (YEA)] at different times of growth of the antagonists (early, co-inoculation with the pathogen and late). Overall antagonistic assays showed differential susceptibility among the pathogens as a function of the type of culture media and time of colonization (P < 0.05). In general, higher suppressive activities were recorded for assays performed on YEA than on PDA; and also when the antagonistic activity (P < 0.05) were found between the different isolates. In general, Bacillus sp. MC3B-22 displayed a greater antagonistic effect than the commercial biocontrol strain Bacillus subtilis G03 (KodiakA (R)). Further incubation studies and scanning electronic microscopy revealed that Bacillus sp. MC3B-22 was able to colonize, multiply, and inhibit C. gloeosporioides ATCC 42374 when tested in a mango leaf assay, showing its potential for fungal biocontrol. Additional studies are required to definitively identify the active isolates and to determine their mode of antifungal action, safety, and biocompatibility.
- Padghan, P. R. and M. M. Baviskar (2009). "Efficacy of bioagent and different root extracts for supression of chickpea wilt in vitro." Asian Journal of Bio Science 4(1): 56-58. udid, sorghum bicolor), groundnut and mung bean and biological control agents (Trichoderma viride, T. harzianum, T lignorum and T. koningii) against the chickpea wilt pathogen, Fusarium oxysporum f.sp. ciceris (FOC), was studied in the laboratory. A lower radial mycelial growth and a higher inhibitory effect were recorded in sorghum root extract medium (28.00 mm and 54.34%), respectively, however, it was at par with groundnut root extract medium (30.00 mm and 51.08%), compared to the control (61.33 mm). In dual culture technique, the growth of FOC was restricted by T. viride (56.16%), followed by T. harzianum (50.57%). T. lignorum recorded the minimum zone of inhibition (40.45%).
- Qiu, W., H. Huang, *et al.* (2009). "Screening of actinomycete against Fusarium oxysporum f. sp. cubense and identification of strain DA07408." Research of Agricultural Modernization 30(1): 126-128. samples, and 8 of these strains showed significant activities against F. oxysporum f.sp. cubense. One actinomycete (DA07408) isolated from an arboretum in Danzhou, Hainan, China, exhibited marked antagonism towards F. oxysporum f.sp. cubense. The conditions for the fermentation of the actinomycete were optimized. Based on the morphological, physiological and biochemical characteristics of the strain, and on the analysis of 16S rDNA and phylogenetic tree, DA07408 was identified as Streptomyces olivochromogenes.
- Raddadi, N., A. Belaouis, *et al.* (2009). "Characterization of polyvalent and safe Bacillus thuringiensis strains with potential use for biocontrol." Journal of Basic Microbiology 49(3): 293-303. Sixteen Bacillus thuringiensis (Bt) strains were screened for their anti-insect, antibacterial and antifungal determinants by phenotypic tests and PCR targeting major insecticidal proteins and complements, chitinases, lactonases, beta-1,3-glucanases and zwittermicin A. Six strains had genes of at least two major insecticidal toxins and of insecticidal complements. With regard to fungal biocontrol, all the strains inhibited Fusarium oxysporum and Aspergillus flavus growth and four strains had all or most of the antifungal determinants examined, with strain Bt HD932 showing the widest antifungal activity spectrum. Autolysins, bacteriocin and AHL-lactonases were produced by all or most of the tested strains with different activity spectra including pathogens like Listeria monocytogenes. Safety evaluation was carried out via PCR by screening the B. cereus psychrotolerance-related genes, toxin genes and the virulence pleiotropic regulator plcR. Diarrheal enterotoxins and other toxin genes were widespread among the collection with strains Bt HD9 and H45 lacking psychrotolerance-related genes, while five strains were positive. Only three strains (BMG1.7, H172, H156) resulted positive with primer sets targeting partial or complete plcR gene. By Vero Cell Assays, Bt HD868 followed by Bt HD9 were shown to be the safest strains. These polyvalent and safe Bt strains could be very promising in field application.
- Rasal, P. H., J. R. Shelar, et al. (2009). "Effect of endophytic antagonist on pigeonpea." Journal of Maharashtra Agricultural Universities 34(1): 52-53.

resistant (ICP 8863) and resistant (BDN2) cultivars of pigeon pea were screened against Fusarium oxysporum f. udum [F. udum]. The inoculation of endophytic antagonists into different cultivars of pigeon pea improved germination, plant height, branching, nodulation, root length and biomass production, and reduced wilt intensity significantly over the un-inoculated control. Among the inoculants, Pseudomonas-2 was the most beneficial, followed by Pseudomonas-3, Bacillus-3, Pseudomonas-1, and Bacillus-1 and -2. Antagonists isolated from resistant cultivar were the most beneficial, followed by antagonists from the moderately resistant cultivar, and antagonists isolated from the susceptible cultivar.

## Nicot et al. (Appendix for Chapter 1)

#### Recep, K., S. Fikrettin, et al. (2009). "Biological control of the potato dry rot caused by Fusarium species using PGPR strains." Biological Control 50(2): 194-198.

In this study, a total of 17 Plant Growth Promoting Rhizobacteria (PGPR) strains, consisting of eight different species (Bacillus subtilis, Bacillus pumilus, Burkholderia cepacia, Pseudomonas putida, Bacillus amyloliquefaciens, Bacillus atrophaeus, Bacillus macerans and Flavobacter balastinium), were tested for antifungal activity in in vitro (on Petri plate) and in vivo (on potato tuber) conditions against Fusarium sambucinum, Fusarium oxysporum and Fusarium culmorum cause of dry rot disease of potato. All PGPR strains had inhibitory effects on the development of at least one or more fungal species on Petri plates. The strongest antagonism was observed in B. cepacia strain OSU-7 with inhibition zones ranging from 35.33 to 47.37 mm. All PGPR strains were also tested on tubers of two potato cultivars 'Agria' and 'Granola' under storage conditions. Only B. cepacia strain OSU-7 had significant effects on controlling potato dry rot caused by three different fungi species on the two potato cultivars. There were no significant differences in rot diameters among the treatments in comparison to the negative control (with water). This is the first study showing that B. cepacia has great potential to be used as effective biocontrol agent of Fusanium dry rot of potatoes (F. oxysporum and F culmorum) under storage conditions. (C) 2009 Elsevier Inc. All rights reserved.

#### Riaz, T., S. N. Khan, et al. (2009). "Effect of co-cultivation and crop rotation on corm rot disease of Gladiolus." Scientia Horticulturae 121(2): 218-222.

Field and pot experiments were conducted to evaluate the effect of co-cultivation and crop rotation on the growth and corm rot disease of gladiolus (Gladiolus grandiflorus sect. Blandus) cv. Aarti caused by Fusarium oxysporum f.sp. gladioli (Massey) Snyd. and Hans. In the field experiment, gladiolus was co-cultivated with 10 agricultural/horticultural crops viz. Allium cepa L., Brassica campestris L., Capsicum annuum L., Eruca sativa Mill., Helianthus annuus L., Tagetes erectus L., Zea mays L., Vinca rosea L. and Rosa indica L., in a soil infested with F. oxysporum. All the crops except V. rosea and R. indica reduced disease incidence. The effect of H. annuus and T. erectus was significant and more pronounced than other co-cultivated crops. In general, root and shoot dry biomass, corm fresh weight, number of cormlets and number of flowers per spike decreased as compared to the un-inoculated monoculture gladiolus treatment (negative control) but these parameters enhanced as compared to the F. oxysporum inoculated monoculture gladiolus treatment (positive control). In a pot experiment, all the crops of the field experiment except V. rosea and R. indica were sown in rotation with gladiolus. Pot grown plants of different species were harvested at maturity and the soil was inoculated with F oxysporum. Gladiolus was cultivated I week after inoculation. Disease incidence was significantly suppressed in all the treatments ranging from 29% to 53%. The highest suppression of disease incidence was recorded in T erectus (53%) followed by B. campestris (49%). The effect of preceding crops on various vegetative parameters was similar in the pot experiment to that of the field experiment. The present study suggests that corm rot disease of gladiolus can be managed by mixed cropping of H. annuus and T erectus or cultivation of T. erectus and B. campestris in rotation. (c) 2009 Elsevier B.V. All rights reserved.

Saidi, N., S. Kouki, et al. (2009). "Characterization and selection of Bacillus sp strains, effective biocontrol agents against Fusarium oxysporum f. sp radicis-lycopersici, the causal agent of Fusarium crown and root rot in tomato." Annals of Microbiology 59(2): 191-198.

The antagonistic activities of 20 Bacillus isolates were tested with dual culture and greenhouse conditions against Fusarium oxysporum f. sp. radicis-lycopersici (FORL) race 0, the causal agent of Fusarium crown and root rot of tomato. Under dual culture, 10 isolates inhibited mycelial growth > 38% and the most effective inhibited fungal growth > 50%. The 20 Bacillus isolates were tested for production of volatiles, cyanide, antibiotics, and phosphorus solubilisation; 15 isolates produced volatiles that inhibited growth of pathogens, 9 isolates produced cyanide, 10 produced antibiotics, and five solubilised phosphorus. Greenhouse experiments with the same 20 isolates revealed the effectiveness of 12 strains, which increased the percentage of healthy plants in the tested cultivar from 66 to 96%. The best disease control was achieved by isolates B11, B5, B17, and B18. However, B11 and B17 were the only isolates that produced cyanide, antibiotics, solubilised phosphate and showed 44% inhibition of fungal growth. The selected strains could be considered in plant growth promotion and biological disease control.

#### Shi, Y. W., K. Lou, et al. (2009). "Isolation, quantity distribution and characterization of endophytic microorganisms within sugar beet." African Journal of Biotechnology 8(5): 835-840.

The present investigation was undertaken in order to document the spectrum of endophytes colonizing healthy leaves of sugar beet cultivars in Xinjiang Province (China) and to determine the degree of colonization at three growth stages. From the 360 sugar beet leaf and root segments incubated, 221 bacterial isolates, 34 fungal isolates and 5 actinomycete isolates were obtained. Of all the isolates, 7 bacterial species and 6 fungal species were identified. The actinomycete isolates were characterized as Streptomyces griseofuscus and Streptomyces globisporus. There were significant differences between microorganisms, stages of growth, and stages of microorganism interaction. The number of microorganisms isolated increased during the growth period of the sugar beet. At the same time, the number of microorganisms affecting different parts of the sugar beet tissue was quite different. The greatest number of microorganisms was found in the secondary root emergence zone of the sugar beet tissue. Endophytic microorganisms in sugar beet promote growth and increase the yield of the beet.

Son, S. H., Z. Khan, *et al.* (2009). "Plant growth-promoting rhizobacteria, Paenibacillus polymyxa and Paenibacillus lentimorbus suppress disease complex caused by root-knot nematode and fusarium wilt fungus." Journal of Applied Microbiology 107(2): 524-532.

Paenibacillus strains against disease complex caused by Meloidogyne incognita and Fusarium oxysporum f. sp. lycopersici interactions. Methods and Results: Paenibacillus strains were collected from rotten ginseng roots. The strains were tested under in vitro and pots for their inhibitory activities, and biocontrol potential against disease complex caused by M. incognita and F. oxysporum f. sp. lycopersici on tomato. In in vitro experiments, among 40 tested strains of Paenibacillus spp., 11 strains showed antifungal and nematicidal activities against F. oxysporum f. sp. lycopersici and M.

incognita, respectively. Paenibacilluspolymyxa GBR-462; GBR-508 and P. lentimorbus GBR-158 showed the strongest antifungal and nematicidal activities. These three strains used in pot experiment reduced the symptom development of the disease complex (wilting and plant death), and increased plant growth. The control effects were estimated to be 90-98%, and also reduced root gall formation by 64-88% compared to the untreated control. Conclusion: The protective properties of selected Paenibacillus strains make them as potential tool to reduce deleterious impact of disease complex plants. Significance and Impact of the Study: The study highlights biocontrol potential of Paenibacillus strains in management of disease complex caused by nematode-fungus interaction.

Srinivasan, K., G. Gilardi, et al. (2009). "BACTERIAL ANTAGONISTS FROM USED ROCKWOOL SOILLESS SUBSTRATES SUPPRESS FUSARIUM WILT OF TOMATO." Journal of Plant Pathology 91(1): 147-154.

Five bacterial E,trains (FC-6B, FC-7B, FC-8B, FC-9B and FC-24B) isolated from used rockwool soilless substrates were identified using 16S ribosomal DNA (16S rDNA) sequence analysis as belonging to the Pseudomonas genus. Seven glasshouse trials were conducted in order to evaluate the efficacy of these bacteria strains (Pseudomonas putida FC-6B, Pseudomonas sp. FC-7B, Pseudomonas sp. FC-9B and Pseudomonas sp. FC-24B) together with Achromobacter sp. AM1 and Serratia sp. DM1 obtained from suppressive sod, against Fusarium wilt of tomato. Two commercial bioproducts, Trichoderma harzianum T22 (RootShield) and Pseudomonas chlororaphis MA 342 (Cedomon) were also evaluated. Different treatment strategies including soil application (10(7) and 10(8) cfu ml(-1)) were adopted in different glasshouse trials (Trial I to VI) to test the efficacy of the bacterial strains against Fusarium wilt. Root dipping was used in Trial VII (10(8) and 10(9) cfu ml(-1)). The lowest: disease incidence (3.3) was recorded with a single application of P. putida FC-6B at 10(8) cfu ml(-1). Similar results were obtained with the same bacteria when the concentration was decreased to 10(7) cfu ml(-1) but an increasing number of applications was required. The highest plant biomass (50.3 g/plant) was recorded in the P. putida FC-8B treatment (Trial III). In conclusion, the current study showed the potential biocontrol activity of bacterial strains FC-6B, FC-7B, FC-8B, FC-9B and FC-24B isolated from re-used rockwool soilless substrates against Fusarium wilt disease, and the growth promoting activity of these strains on tomato plants.

Srivastava, D. K., A. K. Singh, et al. (2009). "Efficacy of bio-control agents and seed dressing fungicides against damping off of tomato." Annals of Plant Protection Sciences 17(1): 257-258. in Unao, Madhya Pradesh, India, during 2005-06 yielded associated pathogen on PDA medium. The antagonistic activity of biological control agents against Fusarium oxysporum f.sp. lycopersici was determined using dual culture method. All the antagonists and fungicide inhibited the mycelial growth of Fusarium, however, Trichoderma viride caused maximum inhibition of mycelial growth. Trichoderma viride, Trichoderma harzianum, Gliocladium virens, carbendazim and thiram, which showed significant in vitro inhibition of Fusarium were tested in the field. Maximum increase in seed germination (83.4%), seedling survival (79.0) and plant height (6.32 cm) over the control was observed when treated with Trichoderma viride followed by Trichoderma harzianum, carbendazim, thiram, and Gliocladium virens.

Thanh, D. T., L. T. T. Tarn, *et al.* (2009). "Biological Control of Soilborne Diseases on Tomato, Potato and Black Pepper by Selected PGPR in the Greenhouse and Field in Vietnam." Plant Pathology Journal 25(3): 263-269.

Bacterial wilt, Fusarium wilt and Foot rot caused by Ralstonia solanacearum, Fusarium oxysporum, and Phytophthora capsici respectively, continue to be severe problems to tomato, potato and black pepper growers in Vietnam. Three bio-products, Bacillus vallismortis EXTN-1 (EXTN-1), Bacillus sp. and Puenibacillus sp. (ESSC) and Bacillus substilis (MFMF) were examined in greenhouse bioassay for the ability to reduce bacterial wilt, fusarium wilt and foot rot disease severity. While these bio-products significantly reduced disease severities, EXTN-1 was the most effective, providing a mean level of disease reduction 80.0 to 90.0% against bacterial wilt, fusarium wilt and foot rot diseases under greenhouse conditions. ESSC and MFMF also significantly reduced fusarium wilt, bacterial wilt and foot rot under field condition at Song Phuong and Thuong Tin locations in Ha Tay province, Vietnam. Under field condition, EXTN-1 provided a mean level of disease reduction more than 45.0% against all three diseases compared to water treated control. Besides, EXTN-1 treatment increased the yield in tomato fruits 17.3% than water treated control plants.

Wu, H., X. Yang, et al. (2009). "Suppression of Fusarium wilt of watermelon by a bio-organic fertilizer containing combinations of antagonistic microorganisms." BioControl 54(2): 287-300. the crop has been grown for many seasons. Its occurrence results in a severely decreased watermelon crop. The goal of this study was to assess the capability of a new product (bio-organic fertilizer) to control the wilt in Fusarium-infested soil. Pot experiments were conducted under growth chamber and greenhouse conditions. The results showed that the fertilizer controlled the wilt disease. Compared with control pots, the incidence rates of Fusarium wilt at 27 and 63 days following treatment of the plants with the bio-organic fertilizer at a rate of 0.5% (organic fertilizer+antagonistic microorganisms, including 3\*109 CFU g-1 respectively, in both the growth chamber and greenhouse settings. The activities of antioxidases (catalase, superoxide dismutase and peroxidase) in watermelon leaves increased by 38.9, 150 and 250%, respectively. In the roots, stems and leaves, the activity of beta-1,3-glucanase (pathogenesis-related proteins) increased by 80, 1140 and 100% and that of chitinase increased by 240, 80, and 20%, respectively, while the contents of malondialdehyde fell by 56.8, 42.1 and 45.9%, respectively. These results indicate that this new fertilizer formula is capable of protecting watermelon from Fusarium oxysporum f.sp. niveum. The elevated levels of defense-related enzymes are consistent with the induction and enhancement of systemic acquired resistance of plant.

## Nicot et al. (Appendix for Chapter 1)

- Wu, Q., H. Zeng, *et al.* (2009). "Stability of fermentation broth of actinomycete strain WZ162 resistance to Fusarium oxysporum f.sp. cubense of banana." Guangxi Agricultural Sciences 40(4): 366-369. The fermentation broth of actinomycete strain WZ162 has strong inhibiting effect against Fusarium oxysporum f.sp. cubense of banana. Under different conditions, the stabilities of fermentation broth of WZ162 were detected. The results showed that the fermentation broth of WZ162 had better heat stability when temperature of water bath was below 80C. The antibiotics ingredient of fermentation broth would not be changed and can maintain the antifungal activity under conditions of sun light and ultraviolet rays. Under acid and neutrality conditions, the inhibition rate of fermentation broth against Focr4 was 24.92%-34.73% and 11.21%-25.39%, respectively. Therefore, the stability of fermentation broth in acid was better than that of neutrality. When the fermentation broth with pH 1-12 were treated with different time in 100C water bath, the inhibition rate was obviously lower than that of the treatments without water bath, and the stability of fermentation broth with pH 1 was the best.
- Yin, X., D. Chen, et al. (2009). "An endophytic Erwinia chrysanthemi strain antagonistic against banana fusarium wilt disease." Chinese Journal of Biological Control 25(1): 60-65. An endophytic strain E353 was obtained from the pseudostem of healthy banana plant in a field heavily infected with Fusarium oxysporum f. sp. cubense (FOC). Antagonism of the strain against FOC was tested via dual-culture, inhibition test on conidia germination, and pot trials. Results showed that E353 effectively inhibited mycelium growth and conidia germination. Efficacy of strain E353 to control the wilt disease was 60.67% in pot trials. Strain E353 was identified as Erwinia chrysanthemi according to its characteristics in morphology, physiology, biochemistry and 16S rDNA sequence.
- Zhong, X., M. Liang, *et al.* (2009). "Study on the inhibition of Trichoderma sp. against Fusarium oxysporum f. sp. cubense in banana." Journal of Fruit Science 26(2): 186-189. effective antagonist against Fusarium oxysporum f. sp. cubens, was isolated and identified as Trichoderma sp. based upon 18S rDNA gene analysis. With solid and liquid cultures, the inhibitive efficacy to the growth of Fusarium oxysporum f. sp. cubens was primarily studied. The experimental results showed that the cells of Fusarium oxysporum f. sp. cubens were completely covered by short fiber-like hyphace and spore stem of G2 within 7 days in the dual culture plate, and in the antagonist plate, the average rate of inhibitory by the culture solution of G2 was about 90.4%, the average rate of the inhibitory by volatile substance reached 68.3%. After 10 days' incubation with 20% (v/v) fungal strain G2, the melt of the pathogenic mycel and spore were observed in the liquid culture containing 1.0\*107 cfu . L-1 G2 can strongly inhibit the growth of Fusarium oxysporum f. sp. cubens.
- Zhu, H., Y. Ma, *et al.* (2009). "Control effect of combining biocontrol strains against Fusarium oxysporium f. sp. niveum and Verticillium dahlae." Journal of Northwest A & F University Natural Science Edition 37(7): 152-156.

Objective: Five actinomycetes strains having certain inhibiting capability were screened as material to study the control effect of the actinomycetes and five combinations on watermelon Fusarium wilt and Eggplant Verticillium wilt, and to filter the combining biocontrol strains which have better biocontrol efficacy and growth promotion. Method: The biocontrol efficacy and growth promotion of single and combining strains were analyzed by antagonistic activity in vitro and manual inoculation in vivo. Result: Strain SC11 and SE2 had significant inhibiting effect on Fusarium oxysporium f. sp. niveum and Verticillium dahlae in vitro. Inhibiting rate on conidia germination was also high; in greenhouse experiment, 84.93% control ratio to Fusarium oxysporium f. sp. niveum and 71. 48% to Verticillium dahlae were found by C2; The fermentation broth of C3 had the most significant effect for every index of watermelon. The effect on reduction intensity of watermelon rootage was obvious. For eggplant, the growth promotion was only inferior to strain SF6. Conclusion: These results suggested that the control effect and growth promotion of combining biocontrol strains are significantly higher than individual, and combining strains express complementary biocontrol activities by collaboration. There is no correlation between the number of strains and control effect, only proper combinations of biocontrol strains can enhance disease control effect.

# Appendix 7. Number of references retrieved by using the CAB Abstracts database in order to review scientific literatures on augmentative biological control in selected crops for Chapter 2.

Key words	1973-2008	1998-2008	
Biological control	1644	-	
Augmentative biological control	7	6	
Augmentation biological control	10	6	
Inoculative biological control	4	1	
Inundative biological control	7	3	
Insects biological control	773	373	
Mites biological control	320	190	
Total references dealing with	607	579	
augmentative biocontrol to be examined			

\* Survey includes records for grapevine, grape and vineyard.

#### APPLE

Key words	1973-2008	1998-2008
Biological control	3971	-
Augmentative biological control	13	10
Augmentation biological control	18	9
Inoculative biological control	5	3
Inundative biological control	10	2
Insects biological control	2310	817
Mites biological control	981	258
Total references dealing with	1145	1099
augmentative biocontrol to be examined		

PEAR

Key words	1973-2008	1998-2008
Biological control	1270	-
Augmentative biological control	3	2
Augmentation biological control	2	1
Inoculative biological control	1	1
Inundative biological control	3	1
Insects biological control	756	325
Mites biological control	174	61
Total references dealing with	400	391
augmentative biocontrol to be examined		

CORI		
Key words	1973-2008	1998-2008
Biological control	6828	-
Augmentative biological control	19	14
Augmentation biological control	38	18
Inoculative biological control	18	8
Inundative biological control	39	17
Insects biological control	4293	1682
Mites biological control	250	66
Total references dealing with	1919	1805
augmentative biocontrol to be examined		
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# CORN\*

\* Survey include records for **corn** and **maize**.

## WHEAT

Key words	1973-2008	1998-2008	
Biological control	5250	-	
Augmentative biological control	9	7	
Augmentation biological control	13	6	
Inoculative biological control	1	1	
Inundative biological control	8	3	
Insects biological control	2307	866	
Mites biological control	157	66	
Total references dealing with	980	949	
augmentative biocontrol to be examined			

# CARROT

Key words	1973-2008	1998-2008
Biological control	360	-
Augmentative biological control	1	1
Augmentation biological control	1	1
Inoculative biological control	1	1
Inundative biological control	0	0
Insects biological control	179	62
Mites biological control	20	8
Total references dealing with	76	73
augmentative biocontrol to be examined		

# ONION

Key words	1973-2008	1998-2008
Biological control	810	-
Augmentative biological control	2	2
Augmentation biological control	3	3
Inoculative biological control	3	3
Inundative biological control	1	1
Insects biological control	532	233
Mites biological control	187	62
Total references dealing with augmentative biocontrol to be examined	313	304

# Appendix 8. Collection of data on augmentative biological control of pests in grapevine. Each table refers to a group of biocontrol agents.

References	Species of biocontrol agent	Species of insect pest	Taxonomic category of pests	Country	Type of augmentation	Type of test	Efficacy of biocontrol agents*	Additional information and results
Remund & Bigler, 1986	T. dendrolimi	<i>Eupoecilia ambiguella</i> (grape berry moth)	Lepidoptera: Tortricidae			Lab		Evaluation of biological parameters
				Switzerland		Field		Evaluation of biological parameters
	T. maidis			Switzerland	Inundative	Field	+	
Segonca & Leisse, 1989	T. semblidis	Eupoecilia ambiguella and Lobesia botrana	Lepidoptera: Tortricidae	Ahr Valley, Germany	Inundative	Field	+	
Glenn & Hoffmann, 1997	T. carverae	<i>Epiphyas postvittana</i> (light brown apple moth)	Lepidoptera: Tortricidae	Victoria, Australia	Inundative	Field (small blocks)	+	
Basso <i>et al.</i> , 1998	T. pretiosum T. exiguum	Argyrotaenia sphaleropa (South American tortricid moth), Bonagota cranaodes (Brasilian apple leafroller)	Lepidoptera: Tortricidae	Uruguay		Lab		Evaluation of biological parameters
Basso et al., 1999	T. pretiosum T. exiguum	A. sphaleropa B. cranaodes	Lepidoptera: Tortricidae	Uruguay	Inundative	Field	+	
Garnier-Geoffroy et al., 1999	T. brassicae	Lobesia botrana	Lepidoptera: Tortricidae			Lab	-	Evaluation of allelocemical relations
Hommay et al., 2002	<i>T. evanescens</i> and <i>T. cacoeciae</i> (two strains)	Lobesia botrana	Lepidoptera: Tortricidae	France	Inundative	Field	+ -	<ul><li>+ as % parasitization.</li><li>- as % grapes attacked.</li></ul>
Nagargatti et al., 2002	T. minutum	<i>Endopiza viteana</i> (grape berry moth)	Lepidoptera: Tortricidae	Pennsylvania, USA		Field	+	+ as natural parasitism. Inundative releases of <i>T</i> . <i>minutum</i> in border rows is suggested
Thomson & Hoffmann, 2002	T. carverae	<i>Epiphyas postvittana</i> (light brown apple moth)	Lepidoptera: Tortricidae	Victoria, Australia		Lab Field		Assessment of quality indicators
Nagargatti et al., 2003	T. minutum	Endopiza viteana	Lepidoptera: Tortricidae	Pennsylvania, USA	Inundative	Field	+	Parasitoids released in border rows
Zimmermann, 2004	Trichogramma spp.	Lobesia botrana and Eupoecilia ambiguella	Lepidoptera: Tortricidae	Germany	Inundative	Field		Commercialized to be used in home garden
Begum et al., 2006	T. carverae	Epiphyas postvittana	Lepidoptera: Tortricidae	Australia	Inundative	Greenho use/ Field	+	Ground-cover plant species identified to improve performance of mass released parasitoids.

# **Giorgini** (Appendix for Chapter 2)

El-Wakeil <i>et al.</i> , 2008 <i>T. evanescens</i>	<i>Lobesia botrana</i> (European grape berry moth)	Lepidoptera: Tortricidae	Egypt	Inundative	Field	+	Parasitism > 97% and reduction percents of infestation reached 96.8%
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\* + means effective, - means not effective biocontrol agent.

#### 8.2 Parasitoid Hymenoptera: Encyrtidae [4 species], Pteromalidae [1 species]

Reference	Species of biocontrol agent	Species of insect pest	Taxonomic category of pests	Cuontry	Type of augmentation	Type of test	Efficacy of biocontrol agents*	Additional information and results
Walton & Pringle, 1999	Coccidoxenoides peregrinus (Encyrtidae)	Planococcus ficus (vine mealybug)	Hemiptera: Pseudococcidae	South Africa		Lab		Compatibility of fungicides and incompatibility of insecticides with augmentative releases
Walton & Pringle, 2004	Coccidoxenoides perminutus (Encyrtidae)	Planococcus ficus (vine mealybug)	Hemiptera: Pseudococcidae	South Africa	Inundative	Field	+	Mass release was at least as effective as the chemical control
Abd-Rabou, 2005	Anagyrus kamali (Encyrtidae)	Maconellicoccus hirsutus	Hemiptera: Pseudococcidae	Egypt	Inundative	Field	+	It is concluded that the releases of parasitoids were suitable for control.
Daane et al., 2006	Anagyrus pseudococci (Encyrtidae)	Planococcus ficus	Hemiptera: Pseudococcidae	California	Inoculative	Field	+	Promising results. Commercial products are not yet available.
Daane et al., 2008	Anagyrus pseudococci (Encyrtidae)	Planococcus ficus	Hemiptera: Pseudococcidae	Israel	Inoculative	Field	+	Promising results. Commercial products are not yet available.
Kapongo et al., 2007	Muscidifurax raptor (Pteromalidae)	<i>Ceratitis capitata</i> (Mediterranean fruit fly)	Diptera: Tephritidae	Canada	Inundative	Field Lab cages	+	<i>M. raptor</i> constitutes a promising biocontrol agent in vineyards.

\* + means effective, - means not effective biocontrol agent.

#### 8.3 Predators of mites. Acari: Phytoseidae.

References	Species of biocontrol agent	Species of mite pest	Taxonomic category of pests	Country	Type of augmentation	Type of test	Efficacy of biocontrol agents*	Additional information and results
Boller <i>et al.</i> , 1988	Typhlodromus pyri	Panonychus ulmi, Tetranychus urticae	Acari: Tetranychidae	Switzerland	Inoculative	Field		Inoculative release of <i>T. pyri</i> along with the increase of the internal ecological diversity achieved by proper management of the green cover plants will have a strong influence on predator densities.

Camporese & Duso, 1996	Typhlodromus pyri, Amblyseius andersoni, Kampimodromus aberrans	Panonychus ulmi	Acari: Tetranychidae	Italy	Inoculative	Field	+	Different colonization patterns on three grape varieties (with different pubescent leaf undersurfaces). The high competitiveness of <i>K</i> . <i>aberrans</i> over the other 2 phytoseid species is a major factor in selecting predatory species for inoculative releases.
Takahashi et al., 1998	Phytoseiulus persimilis	Tetranychus kanzawai	Acari: Tetranychidae	Japan	Inundative	Field (grape in green house)	+	Release of <i>P. persimilis</i> onto the grass ground cover in the spring. No chemical control was required.
Duso & Vettorazzo, 1999	Kampimodromus aberrans, Typhlodromus pyri	Panonychus ulmi, Eotetranychus carpini Calepitrimerus vitis	Acari: Tetranychidae Acari: Eriophyidae	Veneto, Italy	Inoculative	Field (A)	+	Releases were successful and the predators became more abundant on the variety with pubescent leaf under-surface. Native <i>A. andersoni</i> were displaced by <i>T. pyri</i> .
						Field (B)	+	Two grape varieties with different leaf hair density. <i>T. pyri</i> colonization failed; <i>K.</i> <i>aberrans</i> was more successful on glabrous varieties. <i>K. aberrans</i> displaced native <i>P. finitimus</i> .
Marshall & Lester, 2001	Typhlodromus pyri	Panonychus ulmi	Acari: Tetranychidae	Ontario, Canada	Inoculative	Field	+	<i>T. pyri</i> out-competed native <i>Amblyseius fallacies.</i> <i>T. pyri</i> is an effective biocontrol agent and may be introduced by transferring leaves.
Duso <i>et al.</i> , 2006	Typhlodromus pyri strain resistant to organophosphates	Panonychus ulmi, Eotetranychus carpini Calomerus vitis	Acari: Tetranychidae Acari: Eriophyidae	North-eastern Italy	Inoculative	Field		15-years observations. The predator colonized the vineyard and competed successfully with other species. Role of alternative foods, leaf morphology and selective pesticides.

\* + means effective, - means not effective biocontrol agent.

# **Giorgini (Appendix for Chapter 2)**

## 8.4 Predators of insects. Neuroptera: Chrysopidae [3 species] and Coleoptera: Coccinellidae [2 species]

Reference	Species of biocontrol agent	Species of insect pest	Taxonomic category of pests	Country	Type of augmentation	Type of test	Efficacy of biocontrol agents*	Additional information and results
	NEUROPTERA: CHRYSOPIDAE							
Daane et al., 1996	Chrysoperla carnea (common green lacewing)	<i>Erythroneura</i> <i>variabilis</i> , <i>E. elegantula</i> (leafhoppers)	Hemiptera: Cicadellidae	California	Inundative	Field (caged small- plot)	-	Average leafhopper density reduction 29.5%.
						Field (uncaged small-plot)	-	Release rates reflecting commercial recommendations. Average reduction 15.5%.
						Field (on-farm trials)	-	Average reduction 9.6% Not sufficient to lower the leafhopper density below the economic injury threshold.
Daane & Yokota, 1997	Chrysoperla carnea, C. comanche, C. rufilabris	<i>Erythroneura</i> <i>variabilis</i> , <i>E. elegantula</i> (leafhoppers)	Hemiptera: Cicadellidae	California	Inundative	Field	-	Aspects of release strategies evaluated. High mortality of lacewing eggs and neonate larvae.
Wunderlich & Giles, 1999	Chrysoperla rufilabris	<i>Erythroneura</i> <i>variabilis</i> , <i>E. elegantula</i> (leafhoppers)	Hemiptera: Cicadellidae	California	Inundative	Field		A mechanical technique was assessed for releasing eggs in liquid suspensions. Adhesion of eggs to the canopy was an issue.
	COLEOPTERA: COCCINELLIDAE							
Anagnou et al., 2003	Nephus includens	Planococcus citri	Hemiptera: Pseudococcidae	Greece		Field		It is suggested, for combined infestation by <i>L. botrana</i> and mealybugs, the application of <i>B. thuringiensis</i> and the releases of the effective predator <i>N.</i> <i>includens</i> .
Daane <i>et al.</i> , 2008	Cryptolaemus montrouzieri	Pseudococcus maritimus, P. longispinus (mealybugs)	Hemiptera: Pseudococcidae	California	Inoculative	Field		Commonly released in vineyards, but release rates, timing, and expected outcomes have not been scientifically evaluated. It may be best used by releasing at hot spots where the mealybug density is high.
Mani, 2008	Cryptolaemus montrouzieri	Planococcus citri	Hemiptera: Pseudococcidae	India	Inundative	Green house	+	

\* + means effective, - means not effective biocontrol agent.

# 8.5 Fungi [5 species]

Reference	Species of biocontrol agent	Species of insect pest	Taxonomic category of pests	Country	Type of augmentation	Type of test	Efficacy of biocontrol agents*	Additional information and results
Berner & Schnetter, 2002	<i>Beauveria brongniartii</i> (in combination with the nematode <i>H.</i> <i>bacteriophora</i> )	Melolontha melolontha (European cockchafer)	Coleoptera: Scarabeidae	Germany	Inundative	Field (soil)	+	Only under optimum conditions and with high doses control of the white grubs could be reached.
Tsitsipis <i>et al.</i> , 2003	Beauveria bassiana	<i>Frankliniella</i> <i>occidentalis</i> (western flower thrips)	Thysanoptera: Thripidae	Greece	Inundative	Field	+	<ul> <li><i>B. bassiana</i> in combination with mass trapping was compared to mass trapping or insecticides.</li> <li>Less efficient in the control of insect population if compared to some chemicals.</li> </ul>
Al-Jboory <i>et al.</i> , 2006	Beauveria bassiana	grape thrips	Thysanoptera: Thripidae	Iraq		Lab	+	Two isolates of <i>B. bassiana</i> showed 100% mortality after 5 days
Lopes et al., 2002	Metarhizium anisopliae	Frankliniella occidentalis	Thysanoptera: Thripidae	Brazil	Inundative	Field	+	The effect of chemicals (thiacloprid and methiocarb) with or without <i>M.a.</i> was tested. <i>M.a.</i> in combination with methiocarb was the best strategy.
Laengle <i>et al.</i> , 2004	Metarhizium anisopliae	<i>Daktulosphaira</i> <i>vitifoliae</i> (grape phylloxera)	Hemiptera: Phylloxeridae	Austria	Inundative	Field		Non-target effects on soil fauna: no negative effects detected.
Kirchmair <i>et al.</i> , 2004	Metarhizium anisopliae	Daktulosphaira vitifoliae (grape phylloxera)	Hemiptera: Phylloxeridae	Austria	Inundative	Lab	+	<i>M.a.</i> was effective in pot experiments. Potential role of <i>M.a.</i> in grape phylloxera control.
Kirchmair <i>et al.</i> , 2005	Metarhizium anisopliae	Daktulosphaira vitifoliae (grape phylloxera)	Hemiptera: Phylloxeridae	Germany	Inundative	Field	+	<i>M.a.</i> was effective. No target effects on soil fauna (Acari, Collembola, Lumbricida and the Carabidae <i>Harpalus affinis</i> ) and fungi.
Huber & Kirchmair, 2007	Metarhizium anisopliae	Daktulosphaira vitifoliae (grape phylloxera)	Hemiptera: Phylloxeridae	Germany	Inundative	Field	-	Evaluation of efficacy: more difficulties arise in testing the efficacy of <i>M.a.</i> under field conditions because of the uneven distribution of roots and pest insects in the soil.

# Giorgini (Appendix for Chapter 2)

Kirchmair <i>et al.</i> , 2007	Metarhizium anisopliae	Daktulosphaira vitifoliae (grape phylloxera)	Hemiptera: Phylloxeridae	Germany	Inundative	Field	+	3 months after application an increase of the <i>M.a.</i> density in soil was observed. Compared with untreated plots a lower infestation was observed in the <i>M.a.</i> - treated plots. Two years after treatment a control effect was still observed whereas the density of <i>M.a.</i> in soil decreased. Three years after treatment no effect on the pest was detectable and the <i>M.a.</i> density had decreased to a value similar to that in the control . A periodically application is necessary.
Maheshkumar- Katke & Balikai.	Metarhizium anisopliae, Verticillium lecanii,	Maconellicoccus hirsutus	Hemiptera: Pseudococcidae	India	Inundative	Field	+	
	,		rseudococcidae					
2008	Clerodendron inerme	(grape mealybug)						

\* + means effective, - means not effective biocontrol agent.

#### 8.6 Nematodes [5 species]

Reference	Species of biocontrol agent	Species of insect pest	Taxonomic category of pests	Country	Type of augmentation	Type of test	Efficacy of biocontrol agents*	Additional information and results
Saunders & All, 1985	Steinernema carpocapsae	<i>Vitacea polistiformis</i> (grape root borer)	Lepidoptera: Sesiidae	Georgia, USA	Inundative (soil)	Lab, Field	+	Susceptibility of <i>V.p.</i> 1st-instar larvae. Augmentation of nematode populations during the critical period of <i>V.p.</i> oviposition and eclosion is suggested as a control technique.
English-Loeb et al., 1999	Heterorhabditis bacteriophora (Oswego strain), Steinernema glaseri (isolate 326)	Daktulosphaira vitifolia (grape phylloxera - root form)	Hemiptera: Phylloxeridae	NY, USA		Lab	+	<ul> <li>H. bacteriophora: reduced survival of attached phylloxera by up to 80%.</li> <li>S. glaseri had no measurable impact. No evidence that H.b. could successfully reproduce within the bodies of the hosts.</li> <li>Augmentative use in the field in an release programme may be constrained by the need to use high densities, their dependence on moist soils, and their inability to propagate themselves within hosts.</li> </ul>

Berner & Schnetter, 2002	Heterorhabditis bacteriophora, H.bacteriophora + Beauveria brongniartii (fungus)	Melolontha melolontha (European cockchafer)	Coleoptera: Scarabeidae	Germany	Inundative (soil)	Field	+	Only under optimum conditions and with high doses of nematodes control of grubs could be reached. New variant for the application of nematodes proposed.
Williams <i>et al.</i> , 2002	Heterorhabditis bacteriophora, H. zealandica, H. marelata, and Steinernema carpocapsae	<i>Vitacea polistiformis</i> (grape root borer)	Lepidoptera: Sesiidae	Ohio, USA	Inundative	Lab Greenh ouse	+	<i>H. bacteriophora</i> strains GPS11 and Oswego, <i>H. zealandica</i> strain X1, and <i>H. marelata.</i> <i>S. carpocapsae</i> strain All less effective <i>H. zealandica</i> strain X1 <i>H. bacteriophora</i> strain GPS11

\* + means effective, - means not effective biocontrol agent.

#### 8.7 Bacillus thuringiensis

Reference	<i>B. thuringiensis</i> subspecies	Species of Insect pest	Taxonomic category of pests	Country	Type of test	Efficacy	Additional results and information
Caroli et al., 1998	subsp. aizawai	<i>Lobesia botrana</i> (grape berry moth)	Lepidoptera: Tortricidae	Emilia- Romagna, Italy	Field	+	90-95% reduction in damage against severe pest infestations comparable to the standard chemical products.
Keil & Schruft, 1998		L. botrana, Eupoecilia ambiguella (grape berry moths)	Lepidoptera: Tortricidae		Lab		4 Bt products (0.2% Bactospeine FC, 0.1 % Delfin, 0.1% Dipel ES and 0.1% Thuricide HP) were compared. The influence of temperature on the efficacy is discussed.
Morando et al., 1998		L. botrana, E. ambiguella	Lepidoptera: Tortricidae	Piemonte, Italy	Field	+	The efficacy of Bt was compared to 7 insecticides. All the tested insecticides had a significantly good efficacy.
Boselli et al., 2000		L. botrana	Lepidoptera: Tortricidae	Emilia- Romagna, Italy	Field		Bt compared to insecticides.
Fretay & Quenin, 2000		L. botrana	Lepidoptera: Tortricidae	France	Field		Evaluation of new formularions.
Bagnoli & Lucchi, 2001	subsp. <i>kurstaki</i>	<i>Cryptoblabes</i> <i>gnidiella</i> (honey moth)	Lepidoptera : Pyralidae	Toscana, Italy	Field	+	
Boselli & Scannavini, 2001	subsp. <i>kurstaki</i> subsp. <i>aizawai</i>	L. botrana	Lepidoptera: Tortricidae	Emilia- Romagna, Italy	Field		Treatments included Agree (Bt kurstaki and aizawi), flufenoxuron, chlorpyrifos, lufenuron, tebufenozide, methoxyfenozide, indoxacarb and spinosad. The best <b>control</b> was obtained with methoxyfenozide, indoxacarb, and spinosad.
Neves & Frescata, 2001	kurstaki x aizawai	L. botrana	Lepidoptera: Tortricidae	Bairrada, Portugal	Field	+	TUREX was tested to control the <i>L. botrana</i> third generation. Great interest of this Bt product regarding its efficiency and persistence based in a correct spray moment determination.

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Anagnou et al., 2003	subsp. <i>kurstaki</i> subsp. <i>aizawai</i>	L. botrana	Lepidoptera: Tortricidae		Lab	+	Several products incorporated into an artificial diet resulted in >90% larval mortality. The same formulations did not significantly affect the survival of <i>Nephus includens</i> .
Ifoulis & Savopoulou- Soultani, 2003		L. botrana	Lepidoptera: Tortricidae	Greece	Field	+	Two formulations of Bt are significantly more effective than the control, the dusting being more effective in most cultivars and the spraying in a few cultivars.
Roditakis, 2003		L. botrana	Lepidoptera: Tortricidae	Greece	Field		Pest control strategy involves <i>B.t.</i> application, mating disruption, botanical insecticides and minimal use of insecticides
Samoilov, 2003		Sparganothis pilleriana (grape leafroller)	Lepidoptera: Tortricidae	Odessa, Ukraine	Field	+	
Bakr, 2004	subsp. <i>kurstaki</i>	Lobesia botrana	Lepidoptera: Tortricidae	Egypt	Field	+	The addition of sugar as a feeding stimulant to a 50% reduced rate of Dipel-2X resulted in higher control rates (80%) compared to using the recommended field rates of Dipel-2X alone or Actellic [pirimiphos-methyl].
Besnard et al., 2004	subsp. <i>aizawai</i>	Lobesia botrana	Lepidoptera: Tortricidae	France	Field	+	Xen Tari commercial product.
Hera et al., 2004	subsp. <i>kurstaki</i>	Hyphantria cunea (fall webworm)	Lepidoptera: Arctiidae	Romania	Field	+	Dipel 2x WP at 0.075% also showed good protection. The synergenism of mixtures (50:50) of chemical and biological insecticides was effective in controlling the pest.
Laccone et al., 2004	subsp. <i>kurstaki</i>	Lobesia botrana	Lepidoptera: Tortricidae	Calabria, Italy	Field	+	Bt gave satisfactory control if applied at the onset of ovideposition and provided the canopy was managed in such a way as to expose the berries.
Mazzocchetti <i>et al.</i> , 2004		Lobesia botrana	Lepidoptera: Tortricidae	Abruzzo, Italy	Field		Mating disruption was compared with the traditional methods generally used in the area: chemicals (phosphorganic molecules) and <i>B. thuringiensis</i> .
Moiraghi et al., 2004		L. botrana E. ambiguella	Lepidoptera: Tortricidae	Italy	Field	-	In four years, trials were carried out using several commercial products (9 insecticides and Bt). The best <u>control</u> was obtained using insecticides. <u>Control</u> was lower for azadirachtin and less constant for etofenprox and B. thuringiensis.
Delbac <i>et al.</i> , 2006		Lobesia botrana	Lepidoptera: Tortricidae	France	Field	+	<i>L. botrana</i> was well-controlled by the use of <i>B.t.</i> or IGR's, without mating disruption justification
Marchesini et al., 2006	subsp. <i>aizawai</i> subsp. <i>kurstaki</i>	Lobesia botrana	Lepidoptera: Tortricidae	Veneto, Italy	Field	+	<i>Bta</i> compared to <i>Btk</i> and chemicals. High efficacy of <i>B.t. aizawai</i> .
Laccone, 2007		Lobesia botrana	Lepidoptera: Tortricidae	Molise and Calabria, Italy	Field		Pest control with indoxacarb, spinosad and <i>B. thuringiensis</i> applied against the 2nd generation of insects parasitizing fruit is also outlined
Mescalchin, 2007		Lobesia botrana	Lepidoptera: Tortricidae	Trentino, Italy	Field	+	5-years study (2000-2005). Formulations based Bt can be used for controlling tortricids such as <i>L. botrana</i> .
Mitrea <i>et al.</i> , 2007	subsp. kurstaki	Lobesia botrana	Lepidoptera: Tortricidae	Romania	Field	+	Chemical insecticides followed by <i>Btk</i> to control the second or the third generation. Efficiency of the control treatments ranged between 89.4% and 91.4%.

Morandi-Filho <i>et al.</i> , 2007		Argyrotaenia sphaleropa (South American tortricid moth)	Lepidoptera: Tortricidae	Brazil	Lab Field	+ +	Lab: reducition of the insect population by more than 90%. Field: reduced damage between 83.3 and 94.4%. The control efficacy of B.t was equal to that of chemicals.
Pryke & Samways, 2007	subsp. <i>kurstaki</i>	<i>Epichoristodes</i> <i>acerbella</i> (South African carnation tortrix)	Lepidoptera: Tortricidae	South Africa	Field	+	DiPelReg commercial formulation
Ruiz-de-Escudero et al., 2007		Lobesia botrana	Lepidoptera: Tortricidae		Lab	+	The potential of Bt Cry proteins to control L. botrana was explored. Either Cry1Ia or Cry9C could be used in combination with Cry1Ab to control this pest, either as the active components of Bt sprays or expressed together in transgenic plants.
Subic, 2007	subsp. kurstaki	Lobesia botrana	Lepidoptera: Tortricidae	Croatia	Field	+	Over 90% control was achieved.
Dongiovanni <i>et al.</i> , 2008	subsp. kurstaki	Lobesia botrana	Lepidoptera: Tortricidae	Puglia, Italy	Field	+	

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# Appendix 9. References on classical biological control against insect pests (cited in Chapter 3)

Type of work	Pest (genus level)	<b>References*</b>	
Prospective studies (55%)		(88)	
-	Aproaerema	(89)	
	Cameraria	(61)	
	Cryptococcus	(175) (94)	
	Diabrotica	(154)	
	Hypsipyla	(141)	
	Liriomyza	(87)	
	Lymanthria	(70)(72)	
	Scirtothrips	(45)	
	Tetranychus		
Retrospective studies (35%)		(166)	
-	Chilo	(128)	
	Cinara	(56)	
	Cosmopolites	(103)	
	Maconellicoccus	(47)	
	mealybugs	(191)	
	Mononychellus	(97)	
	Phenacoccus		
Other studies (10%)		(82)	
Pest biology	Enarmonia	(88)	

#### 9.1. Biocontrol agents not precisely known

\* Numbers correspond to refernces presented in section 9.4

Pest	BCA lifestyle	BCA	References*
Aceria	Fungus	Hirsutella	(114)
	Predatory mite	Neoseiulus	
Adelges	Predatory Insect	Laricobius	(119)
Anticarsia	Virus	Nucleopolyhedrovirus	(197)
Aphids	Predatory Insect	Harmonia	(48) (127)
Aphis	Fungus	Neozygites	(19) (90) (91) (137)
Coptotermes	Fungus	Beauvaria & Metarhizium	(168)
Lymantria	Fungus	Microspora	(35)
	Virus	Nucleopolyhedrovirus	
Maconellicoccus	Predatory Insect	Cryptolaemus	(165)
		Scymnus	
Mononychellus	Fungus	Neozygites	(16)
	Predatory mite	Neosiulus &Typhlodromalus	
Oryctes	Virus	_	(51) (86)
Prostephanus	Predatory Insect	Teretrius	(51)
Review	Fungus	_	(14) (39) (42) (43)
Review	Nematode	_	(14) (55) (124) (125) (193) (194)
Sirex	Nematode	Deladenus	(81)
Solenopsis	Fungus	Vairimorpha	(73) (169) (170)

9.2. Details on the use of pathogens, nematodes and predators as agents of Classical Biological Control

\* Numbers correspond to refernces presented in section 9.4

#### 9.3 Categorization of publications related to Insect parasitoids as CIBCA according to the type of work

#### **Pest Biology**

Pest rearing : (83, 183)

#### **BCA Biology**

BCA inventories : (30, 34, 65) (67) (88) (157) (178) BCA systematics: (18, 52, 123) (36) (186) BCA molecular characterization: (121, 132) BCA rearing: (21, 58, 92, 163) (171) BCA biology: (6, 10, 37) (74) (77) (85) (98) (100) (102) (104) (105) (158) (159) (160) (172) (190) (195) BCA Evaluation: (12, 44, 46) (57) (80) (108) (151)

#### **BCA Field Implications**

Pre-release survey: (9, 60, 66) (122) (140) (166) BCA introduction : see table 1 Post-release survey : (20, 22, 32) (33) (36) (50) (54) (64) (68) (76) (78) (106) (107) (113) (109) (135) (142) (145) (146) (148) (150) (162) (179)

#### Non-intended effects

(24, 29, 38) (58) (71) (84) (92) (65) (101) (129) (149) (155) (184) (189)

#### **Biocontrol disruption**

(17, 27, 69) (95) (130) (147) (180)

#### Miscellaneous

Economic valuation: (23) Review: (75, 112, 152) (153) Miscellaneous: (111, 115, 116) (139) (176) "Conservation BC-like": (173)

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	Substance	Cipac & incl 2008/ 127 √	Category	List (*)	Inclusion Date	Expiry Date	Legislation
Botanical	Extract from tea tree		RE	A 4	01/09/2009	31/08/2019	2008/127
Botanical	Garlic extract		RE	A 4	01/09/2009	31/08/2019	2008/127
Botanical	Gibberellic acid	'307	PG	A 4	01/09/2009	31/08/2019	2008/127
Botanical	Gibberellin		PG	A 4	01/09/2009	31/08/2019	2008/127
Botanical	Laminarin		EL	С	01/04/2005	31/03/2015	<u>05/3/EC</u>
Botanical	Pepper		RE	A 4	01/09/2009	31/08/2019	2008/127
Botanical	Plant oils / Citronella oil		HB	A 4	01/09/2009	31/08/2019	2008/127
Botanical	Plant oils / Clove oil		RE	A 4	01/09/2009	31/08/2019	2008/127
Botanical	Plant oils / Rape seed oil		IN, AC	A 4	01/09/2009	31/08/2019	2008/127
Botanical	Plant oils / Spearmint oil		PG	A 4	01/09/2009	31/08/2019	2008/127
Botanical	Sea-algae extract (formerly sea- algae extract and seaweeds)		PG	A 4	01/09/2009	31/08/2019	2008/127
Botanical copied by synthesis	Carvone		PG	С	01/08/2008	31/07/2018	2008/44/EC
Botanical copied by synthesis	Ethylene		PG	A 4	01/09/2009	31/08/2019	<u>2008/127</u>
Botanical but excluded	Pyrethrins	'32	IN	A 4	01/09/2009	31/08/2019	2008/127
Chemical	2,4-D	'1	HB, PG	A 1	01/10/2002	30/09/2012	<u>01/103/EC</u>
Chemical	2,4-DB	'83	HB	A 1	01/01/2004	31/12/2013	<u>03/31/EC</u>
Chemical	1-Methyl-cyclopropene		PG	С	01/04/2006	31/03/2016	<u>06/19/EC</u>
Chemical	Acetamiprid		IN	С	01/01/2005	31/12/2014	<u>04/99/EC</u>
Chemical	Acibenzolar-S-methyl (benzothiadiazole)		PA	С	01/11/2001	31/10/2011	<u>01/87/EC</u>
Chemical	Aclonifen	'498	HB	A 3	01/08/2009	31/07/2019	<u>2008/116</u>
Chemical	Alpha-Cypermethrin (aka alphamethrin)	'454	IN	A 1	01/03/2005	28/02/2015	<u>04/58/EC</u>
Chemical	Aluminium ammonium sulfate		RE	A 4	01/09/2009	31/08/2019	2008/127
Chemical	Aluminium phosphide	'227	IN, RO	A 3	01/09/2009	31/08/2019	2008/125
Chemical	Amidosulfuron	'515	HB	A 3	01/01/2009	31/12/2018	2008/40
Chemical	Amitrole (aminotriazole)	'90	HB	A 1	01/01/2002	31/12/2012	<u>01/21/EC</u>
Chemical	Azimsulfuron		HB	С	01/10/1999	01/10/2019	<u>99/80/EC</u>
Chemical	Azoxystrobin		FU	С	01/07/1998	01/07/2008	<u>98/47/EC</u>
Chemical	Beflubutamid		HB	С	01/12/2007	30/11/2017	<u>07/50/EC</u>
Chemical	Benalaxyl	'416	FU	A 1	01/03/2005	28/02/2015	<u>04/58/EC</u>
Chemical	Benfluralin	'285	HB	A 3	01/01/2009	31/12/2018	<u>2008/108</u>
Chemical	Bensulfuron	'502	HB	A 3	01/11/2009	31/10/2019	2009/11
Chemical	Bentazone	'366	HB	A 1	01/08/2001	31/07/2011	<u>00/68/EC</u>
Chemical	Benthiavalicarb		FU	С	01/08/2008	31/07/2018	08/44/EC
Chemical	Beta-Cyfluthrin	'482	IN	A 1	01/01/2004	31/12/2013	<u>03/31/EC</u>
Chemical	Bifenazate		AC	С	01/12/2005	30/11/2015	<u>05/58/EC</u>
Chemical	Bifenox	'413	HB	A 3	01/01/2009	31/12/2018	2008/66
Chemical	Bordeaux mixture		FU	A 3	01/11/2009	30/11/2016	SCoFCAH voted 01.2009
Chemical	Boscalid		FU	С	01/08/2008	31/07/2018	08/44/EC
Chemical	Bromoxynil	'87	HB	A 1	01/03/2005	28/02/2015	<u>04/58/EC</u>

# Appendix 10. Substances included in the "EU Pesticides Database" as of April 21 2009

# Heilig et al (Appendix for Chapter 4)

Chemical	Calcium carbide	r	RE	A 4	01/09/2009	31/08/2019	2008/127
Chemical		'505	RO	A 4 A 3	01/09/2009	31/08/2019	2008/127
Chemical	Calcium phosphide	303 '40	FU	_	01/09/2009	31/08/2019 30/09/2017	
Chemical	Captan Carbendazim	263	FU FU	A 2		30/09/2017 31/12/2009	07/5/EC 06/135/EC
Chemical		203	FU HB	A 1	01/01/2007 01/10/2003	31/12/2009	
	Carfentrazone-ethyl	1111		C			<u>03/68/EC</u>
Chemical	Chloridazon (aka pyrazone)	'111	HB	A 3	01/01/2009	31/12/2018	<u>2008/41</u>
Chemical	Chlormequat (chloride)	'143	PG	A 3	01/12/2009	30/11/2019	05/50/50
Chemical	Chlorothalonil	'288	FU	A 1	01/03/2006	28/02/2016	<u>05/53/EC</u>
Chemical	Chlorotoluron	'217	HB	A 1	01/03/2006	28/02/2016	<u>05/53/EC</u>
Chemical	Chlorpropham	'43	PG, HB	A 1	01/02/2005	31/01/2015	<u>04/20/EC</u>
Chemical	Chlorpyrifos	'221	IN, AC	A 1	01/07/2006	30/06/2016	<u>05/72/EC</u>
Chemical	Chlorpyrifos-methyl	'486	IN, AC	A 1	01/07/2006	30/06/2016	<u>05/72/EC</u>
Chemical	Chlorsulfuron	'391	HB	A 3	01/09/2009	31/08/2019	
Chemical	Cinidon ethyl		HB	С	01/10/2002	30/09/2012	<u>02/64/EC</u>
Chemical	Clodinafop		HB	A 2	01/02/2007	31/01/2017	<u>06/39/EC</u>
Chemical	Clofentezine	'418	AC	A 3	01/01/2009	31/12/2018	<u>2008/69</u>
Chemical	Clomazone	'509	HB	A 3	01/11/2008	01/11/2018	<u>2007/76</u>
Chemical	Clopyralid	'455	HB	A 2	01/01/2007	30/04/2017	<u>06/64/EC</u>
Chemical	Clothianidin		IN	С	01/08/2006	31/07/2016	<u>06/41/EC</u>
Chemical	Copper compounds		FU	A 3	01/11/2009	30/11/2016	SCoFCAH voted 01.2009
Chemical	Copper hydroxide		FU	A 3	01/11/2009	30/11/2016	SCoFCAH voted 01.2009
Chemical	Copper oxychloride		FU	A 3	01/11/2009	30/11/2016	SCoFCAH voted 01.2009
Chemical	Cuprous oxide		FU	A 3	01/11/2009	30/11/2016	SCoFCAH voted 01.2009
Chemical	Cyazofamid		FU	С	01/07/2003	30/06/2013	<u>03/23/EC</u>
Chemical	Cyclanilide		PG	С	01/11/2001	31/10/2011	<u>01/87/EC</u>
Chemical	Cyfluthrin	'385	IN, AC	A 1	01/01/2004	31/12/2013	<u>03/31/EC</u>
Chemical	Cyhalofop-butyl		HB	С	01/10/2002	30/09/2012	<u>02/64/EC</u>
Chemical	Cymoxanil	'419	FU	A 3	01/09/2009	31/08/2019	2008/125
Chemical	Cypermethrin	'332	IN, AC	A 1	01/03/2006	28/02/2016	<u>05/53/EC</u>
Chemical	Cyprodinil	'511	FU	A 2	01/05/2007	30/04/2017	<u>06/64/EC</u>
Chemical	Cyromazine	'420	IN	A 3	01/01/2010	31/08/2019	
Chemical	Daminozide	'330	PG	A 1	01/03/2006	28/02/2016	05/53/EC
Chemical	Deltamethrin	'333	IN	A 1	01/11/2003	31/10/2013	<u>03/5/EC</u>
Chemical	Desmedipham	'477	HB	A 1	01/11/2003	31/10/2013	04/58/EC
Chemical	Dicamba	'85	HB	A 3	01/01/2009	31/12/2018	2008/69
Chemical	Dichlorobenzoic acid methylester		FU, PGR	A 3	01/09/2009	31/08/2019	2008/125
Chemical	Dichlorprop-P	'476	HB	A 2	01/06/2007	31/05/2017	06/74/EC
Chemical	Didecyldimethylammonium chloride	-	FU	A 4			
Chemical	Difenacoum	'514	RO	A 4			
Chemical	Difenoconazole	'687	FU	A 3	01/01/2009	31/12/2018	2008/69
Chemical	Diflubenzuron	'339	IN	A 3	01/01/2009	31/12/2018	2008/69
Chemical	Diflufenican	'462	HB	A 3	01/01/2009	31/12/2018	2008/66
Chemical	Dimethachlor		HB	A 3	01/01/2010	31/08/2019	
Chemical	Dimethenamid ? P		HB	С	01/01/2004	31/12/2013	<u>03/84/EC</u>
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Chemical	Dimethoate	'59	IN, AC	A 2	01/10/2007	30/09/2017	<u>07/25/EC</u>

Chemical	Dimox ystrobin	1	FU	C	01/10/2006	30/09/2016	06/75/EC
Chemical	Dinocap	'98	FU, AC	A 1	01/01/2007	31/12/2009	<u>06/136/EC</u>
Chemical	Diquat (dibromide)	'55	HB	A 1	01/01/2002	31/12/2011	<u>01/21/EC</u>
Chemical	Diuron	'100	HB	A 2	01/10/2008	30/09/2018	<u>08/91/EC</u>
Chemical	Dodemorph	'300	FU	A 3	01/09/2009	31/08/2019	2008/125
Chemical	Epoxiconazole	'609	FU	A 3	01/01/2009	31/12/2018	2008/107
Chemical	Esfenvalerate	'481	IN	A 1	01/08/2001	31/07/2011	00/67/EC
Chemical	Ethephon	'373	PG	A 2	01/08/2007	31/07/2017	06/85/EC
Chemical	Ethofumesate	'233	HB	A 1	01/03/2003	28/02/2013	02/37/EC
Chemical	Ethoprophos	'218	NE, IN	A 2	01/10/2007	30/09/2017	07/52/EC
Chemical	Ethoxysulfuron		HB	С	01/07/2003	30/06/2013	03/23/EC
Chemical	Etofenprox	'471	IN	A 3	01/01/2010	31/12/2019	<u> </u>
Chemical	Etoxazole		IN	С	01/06/2005	31/05/2015	05/34/EC
Chemical	Famoxadone		FU	С	01/10/2002	30/09/2012	<u>02/64/EC</u>
Chemical	Fenamidone		FU	С	01/10/2003	30/09/2013	03/68/EC
Chemical	Fenamiphos (aka phenamiphos)		NE	A 2	01/08/2007	31/07/2017	<u>06/85/EC</u>
Chemical	Fenhexamid	<u> </u>	FU	C	01/06/2001	31/05/2011	01/28/EC
Chemical	Fenoxaprop-P	'484	HB	A 3	01/01/2009	31/12/2018	2008/66
Chemical	Fenpropidin	'520	FU	A 3	01/01/2009	31/12/2018	2008/66
Chemical	Fenpropimorph	'427	FU	A 3	01/01/2009	31/12/2018	2008/107
Chemical	Fenpyroximate		AC	A 3	01/01/2009	31/12/2018	2008/107
Chemical	Fipronil	'581	IN	A 2	01/10/2007	30/09/2017	07/52/EC
Chemical	Flazasulfuron		HB	С	01/06/2004	31/05/2014	04/30/EC
Chemical	Florasulam		HB	C	01/10/2002	30/09/2012	02/64/EC
Chemical	Fluazinam	'521	FU	A 3	01/01/2009	31/12/2018	2008/108
Chemical	Fludioxonil	'522	FU	A 3	01/11/2008	01/11/2018	2007/76
Chemical	Flufenacet (formerly fluthiamide)		HB	C	01/01/2004	31/12/2013	<u>03/84/EC</u>
Chemical	Flumioxazin	-	HB	C	01/01/2003	31/12/2012	02/81/EC
Chemical	Fluoxastrobin	-	FU	C	01/08/2008	31/07/2018	08/44/EC
Chemical	Flupyrsulfuron methyl		HB	C	01/07/2001	30/06/2011	01/49/EC
Chemical	Fluroxypyr	'431	HB	A 1	01/12/2000	30/11/2010	<u>00/10/EC</u>
Chemical	Flurtamone		HB	C	01/01/2004	31/12/2013	03/84/EC
Chemical	Flusilazole	'435	FU	A 1	01/01/2007	30/06/2008	06/133/EC
Chemical	Flutolanil	'524	FU	A 3	01/01/2009	31/12/2018	2008/108
Chemical	Folpet	'75	FU	A 2	01/10/2007	30/09/2017	07/5/EC
Chemical	Foramsulfuron		HB	С	01/07/2003	30/06/2013	03/23/EC
Chemical	Forchlorfenuron		PG	C	01/04/2006	31/03/2016	06/10/EC
Chemical	Formetanate		IN, AC	A 2	01/10/2007	30/09/2017	07/5/EC
Chemical	Fosetyl	'384	FU	A 2	01/05/2007	30/04/2017	<u>06/64/EC</u>
Chemical	Fosthiazate	504	NE	C	01/01/2004	31/12/2013	03/84/EC
Chemical	Fuberidazole	'525	FU	A 3	01/01/2004	31/12/2013	2008/108
Chemical	Glufosinate	'437	HB	A 2	01/01/2007	30/09/2017	07/25/EC
Chemical	Glyphosate (incl trimesium aka sulfosate)	'284	HB	A 1	01/07/2002	30/06/2012	01/99/EC
Chemical	Imazalil (aka enilconazole)	'335	FU	A 1	01/01/1999	31/12/2008	97/73/EC
Chemical	Imazamox		HB	С	01/07/2003	30/06/2013	<u>03/23/EC</u>
Chemical	Imazaquin	'699	PG	A 3	01/01/2009	31/12/2018	2008/69
Chemical	Imazosulfuron		HB	С	01/04/2005	31/03/2015	<u>05/3/EC</u>
Chemical	Imidacloprid		IN	A 3	01/08/2009	31/07/2019	2008/116
Chemical	Indoxacarb	<u> </u>	IN	C	01/04/2006	31/03/2016	06/10/EC
Chemical	Iodosulfuron-methyl-sodium	<u> </u>	HB	C	01/01/2004	31/12/2013	03/84/EC
Chemical	Ioxynil	'86	HB	A 1	01/03/2005	28/02/2015	04/58/EC
Chemical	Iprodione	'278	FU	A 1	01/01/2004	31/12/2013	03/31/EC
	Iprovalicarb		FU		01/07/2002	30/06/2011	

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Chemical	Iron sulphate		HB	A 4	01/09/2009	31/08/2019	2008/127
Chemical	Isoproturon	'336	HB	A 1	01/01/2003	31/12/2012	02/18/EC
Chemical	Isoxaflutole	550	HB	C	01/01/2003	31/12/2012	03/68/EC
Chemical	Kresoxim-methyl		FU	C	01/02/1999	31/01/2009	<u>99/01/EC</u>
Chemical	lambda-Cyhalothrin	'463	IN	A 1	01/01/2002	31/12/2011	00/80/EC
Chemical	Lenacil	'163	HB	A 3	01/01/2002	31/12/2011	2008/69
Chemical	Linuron	'76	HB	A 1	01/01/2003	31/12/2013	03/31/EC
Chemical	Lufenuron	70	IN	A I A 3	01/01/2004	31/12/2013	03/31/EC
Chemical	Magnesium phosphide	'228	IN IN, RO	A 3	01/01/2010	31/12/2019	2008/125
Chemical	Maleic hydrazide	'310	PG	A 3 A 1	01/09/2009	31/08/2019	2008/123 03/31/EC
Chemical	Mancozeb	'34	FU	A 1 A 1	01/01/2004	30/06/2016	05/72/EC
Chemical	Maneb	'61	FU		01/07/2006	30/06/2016	05/72/EC 05/72/EC
		'2		A 1			05/72/EC 05/57/EC
Chemical	MCPA		HB	A 1	01/05/2006	30/04/2016	
Chemical	MCPB	'50	HB	A 1	01/05/2006	30/04/2016	05/57/EC
Chemical	Mecoprop	'51	HB	A 1	01/06/2004	31/05/2014	03/70/EC
Chemical	Mecoprop-P	'475	HB	A 1	01/06/2004	31/05/2014	03/70/EC
Chemical	Mepanipyrim		FU	C	01/10/2004	30/09/2014	<u>04/62/EC</u>
Chemical	Mepiquat	'440	PG	A 3	01/01/2009	31/12/2018	2008/108
Chemical	Mesosulfuron		HB	С	01/04/2004	31/03/2014	<u>03/119/EC</u>
Chemical	Mesotrione		HB	С	01/10/2003	30/09/2013	<u>03/68/EC</u>
Chemical	Metalaxyl-M		FU	С	01/10/2002	30/09/2012	<u>02/64/EC</u>
Chemical	Metamitron	'381	HB	A 3	01/09/2009	31/08/2019	2008/125
Chemical	Metazachlor	'411	HB	A 3	01/08/2009	31/07/2019	2008/116
Chemical	Metconazole		FU	A 2	01/06/2007	31/05/2017	06/74/EC
Chemical	Methiocarb (aka mercaptodimethur)	'165	IN, MO, RE	A 2	01/10/2007	30/09/2017	07/5/EC
Chemical	Methoxyfenozide		IN	С	01/04/2005	31/03/2015	05/3/EC
Chemical	Metiram	'478	FU	A 1	01/07/2006	30/06/2016	05/72/EC
Chemical	Metrafenone		FU	С	01/02/2007	31/01/2017	07/6/EC
Chemical	Metribuzin	'283	НВ	A 2	01/10/2007	30/09/2017	07/25/EC
Chemical	Metsulfuron	'441	НВ	A 1	01/07/2001	30/06/2011	00/49/EC
Chemical	Molinate	'235	НВ	A 1	01/08/2004	31/07/2014	03/81/EC
Chemical	Nicosulfuron	'709	НВ	A 3	01/01/2009	31/12/2018	2008/40
Chemical	Oxadiargyl		HB	C	01/07/2003	30/06/2013	03/23/EC
Chemical	Oxadiazon	'213	HB	A 3	01/01/2009	31/12/2018	2008/69
Chemical	Oxamyl	'342	IN, NE	A 2	01/08/2006	31/07/2016	06/16/EC
Chemical	Oxasulfuron	342	HB	C	01/07/2003	30/06/2013	03/23/EC
Chemical	Penconazole	'446	FU	A 3	01/01/2010	31/08/2019	<u>05/25/LC</u>
Chemical	Pendimethalin	'357	HB	A1	01/01/2004	31/12/2013	03/31/EC
Chemical	Pethoxamid	337	HB	C	01/01/2004	31/07/2016	05/31/EC 06/41/EC
Chemical	Phenmedipham	'77	HB	A 1	01/03/2005	28/02/2015	04/58/EC
Chemical	Phosmet	'318	IN	A 1 A 2	01/03/2003	30/09/2017	04/38/EC 07/25/EC
Chemical	Picloram	'174	HB		01/01/2007	30/09/2017 31/12/2018	2008/69
Chemical	Picolinafen	1/4	HB	A 3	01/01/2009	31/12/2018	2008/69 02/64/EC
				C			
Chemical	Picoxystrobin	1001	FU	C	01/01/2004	31/12/2013	<u>03/84/EC</u>
Chemical	Pirimicarb	'231	IN	A 2	01/02/2007	31/01/2017	06/39/EC
Chemical	Pirimiphos-methyl	'239	IN	A 2	01/10/2007	30/09/2017	07/52/EC
Chemical	Prohexadione-calcium	1995	PG	C	01/10/2000	01/10/2010	<u>00/50/EC</u>
Chemical	Propamocarb	'399	FU	A 2	01/10/2007	30/09/2017	07/25/EC
Chemical	Propaquizafop		HB	A 3	01/12/2009	30/11/2019	
Chemical	Propiconazole	'408	FU	A 1	01/06/2004	31/05/2014	03/70/EC
Chemical	Propineb	'177	FU	A 1	01/04/2004	30/03/2014	03/39/EC
Chemical	Propoxycarbazone		HB	С	01/04/2004	31/03/2014	<u>03/119/EC</u>
Chemical	Propyzamide	'315	HB	A 1	01/04/2004	30/03/2014	03/39/EC

Chemical	Prosulfocarb	'539	HB	A 3	01/01/2009	31/12/2018	2007/76
Chemical	Prosulfuron		HB	С	01/07/2002	30/06/2011	02/48/EC
Chemical	Prothioconazole		FU	С	01/08/2008	31/07/2018	08/44/EC
Chemical	Pymetrozine		IN	С	01/11/2001	31/10/2011	01/87/EC
Chemical	Pyraclostrobin		FU, PG	С	01/06/2004	31/05/2014	04/30/EC
Chemical	Pyraflufen-ethyl		НВ	С	01/11/2001	31/10/2011	01/87/EC
Chemical	Pyridate	'447	HB	A 1	01/01/2002	31/12/2011	01/21/EC
Chemical	Pyrimethanil		FU	A 2	01/06/2007	31/05/2017	06/74/EC
Chemical	Pyriproxyfen	'715	IN	A 3	01/01/2009	31/12/2018	2008/69
Chemical	Quinoclamine	'648	HB, AL	A 3	01/01/2009	31/12/2018	2008/66
Chemical	Quinoxyfen		FU	С	01/09/2004	31/08/2014	04/60/EC
Chemical	Quizalofop-P	'641	HB	A 3	01/12/2009	30/11/2019	SCoFCAH
							voted 01.2009
Chemical	Quizalofop-P-ethyl	'641	HB	A 3	01/12/2009	30/11/2019	
Chemical	Quizalofop-P-tefuryl	'641	HB	A 3	01/12/2009	30/11/2019	
Chemical	Rimsulfuron (aka renriduron)		HB	A 2	01/02/2007	31/01/2017	06/39/EC
Chemical	Silthiofam		FU	С	01/01/2004	31/12/2013	<u>03/84/EC</u>
Chemical	S-Metholachlor		HB	С	01/04/2005	31/03/2015	<u>05/3/EC</u>
Chemical	Sodium 5-nitroguaiacolate		PG	A 3	01/11/2009	31/10/2019	2009/11
chemical	Sodium hypochlorite		BA	A 4	01/09/2009	31/08/2019	2008/127
Chemical	Sodium o-nitrophenolate		PG	A 3	01/11/2009	31/10/2019	2009/11
Chemical	Sodium p-nitrophenolate		PG	A 3	01/11/2009	31/10/2019	2009/11
Chemical	Spiroxamine		FU	С	01/09/1999	01/09/2009	<u>99/73/EC</u>
Chemical	Sulcotrione		HB	A 3	01/09/2009	31/08/2019	2008/125
Chemical	Sulfosulfuron		HB	С	01/07/2002	30/06/2011	<u>02/48/EC</u>
Chemical	Sulphur	'0018	FU, AC, RE	A 4			SCoFCAH voted 03.2009
Chemical	Tebuconazole	'494	FU	A 3	01/09/2009	31/08/2019	2008/125
Chemical	Tebufenpyrad		AC	A 3	01/11/2009	31/10/2019	2009/11
Chemical	Teflubenzuron	'450	IN	A 3	01/12/2009	30/11/2019	
Chemical	Tepraloxydim		HB	С	01/06/2005	31/05/2015	<u>05/34/EC</u>
Chemical	Thiabendazole	'323	FU	A 1	01/01/2002	31/12/2011	01/21/EC
Chemical	Thiacloprid		IN	С	01/01/2005	31/12/2014	<u>04/99/EC</u>
Chemical	Thiamethoxam		IN	С	01/02/2007	31/01/2017	<u>07/6/EC</u>
Chemical	Thifensulfuron-methyl	'452	HB	A 1	01/07/2002	30/06/2012	01/99/EC
Chemical	Thiophanate-methyl	'262	FU	A 1	01/03/2006	28/02/2016	05/53/EC
Chemical	Thiram	'24	FU	A 1	01/08/2004	31/07/2014	03/81/EC
Chemical	Tolclofos-methyl	'479	FU	A 2	01/02/2007	31/01/2017	06/39/EC
Chemical	Tolylfluanid	'275	FU, AC	A 2	01/10/2006	30/09/2016	06/06/EC
Chemical	Tralkoxydim	'544	HB	A 3	01/01/2009	31/12/2018	2008/107
Chemical	Triadimenol	'398	FU	A 3	01/09/2009	31/08/2019	2008/125
Chemical	Tri-allate	'97	HB	A 3	01/01/2010	31/12/2019	
Chemical	Triasulfuron	'480	HB	A 1	01/08/2001	31/07/2011	00/66/EC
Chemical	Tribasic copper sulfate		FU	A 3			
Chemical	Tribenuron (aka metometuron)	'546	HB	A 2	01/03/2006	28/02/2016	05/54/EC
Chemical	Triclopyr	'376	HB	A 2	01/06/2007	31/05/2017	06/74/EC
Chemical	Trifloxystrobin		FU	С	01/10/2003	30/09/2013	<u>03/68/EC</u>
Chemical	Triflusulfuron		HB	A 3	01/01/2010	31/12/2019	
Chemical	Trinexapac (aka cimetacarb ethyl)		PG	A 2	01/05/2007	30/04/2017	<u>06/64/EC</u>
Chemical	Triticonazole	'652	FU	A 2	01/02/2007	31/01/2017	06/39/EC
Chemical	Tritosulfuron		НВ	С	01/12/2008	30/11/2018	08/70/EC
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Chemical	zeta-Cypermethrin		IN	A 3	01/12/2009	30/11/2019	
Chemical	Ziram	'31	FU, RE	A 1	01/08/2004	31/07/2014	<u>03/81/EC</u>
Chemical	Zoxamide		FU	С	01/04/2004	31/03/2014	<u>03/119/EC</u>
Chemical repellent	Denathonium benzoate		RE	A 4	01/09/2009	31/08/2019	2008/127
Chemical repellent	Repellents by smell/ Tall oil crude (CAS 8002-26-4)			A 4	01/09/2009	31/08/2019	2008/127
Chemical repellent	Repellents by smell/Tall oil pitch (CAS 8016-81-7)			A 4	01/09/2009	31/08/2019	2008/127
Microbial	Ampelomyces quisqualis strain AQ10		FU	С	01/04/2005	31/03/2015	<u>05/2/EC</u>
Microbial	Bacillus subtilis str. QST 713		BA, FU	С	01/02/2007	31/01/2017	<u>07/6/EC</u>
Microbial	Bacillus thuringiensis subsp. aizawai (ABTS-1857 and GC-91)		[IN]	A 4	01/01/2009	31/12/2018	2008/113
Microbial	Bacillus thuringiensis subsp. israelensis (AM65-52)		[IN]	A 4	01/01/2009	31/12/2018	2008/113
Microbial	Bacillus thuringiensis subsp. kurstaki (ABTS 351, PB 54, SA 11, SA12 and EG 2348)		[IN]	A 4	01/01/2009	31/12/2018	<u>2008/113</u>
Microbial	Bacillus thuringiensis subsp. tenebrionis (NB 176)		[IN]	A 4	01/01/2009	31/12/2018	2008/113
Microbial	<i>Beauveria bassiana</i> (ATCC 74040 and GHA)		IN	A 4	01/01/2009	31/12/2018	<u>2008/113</u>
Microbial	Coniothyrium minitans		FU	С	01/01/2004	31/12/2013	<u>03/79/EC</u>
Microbial	<i>Cydia pomonella</i> granulosis virus (CpGV)		IN	A 4	01/01/2009	31/12/2018	2008/113
Microbial	<i>Gliocladium catenulatum</i> strain J1446		FU	С	01/04/2005	31/03/2015	<u>05/2/EC</u>
Microbial	Lecanicillimum muscarium (Ve6) (former Verticillium lecanii)		IN	A 4	01/01/2009	31/12/2018	2008/113
Microbial	Metarhizium anisopliae (BIPESCO 5F/52)		IN	A 4	01/01/2009	31/12/2018	2008/113
Microbial	Paecilomyces fumosoroseus Apopka strain 97		FU	С	01/07/2001	30/06/2011	<u>01/47/EC</u>
Microbial	Paecilomyces lilacinus		FU	С	01/08/2008	31/07/2018	<u>2008/44/EC</u>
Microbial	Phlebiopsis gigantea (several strains)		FU	A 4	01/01/2009	31/12/2018	2008/113
Microbial	Pseudomonas chlororaphis strain MA342		FU	С	01/10/2004	30/09/2014	<u>04/71/EC</u>
Microbial	Pythium oligandrum (M1)		FU	A 4	01/01/2009	31/12/2018	2008/113
Microbial	<i>Spodoptera exigua</i> nuclear polyhedrosis virus		FU	C	01/12/2007	30/11/2017	<u>07/50/EC</u>
Microbial	Streptomyces K61 (K61) (formerly Streptomyces griseoviridis)		FU	A 4	01/01/2009	31/12/2018	2008/113
Microbial	<i>Trichoderma aspellerum</i> (ICC012) (T11) (TV1) (formerly <i>T. harzianum</i> )		FU	A 4	01/01/2009	31/12/2018	2008/113
Microbial	<i>Trichoderma atroviride</i> (IMI 206040) (T 11) (formerly <i>Trichoderma harzianum</i> )		FU	A 4	01/01/2009	31/12/2018	2008/113
Microbial	<i>Trichoderma gamsii</i> (formerly <i>T. viride</i> ) (ICC080)		FU	A 4	01/01/2009	31/12/2018	<u>2008/113</u>
Microbial	<i>Trichoderma harzianum</i> Rifai (T- 22) (ITEM 908)		FU	A 4	01/01/2009	31/12/2018	2008/113
Microbial	Trichoderma polysporum (IMI 206039)		FU	A 4	01/01/2009	31/12/2018	<u>2008/113</u>
Microbial	Verticillium albo-atrum (WCS850) (formerly V. dahliae)		FU	A 4	01/01/2009	31/12/2018	<u>2008/113</u>

Natural other	Abamectin (aka avermectin)	'495	AC, IN	A 3	01/01/2009	31/12/2018	2008/107
Natural other	Acetic acid	493	HB	A 3	01/01/2009	31/12/2018	2008/107
Natural other	Aluminium silicate (aka kaolin)		RE	A4 A4	01/09/2009	31/08/2018	2008/127
Natural other	Blood meal		RE	A4 A4	01/09/2009	31/08/2019	2008/127
Natural other	Carbon dioxide		IN, RO	A4 A4	01/09/2009	31/08/2019	2008/127
Natural other	Fat distilation residues		RE	A4 A4	01/09/2009	31/08/2019	2008/127
Natural other	Ferric phosphate		MO	C A 4	01/09/2009	31/08/2019	<u>2008/127</u> 01/87/EC
Natural other	Kieselguhr (diatomaceous earth)		IN	A 4	01/09/2009	31/10/2011 31/08/2019	2008/127
Natural other	Milbemectin		IN IN, AC	A 4 C	01/09/2009	30/11/2015	<u>2008/127</u> 05/58/EC
Natural other	Quartz sand		IN, AC RE	A 4	01/09/2009	31/08/2019	<u>05/38/EC</u> 2008/127
Natural other	Spinosad		IN	A 4 C	01/09/2009	31/08/2019	<u>2008/127</u> 07/6/EC
Natural other	Benzoic acid			C	01/02/2007	31/01/2017 31/05/2014	
by synthesis			BA, FU, OT	C			<u>04/30/EC</u>
Natural other by synthesis	Potassium hydrogen carbonate		FU	A 4	01/09/2009	31/08/2019	2008/127
Natural other by synthesis	Urea		IN	A4	01/09/2009	31/08/2019	2008/127
Natural other fatty acid	Capric acid (CAS 334-48-5)		IN, AC, HB, PG	A 4	01/09/2009	31/08/2019	2008/127
Natural other fatty acid	Caprylic acid (CAS 124-07-2)		IN, AC, HB, PG	A 4	01/09/2009	31/08/2019	2008/127
Natural other fatty acid	Fatty acids C7 to C20		IN, AC, HB, PG	A 4	01/09/2009	31/08/2019	2008/127
Natural other fatty acid	Fatty acids C7-C18 and C18 unsaturated potassium salts (CAS 67701-09-1)		IN, AC, HB, PG	A 4	01/09/2009	31/08/2019	2008/127
Natural other fatty acid	Fatty acids C8-C10 methyl esters (CAS 85566-26-3)		IN, AC, HB, PG	A4	01/09/2009	31/08/2019	2008/127
Natural other fatty acid	Lauric acid (CAS 143-07-7)		IN, AC, HB, PG	A 4	01/09/2009	31/08/2019	2008/127
Natural other fatty acid	Methyl decanoate (CAS 110-42-9)		IN, AC, HB, PG	A 4	01/09/2009	31/08/2019	2008/127
Natural other fatty acid	Methyl octaonate (CAS 111-11-5)		IN, AC, HB, PG	A 4	01/09/2009	31/08/2019	2008/127
Natural other fatty acid	Oleic acid (CAS 112-80-1)		IN, AC, HB, PG	A 4	01/09/2009	31/08/2019	2008/127
Natural other fatty acid	Pelargonic acid (CAS 112-05-0)		IN, AC, HB, PG	A 4	01/09/2009	31/08/2019	2008/127
Natural other repellent	Calcium carbonate		RE	A 4	01/09/2009	31/08/2019	2008/127
Natural other repellent	Limestone		RE	A 4	01/09/2009	31/08/2019	2008/127
Natural other Repellent	Methyl nonyl ketone	V	RE	A 4	01/09/2009	31/08/2019	2008/127
Natural other repellent	Sodium aluminium silicate		RE	A 4	01/09/2009	31/08/2019	2008/127
Natural other repellent	Repellents by smell/Fish oil		RE	A 4	01/09/2009	31/08/2019	2008/127
Natural other repellent	Repellents by smell/Sheep fat		RE	A 4	01/09/2009	31/08/2019	2008/127

Semio	(Z)-13-Hexadecen-11yn-1-yl						2008/127
	acetate	$\checkmark$	AT	A 4	01/09/2009	31/08/2019	
Semio	(Z,Z,Z,Z)-7,13,16,19-						<u>2008/127</u>
	Docosatetraen-1-yl isobutyrate	$\checkmark$	AT	A 4	01/09/2009	31/08/2019	
Semio	Ammonium acetate	$\checkmark$	AT	A 4	01/01/2009	31/12/2018	2008/127
Semio	Hydrolysed proteins	$\checkmark$	IN	A 4	01/09/2009	31/08/2019	2008/127
Semio	Putrescine (1,4-Diaminobutane)	$\checkmark$	AT	A 4	01/09/2009	31/08/2019	2008/127
Semio	Trimethylamine hydrochloride	$\checkmark$	AT	A 4	01/09/2009	31/08/2019	2008/127
Semio	Straight Chain Lepidoptera Pheromones	$\checkmark$	AT	A 4	01/09/2009	31/08/2019	<u>2008/127</u>
Semio/SCLP	(2E, 13Z)-Octadecadien-1-yl acetate	V	AT	A 4	01/09/2009	31/08/2019	2008/127
Semio/SCLP	(7E, 9E)-Dodecadien 1-yl acetate	$\checkmark$	AT	A 4	01/09/2009	31/08/2019	2008/127
Semio/SCLP	(7E, 9Z)-Dodecadien 1-yl acetate	$\sqrt{\sqrt{1}}$	AT	A 4	01/09/2009	31/08/2019	2008/127
Semio/SCLP	(7Z, 11E)-Hexadecadien-1-yl acetate	$\sqrt{}$	AT	A 4	01/09/2009	31/08/2019	2008/127
Semio/SCLP	(7Z, 11Z)-Hexadecdien-1-yl acetate	$\checkmark$	AT	A 4	01/09/2009	31/08/2019	2008/127
Semio/SCLP	(9Z, 12E)-Tetradecadien-1-yl acetate	√	AT	A 4	01/09/2009	31/08/2019	2008/127
Semio/SCLP	(E)-11-Tetradecen-1-yl acetate	$\checkmark$	AT	A 4	01/09/2009	31/08/2019	2008/127
Semio/SCLP	(E)-5-Decen-1-ol	$\sqrt{}$	AT	A 4	01/09/2009	31/08/2019	2008/127
Semio/SCLP	(E)-5-Decen-1-yl-acetate	$\sqrt{}$	AT	A 4	01/09/2009	31/08/2019	2008/127
Semio/SCLP	(E)-8-Dodecen-1-yl acetate	$\sqrt{}$	AT	A 4	01/09/2009	31/08/2019	2008/127
Semio/SCLP	(E,E)-8,10-Dodecadien-1-ol	$\checkmark$	AT	A 4	01/09/2009	31/08/2019	2008/127
Semio/SCLP	(E/Z)-8-Dodecen-1-yl acetate	$\sqrt{}$	AT	A 4	01/09/2009	31/08/2019	2008/127
Semio/SCLP	(Z)-11-Hexadecen-1-ol	$\checkmark$	AT	A 4	01/09/2009	31/08/2019	2008/127
Semio/SCLP	(Z)-11-Hexadecen-1-yl acetate	$\sqrt{}$	AT	A 4	01/09/2009	31/08/2019	2008/127
Semio/SCLP	(Z)-11-Hexadecenal	$\sqrt{\sqrt{\sqrt{1}}}$	AT	A 4	01/09/2009	31/08/2019	2008/127
Semio/SCLP	(Z)-11-Tetradecen-1-yl acetate	$\checkmark$	AT	A 4	01/09/2009	31/08/2019	2008/127
Semio/SCLP	(Z)-13-Octadecenal	$\sqrt{}$	AT	A 4	01/09/2009	31/08/2019	<u>2008/127</u>
Semio/SCLP	(Z)-7-Tetradecenal	$\checkmark$	AT	A 4	01/09/2009	31/08/2019	<u>2008/127</u>
Semio/SCLP	(Z)-8-Dodecen-1-ol	$\sqrt{}$	AT	A 4	01/09/2009	31/08/2019	2008/127
Semio/SCLP	(Z)-8-Dodecen-1-yl acetate	$\sqrt{\sqrt{2}}$	AT	A 4	01/09/2009	31/08/2019	2008/127
Semio/SCLP	(Z)-9-Dodecen-1-yl acetate	422 √ √	AT	A 4	01/09/2009	31/08/2019	2008/127
Semio/SCLP	(Z)-9-Hexadecenal	$\sqrt{}$	AT	A 4	01/09/2009	31/08/2019	2008/127
Semio/SCLP	(Z)-9-Tetradecen-1-yl acetate	$\checkmark$	AT	A 4	01/09/2009	31/08/2019	2008/127
Semio/SCLP	Dodecyl acetate	$\sqrt{}$	AT	A 4	01/09/2009	31/08/2019	2008/127
Semio/SCLP	Tetradecan-1-ol	$\checkmark$	AT	A 4	01/09/2009	31/08/2019	2008/127
	Official Total Included:	334			A: Existing active substances divided into four lists for phased evaluations		
					C: New active		

# Appendix 11. Invertebrate beneficials available as biological control agents against invertebrate pests in five European countries.

Beneficial	Taxonomy	Target	Сгор
Adalia bipunctata	coleoptera	aphids on leaves	orchards: all
Adalia bipunctata	coleoptera	aphids on leaves	vegetable greenhouse: all crops
Delphastus pusillus	coleoptera	whiteflies	vegetables greenhouse and covered
Harmonia axyridis	coleoptera	aphids on leaves	Vegetables, orchards
Aphidoletes aphidimyza	diptera	aphids on leaves	vegetable greenhouse: tomato, cucumber, egg plant, sweet pepper
Feltiella acarisuga	diptera	Tetranychus urticae	vegetable greenhouse: all crops
Anthocoris nemoralis	heteroptera	psylla	orchard: pear
Macrolophus caliginosus	heteroptera	Aleurodina (whiteflies), secondarily against Tetranychus & aphids ( <i>Macrosiphum euphorbiae</i> , <i>Aphis gossypii</i> )	vegetable greenhouse: all crops
Anagrus atomus	hymenoptera	Tomato Leaf-hopper ( <i>Hauptidia maroccana</i> )	vegetables
Aphelinus abdominalis	hymenoptera	aphids: <i>M. euphorbiae</i>	vegetable greenhouse: all, tomato, egg plant, sweet pepper
Aphidius colemani	hymenoptera	aphids: A. gossypii, Myzus persicae (green peach aphid)	vegetable greenhouse: all crops
Aphidius ervi	hymenoptera	aphids: Aulacorthum solani, M. euphorbiae, M. persicae	vegetable greenhouse: all crops
Diaeretiella rapae	hymenoptera	aphids : Brevicoryne brassicae	Cabbage, oil-seed rape
Dacnusa sibirica	hymenoptera	Agromyzidae (leaf-miner flies)	vegetable greenhouse: all crops
Diglyphus isaea	hymenoptera	Agromyzidae (leaf-miner flies)	vegetable greenhouse: all crops
Encarsia formosa	hymenoptera	Aleurodina (whiteflies)	vegetable greenhouse: all crops
Eretmocerus eremicus (syn. Californicus)	hymenoptera	Aleurodina (whiteflies): Bemisia tabaci, Trialeurodes vaporariorum	vegetable greenhouse: all crops
Eretmocerus mundus	hymenoptera	Aleurodina (whiteflies): B. tabaci, T. vaporariorum	vegetable greenhouse: all crops, orchards
Orius insidiosus	hymenoptera	thrips: Frankliniella occidentalis, Thrips tabaci	vegetable greenhouse: all crops
Orius laevigatus	hymenoptera	thrips, partial: Tetranychus	vegetable greenhouse: all crops
Orius majusculus	hymenoptera	thrips, partial: Tetranychus	vegetable greenhouse: all crops

### **11.1.** Invertebrate biocontrol agents used in France

Trichogramma brassicae Bezdenko	hymenoptera	Ostrinia nubilalis	maize
Trichogramma evanescens	hymenoptera	(European corn borer) Noctuidae(Owlet moths), Pyralidae	vegetable greenhouse: all crops
Amblyseius andersoni	mite	N. rubi, P. ulmi, T.urticae	orchards: all
Amblyseius andersoni	mite	Aculops lycopersici, Tetranychus	Vegetables
Amblyseius californicus	mite	Tetranychus, Panonychus	vegetables
Amblyseius cucumeris	mite	Tetranychus, thrips	vegetable covered: tomato, cucumber, sweet pepper, all crops (thrips)
Amblyseius degenerans	mite	thrips	vegetable greenhouse: egg plant, sweet pepper
Amblyseius swirskii	mite	thrips, whiteflies	vegetables
Hypoaspis aculeifer	mite	Sciaridae (fungus gnats), bulb	vegetable greenhouse: all
11ypouspis ucuieijei	mite	mite (Rhyzogliphus robini)	crops
Hypoaspis miles	mite	Sciaridae (fungus gnats), thrips	vegetable greenhouse: all crops
Phytoseiulus persimilis	mite	Tetranychus urticae	vegetables
Heterorhabditis bacteriophora,	nematode	Otiorhynchus salicicola, Otiorhynchus sulcatus (vine weevil), Phyllopertha horticola	orchards: crops & nursery
Heterorhabditis megidis	nematode	Otiorhynchus salicicola, Otiorhynchus sulcatus (vine weevil)	Vegetables, flowers
Phasmarhabditis hermaphrodita	nematode	Limacidae (Slugs)	vegetable: general
Steinernema carpocapsae	nematode	Codling moth	pome fruit
Steinernema kraussei	nematode	Otiorhynchus salicicola, Otiorhynchus sulcatus (vine weevil)	vegetable: general
Steinernema carpocapsae	nematode	Noctuids, Gryllotalpa gryllotalpa (European mole cricket), Tipula paludosa (March crane fly)	vegetable: general
Steinernema feltiae	nematode	codling moth, Cydia molesta	pome fruit
Steinernema feltiae	nematode	Sciaridae (fungus gnats)	vegetable: general young plants
Chrysoperla carnea	nevroptera	aphids on leaves	vegetables and ornements covered
Chrysoperla lucasina	nevroptera	aphids, thrips, scales, whiteflies, acarids eggs, leak moth	vegetables and ornements covered
Micromus angulatus	nevroptera	scales, aphids	vegetables
Franklinothrips vespiformis	thysanoptera	thrips	vegetable greenhouse: all crops

## 11.2. Invertebrate biocontrol agents used in Germany

Beneficial	Pest	
Entomopathogenic nematodes		
Heterorhabditis bacteriophora Poinar	Larvae of vine weevil ( <i>Otiorynchus sulcatus</i> ), caterpillar of ghost moths (genus: <i>Hepialus</i> ), larvae of garden chaffer ( <i>Phyllopertha horticola</i> ) and other insect larvae feeding on roots	
Heterorhabditis megidis Poinar	Larvae of vine weevil ( <i>Otiorynchus sulcatus</i> ) and other insect larvae feeding on roots	
Steinernema carpocapsae Weiser	Larvae of vine weevil ( <i>Otiorynchus sulcatus</i> ) and other insect larvae feeding on roots, mole cricket	
Steinernema feltiae Filipjev	Larvae of fungus gnats (Diptera: Sciaridae) and March flies (Diptera: Bibionidae)	
Steinernema kraussei Steiner	Insect larvae feeding on roots, e.g. vine weevil ( <i>Otiorynchus sulcatus</i> )	
Gastropod pathogenic nematodes		
Phasmarhabditis hermaphrodita A. Schneider	Slugs (Deroceras spp, Agriolimax spp and others)	
Predatory mites	· · · · · · · · · · · · · ·	
Amblyseius andersoni Chant	Spider mites ( <i>Tetranychus spp</i> , <i>Panonychus spp</i> ), gall mites (Eriophyidae) u.a.	
Amblyseius barkeri Hughes	Thrips (Frankliniella occidentalis and others)	
Amblyseius californicus McGregor	Spider mites	
Amblyseius cucumeris Oudemans	Thrips (Frankliniella occidentalis and others)	
Amblyseius degenerans Berlese	Thrips	
Amblyseius swirskii Athias-Henriot	-Henriot White flies (e.g. <i>Bemisia tabaci</i> ), spider mites (Tetranychus spp.) and thrips	
Cheyletus eruditus Schrank	Stored product mites, booklice (Psocoptera)	
Hypoaspis aculeifer Canestrini	Thrips	
Hypoaspis milesBerlese	Thrips	
Phytoseiulus persimilis Athias-Henriot	Spider mites (Tetranychus spp)	
Typhlodromus pyri Scheuten	Spider mites	
Predatory thrips (Thysanoptera)		
Franklinothrips vespiformis Crawford	Thrips (in particular <i>Echinothrips americanus</i> , <i>Parthenothrips dracaenae</i> , <i>Frankliniella occidentalis</i> )	
Parasitic wasps (Hymenoptera)		
Anagrus atomus Linnaeus	Cicadidae eggs	
Anagyrus fusciventris Girault	Wooly aphids (Eriosomatidae) and mealy bugs (Pseudococcididae)	
Anisopteromalus calandrae Howard	Drugstore ( <i>Stegobium paniceum</i> ), Tobacco ( <i>Lasioderma serricorne</i> )	
Aphelinus abdominalis Dalman	Aphids (Macrosiphum euphorbiae, Aulacorthum solani)	
Aphelinus mali Haldeman	Wooly aphid (Eriosoma lanigerum)	
Aphidius colemani Viereck	Aphids (Aphis gossypii, Myzus persicae, M. nicotianae)	
Aphidius ervi Haliday	Aphids (Macrosiphum euphorbiae)	
Aphidius matricariae Haliday	Aphids (Myzus persicae)	
Aprostocetus hagenowii Ratzeburg	Cockroaches (Blatta orientalis, Periplaneta spp)	
Cephalonomia tarsalis Ashmead	Saw-toothed and marchand grain beetles ( <i>Oryzaephilus surinamensis</i> und <i>O. mercator</i> )	
Coccidoxenoides perminutus Girault	Different wooly aphids and mealy bugs (Pseudococcididae)	
Coccophagus licymnia Walker	Scale insects (Coccidae)	
Coccophagus rusti Compere	Scale insects (Coccidae)	

Coccophagus scutellaris Dalman	Scale insects (Coccidae)	
Dacnusa sibirica Telenga	Leaf-miner flies (Agromyzidae: <i>Liriomyza</i> and others	
Diglyphus isaea Walker	Leaf-miner flies (Agromyzidae: <i>Liriomyza</i> and others	
Diaeretiella rapae M'Intosh	Cabbage aphid ( <i>Brevicoryne brassicae</i> )	
Encarsia citrina Craw	Diaspididae	
Encarsia formosa Gahan	White fly ( <i>Trialeurodes vaporarium</i> )	
Encyrtus lecaniorum Mayr	Scale insect ( <i>Saissetia hemisphaerica</i> )	
<i>Eretmocerus californicus</i> Howard (= <i>eremicus</i>	Scale filseet (Suisseita hemisphäerica)	
Rose & Zolnerowich)	White flies (Bemisia spp and others)	
Eretmocerus mundus Mercet	White flies (Aleyrodidae)	
Gyranusoidea litura Prinsloo	Long-tailed mealy bug (Pseudococcus longispinus)	
Habrobracon hebetor Say	Stored product moths (Indian Meal moth, <i>Plodia interpuntella</i> ) and ( <i>Ephestia spp</i> )	
Lariophagus distinguendus Förster	Grain weevils ( <i>Sitophilus spp</i> ), drugstore beetle ( <i>Stegobium paniceum</i> ), tobacco beetle ( <i>Lasioderma</i> <i>serricorne</i> ), shiny spider beetle ( <i>Gibbium psylloides</i> ), golden spider beetle ( <i>Niptus hololeucus</i> )	
Leptomastidea abnormis Girault	Wooly aphids and mealy bugs (Pseudococcididae)	
Leptomastix dactylopii Howard	Wooly aphids and mealy bugs (Pseudococcididae)	
<i>Leptomastix epona</i> Walker	Wooly aphids and mealy bugs (Pseudococcididae)	
Lysiphlebus testaceipes Cresson	Apids (Aphis gossypii)	
	Scale insects (Coccidae: Saissetia oleae, Coccus	
Metaphycus flavus Howard	hesperidum)	
	Scale insects (Coccidae: Saissetia oleae,	
Metaphycus helvolus Compere	Coccus hesperidum)	
Metaphycus lounsburyi Howard	Scale insects (Coccidae: Saissetia oleae)	
Metaphycus stanleyi Compere	Scale insects (Coccidae)	
Microterys flavus Howard	Scale insects (Coccidae: Saissetia oleae)	
Pseudaphycus maculipennis Mercet	Wooly aphids and mealy bugs (Pseudococcididae)	
Theocolax elegans Westwood	Lesser grain borer ( <i>Rhyzopertha dominica</i> )	
Theocolax elegans westwood	Thrips (Hercinothrips femoralis, Heliothrips	
Thripobius semiluteus Boucek	haemorrhoidalis, Echinothrips americanus)	
<i>Trichogramma brassicae</i> Bezdenko	Eggs of corn borer ( <i>Ostrinia nubilalis</i> ) and other moths	
Dezdeliko	Eggs of plum maggot moth ( <i>Cydia funebrana</i> ) and	
Trichogramma cacoeciae Marchal	codling moth (Cydia pomonella)	
Trichogramma dendrolimi Matsumura	Eggs of plum maggot moth ( <i>Cydia funebrana</i> ) and	
-	codling moth ( <i>Cydia pomonella</i> )	
Trichogramma evanescens Westwood	Eggs of pest lepidoptera and stored product moths	
Trichogramma evanescens Westwood (Stamm	Eggs of stored product moths	
"Lager") Venturia canescens Gravenhorst	Eggs of stored product mothes (Indian meal moth, <i>Plodia</i>	
	<i>interpuntella</i> ) and ( <i>Ephestia spp</i> )	
Predatory midges and syrphids (Diptera)		
Aphidoletes aphidimyza Rondani	Aphids	
Diaeretiella rapae DeGeer	Aphids	
Feltiella acarisuga Vallot	Spider mites ( <i>Tetranychus urticae</i> , <i>T. cinnabarinus</i> , <i>Panonychus ulmi</i> )	
Predatory beetles (Coleoptera)		
(colcoptera)		
Adalia bipunctata Linnaeus	Aphids	
	Aphids Parasites of fly pupa	

Coccinella septempunctata Linnaeus	Aphids
Cryptolaemus montrouzieri Mulsant	Wooly aphids (Eriosomatidae) and mealy bugs
Cryptotaemas montroaziert Wallsand	(Pseudococcididae)
Exochomus quadripustulatus Linnaeus	Scale insects
Rhyzobius forestieri Mulsant	Scale insects (Saissetia oleae)
Rhyzobius lophantae Blaisdell	Scale insects (Coccidae), Wooly aphids (Eriosomatidae)
Knyzoolus lophunide Blaisden	and mealy bugs (Pseudococcididae)
Rodolia cardinalis Mulsant	Wooly aphids (Eriosomatidae) and mealy bugs
Kodolid curainalis Mulsant	(Pseudococcididae)
Stethorus punctillum Weise	Spider mites
Predatory true bugs (Heteroptera)	
Anthocoris nemoralis Fabricius	Suckers (Psyllids, Psyllidae)
Dicyphus hesperus Knight	White fly (Trialeurodes vaporariorum)
Macrolophus melanotoma Costa(= caliginosus	White flies (Aleyrodidae)
E. Wagner)	white mes (Aleyfoundae)
Macrolophus pygmaeus Rambur	White flies (Aleyrodidae)
Orius insidiosus Say	Thrips (Thysanoptera)
Orius laevigatus Fieber	Thrips (Thysanoptera)
Orius majusculus Reuter	Thrips (Thysanoptera)
Lacewings	
Chrysoperla carnea Stephens	Aphids
Parasites and predators of stable flies	
Diaeretiella rapae Girault & Sanders	Housefly-related flies
Muscidifurax zaraptor Kogan & Legner	Housefly-related flies
Nasonia vitripennis Walker	Housefly-related flies
Spalangia cameroni Perkins	Housefly-related flies
Spalangia endius Walker	Housefly-related flies
Spalangia nigroaeneus Curtis	Housefly-related flies
Hydrothaea aenescens Wiedemann	Housefly-related flies

Source: http://www.jki.bund.de/

## 11.3. Invertebrate biocontrol agents used in Spain

Beneficial	Taxonomy	Target	Сгор
Adalia bipunctata	coleoptera	Aphids on leaves	orchards: all
Adalia bipunctata	coleoptera	Aphids on leaves	vegetable greenhouse: all crops
Delphastus pusillus	coleoptera	Whiteflies	vegetables greenhouse and covered
Harmonia axyridis	coleoptera	aphids on leaves	Vegetables, orchards
Aphidoletes aphidimyza	diptera	Aphids on leaves	vegetable greenhouse: tomato, cucumber, egg plant, sweet pepper
Feltiella acarisuga	diptera	Tetranychus urticae	vegetable greenhouse: all crops
Anthocoris nemoralis	heteroptera	Psylla	orchard: pear
Macrolophus caliginosus	heteroptera	Aleurodina (whiteflies), secondary vs Tetranychus & Aphids: <i>Macrosiphum</i> <i>euphorbiae, Aphis gossypii</i>	vegetable greenhouse: all crops
Anagrus atomus	hymenoptera	Tomato Leaf-hopper (Hauptidia maroccana)	vegetables
Aphelinus abdominalis	hymenoptera	Aphids: <i>Macrosiphum</i> <i>euphorbiae</i>	vegetable greenhouse: all, tomato, egg plant, sweet pepper
Aphidius colemani	hymenoptera	Aphids: <i>Aphis gossypii,</i> <i>Myzus persicae</i> (green peach aphid)	vegetable greenhouse: all crops
Aphidius ervi	hymenoptera	Aphids: Aulacorthum solani Macrosiphum euphorbiae, myzus persicae	vegetable greenhouse: all crops
Diaeretiella rapae	hymenoptera	Aphids : Brevicoryne brassicae	Cabbage, oil-seed rape
Dacnusa sibirica	hymenoptera	Agromyzidae (leaf-miner flies)	vegetable greenhouse: all crops
Diglyphus isaea	hymenoptera	Agromyzidae (leaf-miner flies)	vegetable greenhouse: all crops
Encarsia formosa	hymenoptera	Aleurodina (whiteflies)	vegetable greenhouse: all crops
Eretmocerus eremicus (syn. Californicus)	hymenoptera	Aleurodina (whiteflies): Bemisia tabaci, Trialeurodes vaporariorum	vegetable greenhouse: all crops
Eretmocerus mundus	hymenoptera	Aleurodina (whiteflies): Bemisia tabaci, Trialeurodes vaporariorum	vegetable greenhouse: all crops, orchards
Orius insidiosus	hymenoptera	Thrips: Frankliniella occidentalis, Thrips tabaci	vegetable greenhouse: all crops
Orius laevigatus	hymenoptera	Thrips, partial: Tetranychus	vegetable greenhouse: all crops
Orius majusculus	hymenoptera	Thrips, partial: Tetranychus	vegetable greenhouse: all crops

Trichogramma brassicae	hymenoptera	Ostrinia nubilalis	maize
Bezdenko Trichogramma brassicae		(European corn borer) Noctuidae(Owlet moths),	vegetable greenhouse: all
Bezdenko	hymenoptera	Pyralidae	crops
Trichogramma	humanantana	Noctuidae(Owlet moths),	vegetable greenhouse: all
evanescens	hymenoptera	Pyralidae	crops
Amblyseius andersoni	mite	N. rubi, P. ulmi, T.urticae	orchards: all
Amblyseius andersoni	mite	Aculops lycopersici,	Vegetables
Amblyseius californicus	mite	TetranychusTetranychus, Panonychus	vegetables
Amblyseius cucumeris	mite	Tetranychus, thrips	vegetable covered: tomato, cucumber, sweet pepper, all crops (thrips)
Amblyseius degenerans	mite	Thrips	vegetable greenhouse: egg plant, sweet pepper
Amblyseius barkeri (mackenziei)	mite	Thrips	vegetables
Amblyseius swirskii	mite	Thrips, whiteflies	vegetables
Hypoaspis aculeifer	mite	Sciaridae (fungus gnats), bulb mite (Rhyzogliphus robini)	vegetable greenhouse: all crops
Hypoaspis miles	mite	Sciaridae (fungus gnats), thrips	vegetable greenhouse: all crops
Phytoseiulus persimilis	mite	Tetranychus urticae	vegetables
Heterorhabditis bacteriophora,	nematode	Otiorhynchus salicicola, Otiorhynchus sulcatus (vine weevil),Phyllopertha horticola	orchards: crops & nursery
Heterorhabditis megidis	nematode	Otiorhynchus salicicola, Otiorhynchus sulcatus (vine weevil)	Vegetables, flowers
Phasmarhabditis hermaphrodita	nematode	Limacidae (Slugs)	vegetable: general
Steinernema carpocapsae	nematode	Codling moth	Apple, pear
Steinernema kraussei	nematode	Otiorhynchus salicicola, Otiorhynchus sulcatus (vine weevil)	vegetable: general
Steinernema carpocapsae	nematode	Noctuids, Gryllotalpa gryllotalpa (European mole cricket), Tipula paludosa (March crane fly )	vegetable: general
Steinernema feltiae	nematode	Codling moth, cydia molesta	Apple, pear
Steinernema feltiae	nematode	Sciaridae (fungus gnats)	vegetable: general young plants
Chrysoperla carnea	nevroptera	aphids on leaves	vegetables and ornements covered
Chrysoperla lucasina	nevroptera	Aphids, thrips, scales, whiteflies, acarids eggs, leak moth	vegetables and ornements covered
Micromus angulatus	nevroptera	Scales, aphids	vegetables
Franklinothrips vespiformis	Thrips	Thrips	vegetable greenhouse: all crops
Adalia bipunctata	coleoptera	Aphids on leaves	orchards: all
Adalia bipunctata	coleoptera	Aphids on leaves	vegetable greenhouse: all

			crops
Delphastus pusillus	coleoptera	Whiteflies	vegetables greenhouse and covered
Harmonia axyridis	coleoptera	aphids on leaves	Vegetables, orchards
Aphidoletes aphidimyza	diptera	Aphids on leaves	vegetable greenhouse: tomato, cucumber, egg plant, sweet pepper
Feltiella acarisuga	diptera	Tetranychus urticae	vegetable greenhouse: all crops
Anthocoris nemoralis	heteroptera	Psylla	orchard: pear
Macrolophus caliginosus	heteroptera	Aleurodina (whiteflies), secondary vs Tetranychus & Aphids: <i>Macrosiphum</i> <i>euphorbiae</i> , <i>Aphis gossypii</i>	vegetable greenhouse: all crops
Anagrus atomus	hymenoptera	Tomato Leaf-hopper (Hauptidia maroccana)	vegetables
Aphelinus abdominalis	hymenoptera	Aphids: Macrosiphum euphorbiae	vegetable greenhouse: all, tomato, egg plant, sweet pepper

## 11.4. Invertebrate biocontrol agents used in Switzerland

Beneficial	Taxonomy	Target	Сгор
Adalia bipunctata	coleoptera	Aphids on leaves	orchards: all
Adalia bipunctata	coleoptera	Aphids on leaves	vegetable greenhouse: egg
			plant, cucumber, sweet
			pepper
Aphidoletes aphidimyza	diptera	Aphids on leaves	vegetable greenhouse:
			tomato,
			cucumber, egg plant,
			sweet pepper
Aphidoletes aphidimyza	diptera	Aphids on leaves	vegetable covered: all
Feltiella acarisuga	diptera	Tetranychus urticae	vegetable greenhouse:
			cucumber, egg plant,
			sweet pepper
Anthocoris nemoralis	heteroptera	Psylla	orchard: pear
Macrolophus	heteroptera	Aleurodina (whiteflies),	vegetable greenhouse:
caliginosus		secondary vs	tomato,
		Tetranychus &	egg plant, sweet pepper
		Aphids: Macrosiphum	
		euphorbiae, Aphis gossypii	
Aphelinus abdominalis	hymenoptera	Aphids:	vegetable greenhouse: all,
		Macrosiphum	tomato,
		euphorbiae,	egg plant, sweet pepper
		Aulacorthum solani	
		<i>Myzus persicae</i> (green peach	
		aphid)	
Aphidius colemani	hymenoptera	Aphids:	vegetable greenhouse: all
		Aphis gossypii,	crops
		Aphis fabae,	
		<i>Myzus persicae</i> (green peach	
	humanantara	aphid) Aphids:	vegetable greenhouse: all
	hymenoptera	Aulacorthum solani	crops
Aphidius ervi		Macrosiphum euphorbiae	crops
Inplicatus ervi	hymenoptera	Agromyzidae	vegetable greenhouse: all
Dacnusa sibirica	nymenoptera	(leaf-miner flies)	crops
Duchusu sibiricu	hymenoptera	Agromyzidae	vegetable greenhouse: all
Diglyphus isaea	nymenopteru	(leaf-miner flies)	crops
Encarsia formosa	hymenoptera	Aleurodina (whiteflies)	vegetable greenhouse: all
Litearstaformosa	inginenopteru	Theatoania (whitemes)	crops
	hymenoptera	Aleurodina (whiteflies):	vegetable greenhouse:
	,	Bemisia tabaci,	tomato,
Eretmocerus eremicus		Trialeurodes vaporariorum	cucumber, egg plant,
(syn. Californicus)		1	sweet pepper
Orius insidiosus	hymenoptera	Thrips:	vegetable greenhouse:
		Frankliniella occidentalis,	sweet pepper
		Thrips tabaci	* * *
	hymenoptera	Thrips,	vegetable greenhouse: all
Orius laevigatus		partial: Tetranychus	crops
	hymenoptera	Thrips,	vegetable greenhouse:
Orius majusculus	_	partial: Tetranychus	sweet pepper
Trichogramma brassicae	hymenoptera	Ostrinia nubilalis	maize
Bezdenko		(European corn borer)	
Trichogramma brassicae	hymenoptera	Noctuidae(Owlet moths),	vegetable greenhouse: all

Bezdenko		Pyralidae	crops
Amblyseius barkeri (mackenziei)	mite	Thrips	vegetable greenhouse: all crops, tomato, cucumber,
Amblyseius californicus	mite	Tetranychus	egg plant, sweet pepper vegetable greenhouse: sweet pepper
Amblyseius cucumeris	mite	Tetranychus, thrips	vegetable covered: tomato, cucumber, sweet pepper, all crops (thrips)
Amblyseius degenerans	mite	Tetranychus, thrips	vegetable greenhouse: egg plant, sweet pepper
Hypoaspis aculeifer	mite	Sciaridae (fungus gnats)	vegetable greenhouse: all crops
Hypoaspis miles	mite	Sciaridae (fungus gnats)	vegetable greenhouse: all crops
Phytoseiulus persimilis	mite	Tetranychus urticae	vegetable greenhouse: all crops, tomato, cucumber, egg plant, sweet pepper
Heterorhabditis bacteriophora,	nematode	Otiorhynchus salicicola, Otiorhynchus sulcatus (vine weevil)	orchards: nursery
Heterorhabditis megidis	nematode	Otiorhynchus salicicola, Otiorhynchus sulcatus (vine weevil)	orchards: nursery
Heterorhabditis megidis	nematode	Otiorhynchus salicicola, Otiorhynchus sulcatus (vine weevil)	vine: young plants
Phasmarhabditis hermaphrodita	nematode	Limacidae (Slugs)	vegetable: general
Photorhabdus luminescens	nematode	Otiorhynchus salicicola, Otiorhynchus sulcatus (vine weevil)	orchards: nursery
Photorhabdus luminescens	nematode	Otiorhynchus salicicola, Otiorhynchus sulcatus (vine weevil)	vine: young plants
Steinernema carpocapsae	nematode	Otiorhynchus salicicola, Otiorhynchus sulcatus (vine weevil)	orchards: all
Steinernema carpocapsae	nematode	Otiorhynchus salicicola, Otiorhynchus sulcatus (vine weevil)	vine: young plants
Steinernema carpocapsae	nematode	Noctuids, Gryllotalpa gryllotalpa (European mole cricket)	vegetable: general
Steinernema feltidae	nematode	Sciaridae (fungus gnats)	vegetable: general young plants
Xenorhabdus bovienii	nematode	Sciaridae (fungus gnats)	vegetable: general young plants

Steinernema feltiae Steinernema kraussei	Nemasys	Entompathogenic	Sciarids, leafminer,
	Tternasys		
Steinernema kraussei	· ·	nematode	WFT
Stelliernenna kraussei	Nemasys L	Entompathogenic	vine weevil
		nematode	vine weevin
Heterorhabditis megidis	Nemasys H	Entompathogenic	vine weevil,
neterornabattis megiais		nematode	vine weevil,
Heterorhabditis megidis	Nemasys H	Entompathogenic	Grubs
neterornabattis megiais	Ttellidsys II	nematode	
Steinernema carpocasae	Nemasys C	Entompathogenic	codling moth (occasional
Siemernema carpocasae	Ttellidsys C	nematode	cutworms)
Steinernema carpocasae	Nemasys C	Entompathogenic	Hylobius
_	Themasys C	nematode	Trytoblus
Phasmarhabditis	Nemaslug	slug parasitic nematode	Slugs
hermaphrodita	Iveniasiug	slug parasitie hematode	Slugs
Adalia bipunctata	Adalsure	Natural enemy	Aphids
Amblyseius californicus	Ambsure	Natural enemy	Thrips, rsm
Trichogramma evanescans	Trichogramma	Natural enemy	Caterpillars
Anagrus atomus	Anagsure	Natural enemy	Leaf hopper
Amblyseius cucumeris	Ambsure	Natural enemy	Thrips
TT · · · 1	TT	Ni-town 1 and another	Thrips, bulb mite,
Hypoaspis miles	Hyposure	Natural enemy	sciarids
Orius laevigatus	Orisure	Natural enemy	Thrips
Aphelinus abdominalis	Aphelsure	Natural enemy	Aphids
Aphidius ervi	Aphissure (e)	Natural enemy	Thrips
Aphidius colemani	Aphisure (c)	Natural enemy	Thrips
Aphidoletes aphidimyza	Aphidosure	Natural enemy	Aphids
Chilocorus nigritus	Chilosure(n)	Natural enemy	Scale insect
Chrysoperla carnea	Chrysosure (c)	Natural enemy	Aphids
Cryptolaemus montrouzieiri	Cryptosure (m)	Natural enemy	Mealy bug
Dacnusa sibirica	Dacsure (si)	Natural enemy	Leaf miner
Diglyphus isaea	Digsure (i)	Natural enemy	Leaf miner
Encarsia formosa	Encsure	Natural enemy	Whitefly
Encarsia formosa and		•	
Eretmocerus eremicus	Enersure	Natural enemy	Whitefly
Eretmocerus eremicus	Eretsure (f)	Natural enemy	Whitefly
Feltiella acarisuga	Felsure (a)	Natural enemy	rsm
Macrolophus caliginosus	Macsure (c)	Natural enemy	Whitefly
Phytoseiulus persimilis	Phytosure (p)	Natural enemy	rsm
Bombus terrestris	Beesure	Pollinator	Pollination
Metaphycus helvolus,			
Encarsia citrina,			
Coccophagus lycimnia and	Scalesure	Natural enemy	Scale insect
Encyrtus infelix			
Leptomastix dactilopii	Leptosure (d)	Natural enemy	Mealy bug
Leptomastix dactylopii,			
Anagyrus pseudococci and	Mealysure	Natural enemy	Mealy bug
Leptomastidea abnormis	licarjouro		1.1001, 00g
Metaphycus helviolus	Metasure (h)	Natural enemy	Scale insect
Encarsia formosa	EN-STRIP	parasitic wasp	Whitefly
Encarsia formosa +		· · ·	•
Linearsia jorniosa 1	ENERMIX	parasitic wasp	Whitefly

## 11.5. Invertebrate biocontrol agents used in the United Kingdom

Eretmocerus eremicus	ERCAL	parasitic wasp	Whitefly
Macrolophus caliginosus	MIRICAL	predatory bug	Whitefly/spidermite
Macrolophus caliginosus	MIRICAL NYMPH	predatory bug	Whitefly/spidermite
Feltiella acarisuga	SPIDEND	predatory bug	Spidermite
Phytoseiulus persimilis	SPIDEX	predatory bug	Spidermite
Amblyseius californicus	SPICAL	predatory mite	Spidermite
Ambiysetus catijornicus Aphidoletes aphidimyza	APHIDEND	predatory bug	Aphids
Aphelinus abdominalis	APHILIN	· · ·	Aphids
Aphelinus abaominalis Aphidius colemani	APHILIN	parasitic wasp	Aphids
1		parasitic wasp	1
Chrysoperla carnea	CHRYSOPA	predatory bug	Aphids
Aphidius ervi	ERVIPAR	parasitic wasp	Aphids
Episyrphus balteatus	SYRPHIDEND	predatory bug	Aphids
Adalia bipunctata	ADALIA larvae	predatory beetle	Aphids
Amblyseius cucumeris	THRIPEX	predatory mite	Thrips + some mites
Orius laevigatus	THRIPOR	predatory bug	Thrips
Amblyseius swirski	SWIRSKI MITE	predatory mite	Thrips and Whiteflies
Dacnusa sibirica +	DIMINEX	parasitic wasp	Leafminers
Diglyphus isaea			
Diglyphus isaea	MIGLYPHUS	parasitic wasp	Leafminers
Dacnusa sibirica	MINUSA	parasitic wasp	Leafminers
Hypoaspis aculeifer	ENTOMITE aculeifer	predatory mite	Sciarids
Steinernema feltiae	ENTONEM	parasitic nematode	Sciarids
Steinernema feltiae	SCIA-RID	parasitic nematode	Mushroom flies
Steinernema carpocapsae	CAPSANEM 50 million	parasitic nematode	Cranefly, Caterpillar
Trichogramma sp.	TRICHO-STRIP	parasitic wasp	Caterpillar
Heterorhabditis megidis	LARVANEM	parasitic nematode	Vine Weevil, Chafer
Cryptolaemus montrouzieri	CRYPTOBUG	predatory beetle	Mealybug
Adalia bipunctata	Adaline b	Predator	Aphids
Amblyseius (Euseius) ovalis	Ovaline	Predator	Whitefly and thrips
Amblyseius (Iphiseius) degenerans	Amblyline d	Predator	Thrips
Amblyseius (Neoseiulus ) californicus	Amblyline cal	Predator	Spider mites
Amblyseius (Neoseiulus) cucumeris	Amblyline cu	Predator	Thrips
Amblyseius (Typhlodromips)	Amblyline m	Predator	Thrips
montdorensis Amblyseius (Typhlodromips)	Swirskiline	Predator	Whitefly and thrips
swirskii			
Amblyseius andersoni	Anderline aa	Predator	Spider mites
Anagrus atomus	Anagline a	Parasitoid	Leaf Hoppers
Anthocoris nemoralis	Antholine n	Predator	Pear Psylla
Aphelinus abdominalis	Apheline a	Parasitoid	Aphids
Aphidius colemani	Aphiline c	Parasitoid	Small aphids
Aphidius ervi	Aphiline e	Parasitoid	Large aphids
Aphidoletes aphidimyza	Aphidoline a	Predator	Aphids
Atheta coriaria	Staphyline c	Predator	Sciarid and Shore Flies
Bombus terrestris	Beeline Total System	Pollinator	-
Chrysoperla carnea	Chrysoline c	Predator	Aphids
Chi ysoperia cumea			
Cryptolaemus montrouzieri	Cryptoline m	Predator	Mealybugs

Diglyphus isaea	Digline i	Parasitoid	Leafminers
Encarsia formosa	Encarline f	Parasitoid	Trialeurodes
Eretmocerus eremicus	Eretline e	Parasitoid	<i>Trialeurodes</i> and <i>Bemisia</i>
Feltiella acarisuga	Feltiline a	Predator	Spider mites
Heterorhabditis megidis	Nemasys H	Entomopathogenic nematode	Vine Weevils
Hypoaspis miles	Hypoline m	Predator	Sciarid Flies
Macrolophus caliginosus (also known as M. pygmaeus)	Macroline c	Predator	Whiteflies
Orius laevigatus	Oriline 1	Predator	Thrips
Orius majusculus	Oriline m	Predator	Thrips
Phasmarhabditis hermaphrodita	Nemaslug	Entomopathogenic nematode	Slugs
Phytoseiulus persimilis	Phytoline p	Predator	Spider mites

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