DOI: 10.1007/s13592-017-0506-6

# Toxicity of thiametoxam on in vitro reared honey bee brood

Giacomo Grillone, Daniela Laurino, Aulo Manino, Marco Porporato

Dipartimento di Scienze Agrarie, Forestali e Alimentari, Università di Torino, Largo Paolo Braccini 2, Grugliasco, 10095, Turin, Italy

Received 22 November 2016 - Revised 22 February 2017 - Accepted 14 March 2017

**Abstract** – Pesticides are a possible cause of pollinator decline and honey bee colony losses experienced in several countries in recent years. In the past years, many north-west Italian beekeepers reported the presence of dead brood in field apiaries during neonicotinoid-coated maize sowing; therefore, a possible role of these insecticides was suspected. The objective of this study was to test this hypothesis. Laboratory repeated dose toxicity tests on in vitro reared larvae were carried out using thiamethoxam. The repeated treatment median lethal concentration ( $LC_{50}$ ) and the median lethal dose ( $LD_{50}$ ) 14 and 19 days after grafting were calculated and resulted of the same order of magnitude of realistic brood exposure under a worst-case scenario. Various sublethal effects, like brownish larvae, duplication of the pupal integument, delay in development, and deformed adults were also observed.

Apis mellifera / artificial rearing / cumulative toxicity / neonicotinoids / sublethal doses / thiametoxam

## 1. INTRODUCTION

An extensive pollinator decline (Gallai et al., 2009; Potts et al., 2010; Nieto et al., 2014) and heavy honey bee colony losses (Janke and Rosenkranz, 2009; Neumann and Carreck, 2010) have been experienced in several countries lately. In the effort to bring light on the origin of these phenomena, several factors have so far been investigated. Furthermore, new features like the colony collapse disorder (CCD) syndrome have been described (vanEngelsdorp et al., 2009), nevertheless not a single cause was definitively identified (Maini et al., 2010; Ratnieks and Carreck, 2010; Williams et al., 2010). Therefore, it is most likely that some multifactorial effect is involved, and lately, the cumulative action of several stressors on the honey bee's immune response has been proposed as a unifying model (Nazzi et al., 2012; Di Prisco et al., 2013).

Corresponding author: D. Laurino, daniela.laurino@unito.it

Manuscript Editor: Monique Gauthier

Published online: 14 April 2017

Among the various factors suggested, pesticides are one of the most relevant and in-depth investigated causes of colony losses (Blacquière et al., 2012; Krupke et al., 2012; Lu et al., 2012; van der Sluijs et al., 2013; European Academies Science Advisory Council, 2015). In particular, neonicotinoid insecticides—especially when used in seed coating—were greatly debated since heavy honey bee mortalities were observed in Italy and other European countries during maize sowing (Greatti et al., 2003 and 2006; Forster, 2009; Pistorius et al., 2009; Chauzat et al., 2010; van der Geest, 2012). At last, these facts led in 2008 to the ban of the use of neonicotinoids and fipronil as coating active ingredients for maize and sunflower. As a result, a dramatic reduction in the poisoning incidents reported in Italy was observed (APENET, 2009). In the years before the ban, many north-west Italian beekeepers reported dead brood with symptoms reminding the European foul brood (EFB) in their hives during maize sowing, but an EFB diagnosis was not confirmed. After the neonicotinoid and fipronil use had been restricted, this syndrome nearly disappeared; therefore, a possible role of these insecticides in causing such symptoms was suspected, and the objective of this study is to test this hypothesis.

Pesticide risk assessment on honey bee brood is far more complex than on adult workers, especially when chronic or sublethal effects are involved, since the honey bee brood develops in the hives and carefully reared inside comb cells, and exposure to pesticides is mostly mediated by both forager and nurse adult workers. Due to such difficulties, several laboratory (Czoppelt 1991; Davis et al., 1988; Mussen et al., 2004; Aupinel et al., 2007a, b: Gregore and Ellis, 2011: Hendriksma et al., 2011; Tavares et al., 2015), cage (Schur et al., 2003), and field procedures (Oomen et al., 1992; Everich et al., 2009) have been proposed for testing pesticide toxicity on honey bee brood and to assess experimental results. Presently, a universal consensus on a specific methodology for testing pesticides on honey bee brood has not yet been achieved (Aupinel et al., 2009; European Food Safety Authority, 2013a, b). Nonetheless, as a first step to detect any possible relationship between neonicotinoids and honey bee brood anomalies, we planned some laboratory toxicity tests on in vitro reared larvae hoping that the results could contribute to the debate. A repeated dose exposure test modified from Aupinel's test (Aupinel et al., 2007a, b) was developed, and thiamethoxam was used as toxic agent, since it is extensively used on crops and its toxicity to honey bees is well known even though the few available data on larvae are controversial (European Food Safety Authority, 2013a).

## 2. MATERIALS AND METHODS

## 2.1. Brood rearing

First instar worker larvae were collected from three *A. m. ligustica* colonies. Computer-assisted wing venation morphometric analysis (Meixner et al., 2013) was used to assess the honey bee subspecies. The hives were periodically checked to exclude the presence of the most common honey bee diseases (Shimanuki and Knox, 2000).

The larvae were grafted from a single brood comb into the wells of a sterile 48-well tissue culture plate; different plates were used for each tested colony. After grafting, the brood was reared in a dark incubator at 34.5 ( $\pm 0.5$ ) °C for 21 days to allow adult emergence. Larvae were kept at 96% relative humidity in hermetic desiccators containing dishes filled with a  $K_2SO_4$  saturated solution, and pupae were kept at 80% relative humidity in hermetic desiccators containing dishes filled with a NaCl saturated solution.

The Aupinel's (Aupinel et al., 2005) semiartificial diet was adopted. Larvae were fed daily for 6 days putting a food drop onto the well's bottom; diet composition and amounts fed to larvae are shown in Table I. Fresh royal jelly produced in DISAFA's apiary, yeast extract from Biokar diagnostics (Beauvais, France), and sugars from Serva Feinbiobiochemica (Heiderberg, Germany) were used.

# 2.2. Toxicity tests

A thiamethoxam commercial formulation available in Italy (Actara® 25.0% pure thiamethoxam, hydro-dispersible granules) was used. It was mixed with the larval food so that known amounts of thiamethoxam were ingested by the larvae. Since the larvae were in contact with the artificial food, some contact action could not be completely excluded.

Preliminary tests were carried out to determine the concentration range between 100% mortality and a mortality level not significantly different from that of the untreated controls. On this basis, the following thiamethoxam concentrations were used to determine the LC<sub>50</sub>: 20, 10, 5, 2, 1, 0.5, and 0.2 mg/L. Three replications—one 48-well tissue plate per replicate—were done for each tested concentration and the untreated control, taking care that larvae from a different colony were used for each replicate.

From day 1 to day 5—day 0 being the day the larvae were grafted—1  $\mu L$  of a thiamethoxam suspension in water was added to the food in each well with the treated larvae, and 1  $\mu L$  of water was added in the wells with the untreated control larvae. The thiamethoxam suspension concentrations were such as to obtain the desired test concentration after dispersion of the given 1  $\mu L$  drop in the larval food. Doses fed to larvae are shown in Table I.

**Table I..** Daily food volumes provided to the larvae, composition of the diet according to their age, and doses of thiamethoxam fed to larvae for each tested concentration from day 1 to day 5 after grafting, day 0 being the grafting day.

Rearing day	Day 0	Day 1	Day 2	Day 3	Day 4	Day 5	Total
Larval food amount (µL)	10	10	20	30	40	50	160
diet composition							
Royal jelly (%)	50	50	50	50	50	50	_
Yeast extract (%)	1	1	1.5	2	2	2	_
D-glucose (%)	6	6	7.5	9	9	9	_
D-fructose (%)	6	6	7.5	9	9	9	_
Dry matter <sup>a</sup> (%)	29.55	29.55	33.05	36.55	36.55	36.55	_
Thiamethoxam dose (ng/larv	ra)						
20 mg/L	_	200	400	600	800	1000	3000
10 mg/L	_	100	200	300	400	500	1500
5 mg/L	_	50	100	150	200	250	750
2 mg/L	_	20	40	60	80	100	300
1 mg/L	_	10	20	30	40	50	150
0.5 mg/L	_	5	10	15	20	25	75
0.2 mg/L	_	2	4	6	8	10	30

<sup>&</sup>lt;sup>a</sup> Calculated from Crane (1990) for royal jelly and manufacturer's specifications for the remaining ingredients

Mortality and any morphological and/or developmental anomaly that could be considered as a sublethal effect were visually checked and recorded daily immediately before feeding. Larvae that died within 24 h after grafting were removed from the experiments and all computations in order to exclude grafting effects.

Two toxicity endpoints were evaluated: (i) cumulative mortality at day 14 after grafting as an assessment of living pupae (individual not molted from prepupa to pupa on day 14 after grafting were considered as dead in the pre-pupal phase); (ii) number of emerged adults (non-emerged individuals on day 21 after grafting were counted as dead during the pupal stage).

# 2.3. Statistical analysis

The larval survival was analyzed with the survival analysis module of the statistical package PAST 3.05 (Hammer et al., 2001). The Kaplan-Meier survival curves for the tested concentration and the untreated control were drawn, and each concentration curve was compared with that of the untreated control by means of the log-rank test.

The Kaplan-Meier curves and the log-rank tests were computed according to Kleinbaum and Klein (2005). The obtained P values were corrected with the Holm-Bonferroni procedure (Holm, 1979) to provide a multicomparison correction.

The median lethal concentration (LC<sub>50</sub>) at day 14 and day 19 after grafting and the relative 95% confidence intervals were calculated by means of probit analysis; the procedure devised by Milani (1995) was adopted. Since in the tests each honey bee larva ingests a total of 150 µL of food from day 1 to day 5 (Table I), the repeated treatment cumulative median lethal dose (LD<sub>50</sub>) was obtained from the relative LC<sub>50</sub>. Moreover, the LC<sub>50</sub> at day 14 and day 19 after grafting were calculated from mortality data relative to each tested colony; subsequently, the three pairwise LC<sub>50</sub> ratios were calculated and their statistical significance determined, under the null hypothesis that they are identical, following the procedure given by Robertson and Preisler (1992) and the computational details adopted by Laurino et al. (2013).

The frequency of individuals showing sublethal effects below LC<sub>50</sub> concentrations were

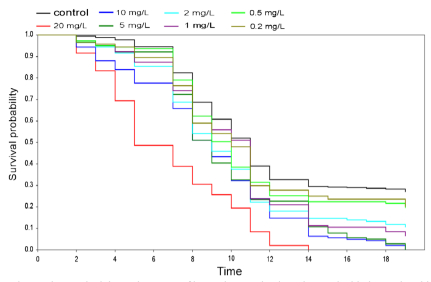


Figure 1. Kaplan-Meier survival dynamic curves of honey bee worker brood treated with decreasing thiamethoxam concentrations and of the untreated controls.

compared with those observed in the untreated controls by means of Pearson's  $\chi^2$  test of independence with d.f. = 1 and significance P values were calculated with a Monte Carlo permutation test with 9999 replicates. The statistical package PAST 3.05 (Hammer et al., 2001) was used for these computations.

### 3. RESULTS

Larval mortality 24 h after grafting was always very low (0–3 dead larvae/plate; average survival  $\pm$  standard deviation = 99.2  $\pm$  1.64%). In the following days, mortality increased

throughout the experiments, although feeding and, therefore, thiamethoxam administration had ceased 5 days after grafting to allow fully grown larvae to transform into prepupae.

Kaplan-Meier survival dynamic curves show a regular increase in mortality from the untreated controls to the highest tested thiamethoxam concentration (Figure 1). All tested concentrations resulted statistically highly different from those of the untreated controls (P < 0.01) except for the lowest one (0.2 mg/L) which was hardly significant (P < 0.05) as a consequence of the Holm-Bonferroni correction (Table II).

**Table II..** Survival dynamics of in vitro reared honey bee worker brood treated with decreasing thiamethoxam concentrations compared with those of the untreated controls.

Thiamethoxam concentration comparisons	Log-rank $\chi^2$	Bonferroni-Holm corrected $\alpha$		P (same) value	
20 mg/L vs control	164.91	0.05	0.01	$9.57 \times 10^{-38}$ **	
10 mg/L vs control	79.45	0.025	0.005	$4.96 \times 10^{-19} **$	
5 mg/L vs control	120.28	0.0166	0.00333	$5.49 \times 10^{-28} **$	
2 mg/L vs control	64.45	0.0125	0.0025	$9.92 \times 10^{-16} **$	
1 mg/L vs control	43.23	0.01	0.002	$4.86 \times 10^{-11} **$	
0.5 mg/L vs control	11.60	0.0083	0.00166	0.00066**	
0.2 mg/L vs control	8.35	0.0072	0.00142	0.0039*	

**Table III..** Thiametoxam repeated treatment median lethal concentration (LC<sub>50</sub>) and median lethal dose (LD<sub>50</sub>) for in vitro reared honey bee worker brood at day 14 and day 19 after grafting. In brackets LC<sub>50</sub> and LD<sub>50</sub> upper and lower limits at P=0.95

	Day 14	Day 19
LC <sub>50</sub> (mg/L)	1.53 (0.33–3.26)	0.79 (0.25–1.39)
LD <sub>50</sub> (ng/larva)	229 (50–489)	118 (37–209)

 $LC_{50}$  14 days after grafting, when pupation was achieved by all surviving individuals, resulted in 1.53 mg/L of larval food, nearly the double of the value of 0.79 mg/L of larval food reached 5 days later after adult emergence (Table III). Food intake

was substantially completed by all larvae reaching the prepupal phase, and therefore, an individual intake of 150  $\mu$ L of food was assumed to compute the LD<sub>50</sub> values shown in Table III even if the larvae that died in the first days did not evidently ingest so much food. The three LD<sub>50</sub> values calculated for each colony used in the tests were substantially similar, and no statistically significant difference emerged from the pairwise LC<sub>50</sub> ratios.

Various anomalies and sublethal effects were observed in the reared brood (Figure 2). In general, sublethal effects were most abundant at lower concentrations than at higher ones, as far as the number of dead larvae and pupae diminished. They were categorised as shown in Table IV.

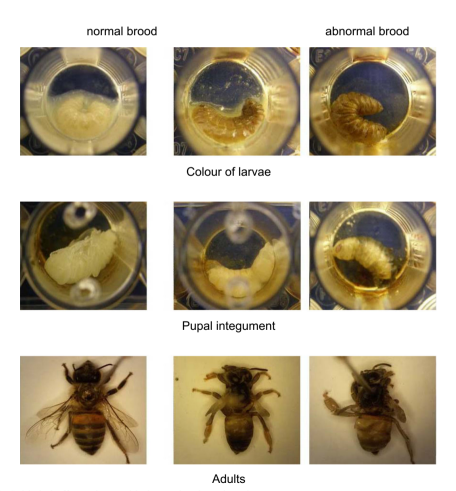


Figure 2. Sublethal effects observed in honey bee brood rearing tests.

**Table IV..** Number of sublethal effects showed by in vitro reared honey bee worker brood treated below  $LC_{50}$  thiamethoxam concentrations (0.5 and 0.2 mg/L) and in the untreated controls. A number of normal individuals are reported for comparison purposes

Sublethal effect	Larvae		$\chi^2$	Monte Carlo P	
	Treated	Control			
Color of larvae					
Brownish	141	114	4.98	0.0281*	
Pearly-white	146	172			
Pupation time					
Delayed	40	27	10.11	0.0017**	
Normal	26	52			
Pupal integument					
Double	3	0	3.42	0.1091 n.s.***	
Normal	66	77			
Adult workers					
Deformed	36	62	8.18	0.0073**	
Normal	21	11			

<sup>\*</sup>Significant (P < 0.01); \*\*highly significant (P < 0.01); \*\*\*n.s. not significant at p = 0.05

The number of brownish worker larvae treated with thiamethoxam concentrations below  $LC_{50}$  was significantly higher than that of untreated controls; pupation time was highly significantly delayed in treated brood, and significantly more deformed adults emerged from it in comparison with untreated controls. Moreover, three pupae still enclosed in the larval integument were also observed in the treated brood.

### 4. DISCUSSION

The very low larval mortality observed 24 h after grafting indicates that grafting effects were negligible throughout the experiment; thanks to the implementation of sound beekeeping practices in the management of the colonies the larvae were taken from and the correct grafting technique.

Feeding the larvae for 5 days with a constant thiamethoxam concentration instead of feeding them with a single dose in an early development stage, as suggested in most recently proposed protocols for testing pesticides on brood (Aupinel et al., 2007a, b; Hendriksma et al., 2011), seems a more realistic simulation of what may happen in a honey bee colony growing in an intensive farming area. The adopted repeated dose

approach could anyhow be considered more as a chronic toxicity test than as an acute toxicity test; therefore, the use of Kaplan-Meier survival dynamic curves to depict the mortality increase, and of log-rank tests to compare them, appear fully justified. Since the untreated control mortality was higher than the optimal levels suggested in the literature (Aupinel et al., 2005 and 2009), the more restrictive value of  $\alpha=0.01$  was preferred to the usual  $\alpha=0.05$  to evidence statistical significance. Under such conditions, the lowest tested concentration could be regarded as the NOAEC for thiamethoxam on brood in the adopted experimental conditions.

LC<sub>50</sub> figures obtained by us are of an order of magnitude lower than those reported by Tavares et al. (2015) for Africanized honey bees. Different responses to toxic compounds have been evidenced between honey bee strains (Ladas, 1972; Mansour and Al-Jalil, 1985; Suchail et al., 2000 Laurino et al., 2013), but in this case, the adopted experimental designs—single dose vs repeated doses—should account for the difference.

The use of LC<sub>50</sub>—and/or the derived LD<sub>50</sub>—values to express the results of a repeated dose toxicity test could be somehow questionable, since they are more often derived from acute

toxicity tests. In any case, they prove to be rather useful when comparing laboratory results with a realistic landscape-level exposure to pesticides. Honey bee larvae are exposed to pesticides chiefly through the food nurse bees give them since the amounts of pollen and honey given to them is very low. Nurse bees may eat contaminated pollen and nectar and thus produce larval food containing pesticides (Rortais et al., 2005). Therefore, effective exposure levels are difficult to determine. To overcome these difficulties, the EFSA guidance document on the risk assessment of plant protection products on bees (European Food Safety Authority, 2013b) proposes various shortcut values (SV) to be used for the estimation of oral exposure via nectar and pollen consumption under different scenarios. Since the reported EFB-like symptoms had been observed in field apiaries during maize sowing, the SVs proposed by EFSA for the adjacent crop (4.4 µg/larva) and/or field margin (2.2 µg/larva) scenarios can be considered as the most appropriate and should be chosen among the various SVs relative to honey bee larvae. These SVs refer to an a.i. application rate of 100 mg/m<sup>2</sup>, while the maximum application rate allowed for maize seed dressing in Italy was 6.3 mg/m<sup>2</sup> only (European Food Safety Authority, 2013a); therefore, the SVs should be reduced to 272 and 136 ng/larva respectively. In any case, it should be noted that such SVs are to be considered as a worst-case scenario. Since the LD<sub>50</sub> values obtained in the present study are lower than the suggested SVs, it is likely that a widespread use of dressed seeds could expose honey bee larvae to dangerous thiamethoxam concentrations. Therefore, extensive brood mortality and colony weakening would likely occur.

Various sublethal effects of pesticides on honey bee brood have been reported in the literature (Desneux et al., 2007; Wu et al., 2011; Blacquière et al., 2012; Simon-Delso et al., 2014), and some of them, like the delay in pupation and in adult emergence, were observed also during this research. On the contrary, developmental troubles leading to the emergence of deformed adults and the extremely rare—and therefore not statistically significant—appearance of pupae that are still enclosed in the larval integument were not

commonly observed and seem to be rather unlikely for neurotoxic insecticides like thiamethoxam.

#### **ACKNOWLEDGEMENTS**

The authors wish to thank Dr. Noa Simon-Delso for her helpful criticism on the earlier draft of the manuscript.

Toxicité du thiamétoxame sur le couvain d'abeille élevé in vitro

Apis mellifera / élevage artificiel / toxicité cumulative / néonicotinoïde / dose sublétale

Toxizität von Thiametoxam für in vitro aufgezogene Brut der Honigbiene

Apis mellifera / künstliche Aufzucht / kumulative Toxizität / Neonikotinoide / sublethale Dosis / Thiametoxam

#### REFERENCES

APENET (2009) Effects of coated maize seed on honey bees. Report based on results obtained from the first year of activity of the APENET project [on line] http://www.cra-api.it/online/immagini/Apenet\_2009\_eng.pdf (accessed on 28 April 2015)

Aupinel, P., Fortini, D., Dufour, H., Tasei, J., Michaud, B., Odoux, F., Pham-Delegue, M. (2005) Improvement of artificial feeding in a standard in vitro method for rearing Apis mellifera larvae. Bull. Insectol. 58, 107–111

Aupinel, P., Fortini, D., Dufour, H., Michaud, B., Marolleau, F., Tasei, J.N., Odoux, J.F. (2007a) Toxicity of dimethoate and fenoxycarb to honey bee brood (*Apis mellifera*), using a new in vitro standardized feeding method. Pest Manag. Sci. 63, 1090–1094

Aupinel, P., Medrzycki, P., Fortini, D., Dufour, H., Michaud, B., Tasei, J.N., Odoux, J.F. (2007b) A new larval in vitro rearing method to test effects of pesticides on honey bee brood. Redia 90, 91–94

Aupinel, P., Fortini, D., Michaud, B., Medrzycki, P., Padovani, E. et al. (2009) Honey bee brood ring-test: method for testing pesticide toxicity on honeybee brood in laboratory conditions. Julius-Kühn-Archiv 423, 96–102

Blacquière, T., Smagghe, G., van Gestel, C.A.M., Mommaerts, V. (2012) Neonicotinoids in bees: a review on concentrations, side-effects and risk assessment. Ecotoxicology 21, 973–992

- Crane, E. (1990) Bees and beekeeping: science, practice and world resources. Heinemann Newnes, Oxford
- Chauzat, M.-P., Martel, A.-C., Blanchard, P., Clément, M.-C., Schurr, F., Lair, C., Ribière, M., Wallner, K., Rosenkranz, P., Faucon J.-P. (2010) A case report of a honey bee colony poisoning incident in France. J. Apic Res. 49, 113–115
- Czoppelt, C. (1991) Evaluation of toxicity of the juvenoid phenoxycarb to honeybee larvae reared *in vitro*. Apidologie 22, 457–459
- Davis, A.R., Solomon, K.R., Shuel, R.W. (1988) Laboratory studies of honeybee larval growth and development as affected by systemic insecticides at adult-sublethal levels. J. Apic. Res. 27, 146–161
- Desneux, N., Decourtye, A., Delpuech J.-M. (2007) The Sublethal Effects of Pesticides on Beneficial arthropods. Annu. Rev. Entomol. 52, 81–106
- Di Prisco, G.D., Cavaliere, V., Annoscia, D., Varricchio, P., Caprio, E., Nazzi, F., Gargiulo, G., Pennacchio, F. (2013) Neonicotinoid clothianidin adversely affects insect immunity and promotes replication of a viral pathogen in honey bees. PNAS 110 (46), 18466–18471
- European Academies Science Advisory Council (2015) Ecosystem services, agriculture and neonicotinoids. German National Academy of Sciences Leopoldina, Halle (Saale)
- European Food Safety Authority (2013a) Conclusion on the peer review of the pesticide risk assessment for bees for the active substance thiamethoxam. EFSA Journal 11(1): 3067
- European Food Safety Authority (2013b) EFSA guidance document on the risk assessment of plant protection products on bees (*Apis mellifera*, *Bombus* spp. and solitary bees). EFSA Journal 11(7): 3295
- Everich, R., Schiller, C., Whitehead, J., Beavers, M., Barrett, K. (2009) Effects of Captan on Apis mellifera Brood Development Under Field Conditions in California Almond Orchards. J. Econ. Entomol. 102, 20–29
- Forster, R., (2009) Bee poisoning caused by insecticidal seed treatment of maize in Germany in 2008. Julius-Kühn-Archiv 423, 126–131
- Gallai, N., Salles, J.M., Settele, J., Vaissière, B.E. (2009) Economic valuation of the vulnerability of world agriculture confronted with pollinator decline. Ecol. Econ. 68, 810–821
- van der Geest B (2012) Bee poisoning incidents in the Pomurje region of Eastern Slovenia in 2011. Julius-Kuhn-Archiv 437, 124
- Greatti, M., Sabatini, A.G., Barbattini, R., Rossi, S., Stravisi, A. (2003) Risk of environmental contamination by the active ingredient imidacloprid used for corn seed dressing. Preliminary results. Bull. Insectol. 56, 69–72
- Greatti, M., Barbattini, R., Stravisi, A., Sabatini, A.G., Rossi, S. (2006) Presence of the a.i. imidacloprid on vegetation near corn fields sown with gaucho® dressed seeds. Bull. Insectol. 59, 99–103
- Gregorc, A., Ellis, J.D. (2011) Cell death localization in situ in laboratory reared honey bee (*Apis mellifera* L.)

- larvae treated with pesticides. Pest. Biochem. Physiol. 99, 200–207
- Hammer, Ø., Harper, D.A.T., Ryan, P.D. (2001) PAST:
   Paleontological Statistics Software Package for
   Education and Data Analysis. Palaeontol. Electron.
   4(1), 9pp
- Hendriksma, H.P., Härtel, S., Steffan-Dewenter, I. (2011) Honey bee risk assessment: new approaches for in vitro larvae rearing and data analyses. Methods Ecol. Evol., 2, 509–517
- Holm, S., (1979) A Simple Sequentially Rejective Multiple Test Procedure. Scand. J. Stat. 6, 65–70
- Janke, M., Rosenkranz, P. (2009) Periodical honey bee colony losses in Germany: preliminary results from a four years monitoring project. Julius-Kühn-Archiv 423, 108–117
- Kleinbaum, D.G., Klein, M. 2005 Survival analysis: a self-learning text. Springer
- Krupke, C.H., Hunt, G.J., Eitzer, B.D., Andino, G., Given, K. (2012) Multiple Routes of Pesticide Exposure for Honey Bees Living Near Agricultural Fields. PLoS ONE 7(1), e29268
- Ladas, A. (1972) Der Einfluss verschiedener Konstitutionsund Umweltfaktoren auf die Anfälligkeit der Honigbiene (Apis mellifica L.) gegenüber zwei insektiziden Pflanzenschutzmitteln. Apidologie, 3, 55–78
- Laurino, D., Manino, A., Patetta, A., Porporato, M. (2013) Toxicity of neonicotinoid insecticides on different honey bee genotypes. Bull. Insectol. 66, 119–126
- Lu, C., Warchol, K.M., Callahan, R.A. (2012) In situ replication of honey bee colony collapse disorder. Bull. Insectol. 65, 99–106
- Maini, S., Medrzycki, P., Porrini, C. (2010) The puzzle of honey bee losses: a brief review. Bull. Insectol. 63, 153–160
- Mansour, S.A., Al-Jalil, M.K. (1985) Pesticides and beneficial organisms: 1. - the response of two honey bee races to certain insecticides. Pesticides 19(3), 39–40
- Meixner, M.D., Pinto, M.A., Bouga, M., Kryger, P., Ivanova, E., Fuchs S. (2013) Standard methods for characterising subspecies and ecotypes of *Apis mellifera*. J. Apic. Res. 52(4), 1–28, DOI: 10.3896 /IBRA.1.52.4.05
- Milani, N. (1995) The resistance of *Varroa jacobsoni* oud to pyrethroids: A laboratory assay. Apidologie 26, 415–429
- Mussen, E.C., Lopez, J.E., Peng, C.Y.S (2004) Effects of selected fungicides on growth and development of larval honey bees, *Apis mellifera* L. (Hymenoptera: Apidae). Environ. Entomol. 33, 1151–1154
- Nazzi, F., Brown, S.P., Annoscia, D., Del Piccolo, F., Di Prisco, G., Varricchio, P., Della Vedova, G., Cattonaro, F., Caprio, E., Pennacchio, F. (2012) Synergistic Parasite-Pathogen Interactions Mediated by Host Immunity Can Drive the Collapse of Honeybee Colonies. PLoS Pathog 8(6), e1002735
- Neumann, P., Carreck, N.L. (2010) Honey bee colony losses. J. Apic. Res. 49, 1–6

- Nieto, A., Roberts, S.P.M., Kemp, J., Rasmont, P., Kuhlmann, M. et al. (2014) European Red List of bees. Publication Office of the European Union, Luxembourg
- Oomen, PA, de Ruijter, A., van der Steen J. (1992) Method for honeybee brood feeding tests with insect growthregulating insecticides. Bull. OEPP/EPPO Bull. 22, 613–616
- Pistorius, J., Bischoff, G., Heimbach, U., Stähler, M. (2009)
  Bee poisoning incidents in Germany in spring 2008
  caused by abrasion of active substance from treated
  seeds during sowing of maize. Julius-Kühn-Archiv
  423, 118–126
- Potts, S.G., Biesmeijer, J.C., Kremen, C., Neumann, P., Schweiger, O., Kunin, W.E. (2010) Global pollinator declines: trends, impacts and drivers. Trends Ecol. Evol. 25, 345–353
- Ratnieks, F.L.W., Carreck, N.L. (2010) Clarity on honey bee collapse? Science 327, 152–153
- Robertson, J.L., Preisler, H.K. (1992) Pesticide bioassays with Arthropods. CRC Press, Boca Raton
- Rortais, A., Arnold, G., Halm, M.P., Touffet-Briens, F. (2005) Modes of honeybees exposure to systemic insecticides: estimated amounts of contaminated pollen and nectar consumed by different categories of bees. Apidologie 36, 71–83
- Schur, A., Tornier, I., Brasse, D., Mühlen, W., von der Ohe, W., Wallner, K., Wehling, M. (2003) Honey bee brood ring-test in 2002: method fort the assessment of side effects of plant protection products on the honey bee brood under semi-field conditions. Bull. Insectol. 56, 91–96

- Shimanuki, H., Knox, D.A. (2000) Diagnosis of Honey Bee Diseases. U.S. Department of Agriculture, Agriculture Handbook No. AH– 690
- Simon-Delso, N., San Martin, G., Bruneau, E., Minsart, L.-A., Mouret, C., Hautier, L. (2014) Honeybee Colony disorder in crop areas: The role of pesticides and viruses. PLoS ONE 9(7), e103073
- van der Sluijs, J.P., Simon-Delso, N., Goulson, D., Maxim, L., Bonmatin, J.-M., Belzunces, L.P. (2013) Neonicotinoids, bee disorders and the sustainability of pollinator services. Curr. Opin. Environ. Sust. 5, 293–305
- Suchail, S., Guez, D., Bbelzunces, L.P. (2000) Characteristics of imidacloprid toxicity in two Apis mellifera subspecies. Environ, Toxicol. Chem. 19, 1901–1905
- Tavares, D.A., Roat, T.C., Carvalho, S.M., Silva-Zacarin, E.C.M., Osmar Malaspina, O. (2015) *In vitro* effects of thiamethoxam on larvae of Africanized honey bee *Apis mellifera* (Hymenoptera: Apidae). Chemosphere 135, 370–378
- vanEngelsdorp, D., Evans, J.D., Saegerman, C., Mullin, C., Haubruge, E. et al. (2009) Colony collapse disorder: a descriptive study. PLoS ONE 4, e6481
- Williams, G.R., Tarpy, D.R., vanEngelsdorp, D., Chauzat, M.-P., Cox-Foster, D.L., Delaplane, K. S., Neumann, P., Pettis, J.S., Rogers, R.E.L., Shutler, D. (2010) Colony Collapse Disorder in context. Bioessays 32, 845–846
- Wu, J.Y., Anelli, C.M., Sheppard, W.S. (2011) Sub-Lethal Effects of Pesticide Residues in Brood Comb on Worker Honey Bee (*Apis mellifera*) Development and Longevity. PLoS ONE 6(2), e14720