Antimicrobial activity of four essential oils against pigmenting *Pseudomonas fluorescens* and biofilm-producing *Staphylococcus aureus* of dairy origin

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Abstract

Essential oils (EOs) are aromatic and volatile mixtures of secondary metabolites of plant origin with many useful properties, among which the antimicrobial activity is also of interest for the food industry. EOs can exert their antimicrobial potential both directly, in food products and active packaging, and indirectly, as sanitizing and anti-biofilm agents of food facility surfaces. Aim of this research was to evaluate the antimicrobial activity of four EOs (bergamot, cinnamon, manuka and thyme) against *Pseudomonas fluorescens* and *Staphylococcus aureus* isolated from milk and dairy products. The chemical composition of EOs was evaluated by Gas Chromatography-Mass Spectrometry analysis. Minimum Inhibitory Concentration values were determined by a microplate method against 9 *Ps. fluorescens* from marketed mozzarella with blue discoloration defect, and 3 biofilm-producing *S. aureus* from milk. Reference ATCC strains were included. Pigment production activity by *Ps. fluorescens* was assessed both in culture and in cheese. EOs of manuka (leptosperm, cinnamon, manuka and thyme) against *Pseudomonas fluorescens* and *Staphylococcus aureus* isolated from milk and dairy products. The present work was aimed to evaluate the antimicrobial activities of four EOs (*Citrus bergamia* Risso, *Cinnamomum zeylanicum* L., *Leptospermum scoparium* J.R. et G. Forst and *Thymus vulgaris* L.) on food-related bacteria: *Pseudomonas fluorescens*, isolated from mozzarella cheese with blue discoloration, and biofilm-producing *Staphylococcus aureus* from milk. Gas chromatography–mass spectrometry (GC-MS) analyses were carried out to perform the chemical characterization of the four tested EOs.

Materials and Methods

**Bacterial cultures**

A total of 10 *Ps. fluorescens* was used in this study: 9 wild isolates (PS59, PS60, PS63, PS70, PS71, PS231, PS249, PS251 and PS282) from commercial samples of spoiled mozzarella cheeses with blue pigmentation and the reference strain *Ps. fluorescens* ATCC 13525. The wild isolates, provided by Istituto Zooprofilattico Sperimentale Lazio-Toscana (Pisa), were phenotypically identified by API 20 NE (bioMérieux, Marcy-l’Etoile, France). Moreover, the identification was confirmed genetically according to Scardellini et al. (2004). After DNA extraction by boiling at 95°C for 10 min, the species-specific PCR was performed using the KAPA Taq ReadyMix PCR kit (Kapa Biosystems, Boston, MA, USA).

The pigmentogenesis of pseudomonads was studied in mozzarella cheese and in a lab-made minimal nutritive broth used for the cultivation and maintenance of *Pseudomonas fluorescens* and other microorganisms (Yeast Extract Glucose Broth, YEGB, https://www.bioworld.com/productinfo/3_43_287_688/10174/Yeast-Extract-Glucose-Broth.html). YEGB had the following composition: glucose 1.5 g/L, di-potassium hydrogen orthophosphate 5.2 g/L, potassium dihydrogen phosphate 3.18 g/L, magnesium sulfate 0.12 g/L, yeast extract 0.5 g/L, ammonium chloride 0.54 g/L. Marketed citric mozzarella samples were inoculated with a final count of about 10^6 CFU/mL in preservation liquid. Samples were stored at 10°C for 72 hours in their packaging and then opened, maintained at the same temperature and observed daily up to 10 days. *Pseudomonas* quantitative determination on Pseudomonas Agar with CFC Supplement (Oxoid, Basingstoke, UK) was carried out to exclude the presence of pseudomonads in the preservation liquid. As for YEGB assay, each isolate was grown overnight at 25°C in YEGB, and ten-fold serial dilutions in sterile saline solution were prepared. Ten µL of the dilutions were seeded in 5 different Eppendorf tubes per isolate, to obtain final counts of pseudomonads ranging from 10^1 to 10^6 CFU/mL. Each tube, of 1.5 mL declared capacity, was filled to the top with 1.8 mL of YEGB to maximize the surface of broth culture exposed to air. Eppendorf tubes were left open, incubated at 25°C and observed daily up to 7 days.

Among biofilm-producing *S. aureus*, 3 strains, isolated from raw milk, came from the collection of Food and Drug Department, Parma (PA1 and PA2) and from that of Veterinary Science.

**Introduction**

Essential oils (EOs) are aromatic and volatile mixtures of secondary metabolites obtained from different parts of plants. EOs are composed of many bioactive molecules, mainly terpenes, terpenoids and phenyl-propanes, and their actual antimicrobial effect depends on many intrinsic and environmental factors (Hyldgaard et al., 2012). Their use in food industry is aimed to provide flavours and to improve safety and quality of products. They can be employed directly in food, or as components of food packaging (Lucera et al., 2012) and even in sanitization of food processing environments and food-contact surfaces due to their antimicrobial and anti-biofilm potential (Valeriano et al., 2012).

The present work was aimed to evaluate the antimicrobial activities of four EOs (*Citrus bergamia* Risso, *Cinnamomum zeylanicum* L., *Leptospermum scoparium* J.R. et G. Forst and *Thymus vulgaris* L.) on food-related bacteria: *Pseudomonas fluorescens*, isolated from mozzarella cheese with blue discoloration, and biofilm-producing *Staphylococcus aureus* from milk.

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**Conflict of interest**

The authors declare no potential conflict of interest.

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Department, Pisa (P11) and their biofilm-producing capacity had been previously assessed following Di Cicco et al. (2015), the other 2 were reference strains, *S. aureus* ATCC 35556 (strong producer) and *S. aureus* ATCC 12600 (intermediate producer). *S. aureus* PA1 and *S. aureus* PA2 had been previously characterized as methicillin-resistant (MRSA) (Di Cicco et al., 2016); *S. aureus* P11 was a methicillin-susceptible, enterotoxin C producer (Pedone et al., 2014).

**Essential oils**

EOs (from Flora s.r.l., Lorenzana, Pisa, Italy) of bergamot (*C. bergamia* Risso, Cb), cinnamon bark (*C. ceylanicum* L., in 40% ethanol, Cz), manuka leaves (*L. scoparium* J. R. et G. Forst, Ls) and thyme (*T. vulgaris* L., Tv) were used.

**Gas chromatography-mass spectrometry analysis**

The chemical composition of EOs was determined by using Gas Chromatography-Electron Impact Mass Spectrometry (GC-MS). Analyses were performed with a Varian CP-3800 gas chromatograph equipped with a DB-5 capillary column (30 m × 0.25 mm; coating thickness 0.25 μm) and a Varian Saturn 2000 ion trap mass detector. Analytical conditions were those reported by Fratini et al. (2014). The identification of the constituents was based on comparison of retention times with those of pure authentic samples, comparing their retention indices relative (LRI) to the series of *n*-hydrocarbons, by computer matching against commercial libraries (NIST 98 and ADAMS) (Adams, 1995) and homemade library mass spectra, built up from pure substances, known oils and MS literature data (Stenhagen et al., 1974).

**Minimal inhibitory concentration determination assay**

MIC values of EOs, i.e. the lowest concentration that inhibits the visible microbial growth, were determined following Wiegand et al. (2008) with minor modifications. The assay was performed in sterile 96-well microtiter plates using 10 μL of bacterial inoculum and 190 μL of each EOs dilution. EOs dilutions were prepared in Tryptone Soy Broth (Oxoid) supplemented with dimethyl sulfoxide (DMSO) to a final ratio of 1:3:4 (v/v) and a two-fold dilution series was performed from 1/8 to 1/16384. For the bacterial *inoculum* an overnight broth culture of each microorganism, spec-trophotometrically (Ultrospec 2100 Pro, Amersham Biosciences, Buckinghamshire, GB) adjusted at 550 nm at about 1.5 x 10⁸ CFU/mL, was used. Positive and negative controls, and for Cz also ethanol control, were included. 

![Image](image-url)

| Table 1. Chemical composition of essential oils detected by gas chromatography-mass spectrometry analysis. |
|-----------------------------------------------|
| Compound                   | Class | LRI | Cb | Cz | Ls | Tv |
|-----------------------------|-------|-----|----|----|----|----|
| α-pinene                    | mh    | 940 | 1.1| 1.7| 1.1|   |
| Sabine                     | mh    | 978 | 1.1|    |    |   |
| β-pinene                    | mh    | 981 | 6  |    |    |   |
| Myrcene                    | mh    | 993 | 1.3|    |    |   |
| α-phellandrene             | mh    | 1006| 1.1|    |    |   |
| α-terpine|                | mh    | 1019| 1.5|    |    |   |
| p-cymene                   | mh    | 1026| 20.2|    |    |   |
| o-cymene                   | mh    | 1026| 2.5|    |    |   |
| Limonene                   | mh    | 1032| 30.8|    |    |   |
| β-phellandrene             | mh    | 1033| 4.4|    |    |   |
| Y-terpine                   | mh    | 1062| 7.4| 8.3|    |   |
| Linalool                   | om    | 1102| 12.5| 5  |    |   |
| Camphor                    | om    | 1148| 1  |    |    |   |
| 4-terpineol                | om    | 1180| 1.8|    |    |   |
| (Z)-cinnamaldehyde         | nt    | 1223| 1.2|    |    |   |
| Linalool acetate           | om    | 1260| 34.9|    |    |   |
| (E)-cinnamaldehyde         | nt    | 1274| 55 |    |    |   |
| Thymol                     | om    | 1290| 15.3|    |    |   |
| Carvacrol                  | om    | 1301| 29.7|    |    |   |
| α-cubebene                 | sh    | 1351| 3.2|    |    |   |
| Eugenol                    | pp    | 1361| 4.6|    |    |   |
| α-copaene                  | sh    | 1376| 1.6| 4.8|    |   |
| α-gurjune                  | sh    | 1410| 1  |    |    |   |
| β-caryophyllene            | sh    | 1418| 10.2| 2.4| 4.8|   |
| Aromadendrene              | sh    | 1441| 1.8|    |    |   |
| (E)-cinnamyl acetate       | nt    | 1449| 1.1|    |    |   |
| cis-murola-3,5-diene       | sh    | 1450| 5  |    |    |   |
| α-humulene                 | sh    | 1456| 2.8|    |    |   |
| trans-cadina-1(4)-diene    | sh    | 1470| 3.7|    |    |   |
| γ-murolene                 | sh    | 1477| 1.1|    |    |   |
| β-selinene                 | sh    | 1485| 4.8|    |    |   |
| trans-murola-4(14)-diene   | sh    | 1494| 1  |    |    |   |
| Virdifllorene              | sh    | 1495| 5.3|    |    |   |
| trans-calamenene           | sh    | 1522| 15.2|    |    |   |
| δ-cadinene                 | sh    | 1523| 4.5|    |    |   |
| α-calicorene               | sh    | 1542| 6.3|    |    |   |
| Caryophyllene oxide        | os    | 1582| 1.4|    |    |   |
| iso-leptosperone           | os    | 1623| 5.6|    |    |   |
| Leptosperme oxide          | os    | 1631| 22.9|    |    |   |
| Cubenol                    | os    | 1647| 1.1|    |    |   |
| Benzylbenzoate             | nt    | 1766| 1.8|    |    |   |
| Unknown                    |       |     | 0.3| 3.2| 0| 0.1|
| Mh                         |       |     | 48.4| 10.3| 2.7| 34.4|
| Om                         |       |     | 49.2| 6.2| 0.2| 58.1|
| Sh                         |       |     | 1.8| 14.6| 6.49| 6.5|
| Os                         |       |     | 0  | 1.6| 31.7| 0.4|
| Nt                         |       |     | 0.3| 59.5| 0.5| 0.5|
| Pp                         |       |     | 0.4| 4.6| 0| 0|
| Total (unknown excluded)   |       |     | 99.7| 96.8| 100.0| 99.9|

LRI, linear retention index; Th, *Citrus bergamia*; Cz, *Cinnamomum zeylanicum; Ls, Leptospermeae sequoia; Tv, *Thymus vulgaris*; mh, monoterpene hydrocarbons; Om, oxygenated monoterpene; sh, sesquiterpenes hydrocarbons; Os, oxygenated sesquiterpenes; nt, no terpene derivatives; pp, phenylpropanoids. Other compounds were detected at <1%. They were: for Cb, α-thujene, myrcene, p-cymene, (E)-β-ocimene, terpinolene, α-terpinene, l-ocinal acetate, neral, geraniol, α-terpinyl acetate, nerol acetate, geranial acetate, trans-α-bisabolene, (E)-β-lancifolene, β-isosalbenol, bergapten, for Cz, α-thujene, β-pinene, camphen, benzaldehyde, β-pinene, α-pinene, p-methan-2,4(8)-diene, 4-terpinene, α-terpinene, (E)-ocimene; humulene epoxide II, tetradecanal; for Ls, β-pinene, myrcene, p-cymene, limonene, 1,8-cineole, γ-terpinene, isopentanoleterate, methylthio clave, cyclooctavene, β-elemene, β-copaene, α-neoclovene, alloaromadendrene, α-omorphone, α-murolene, α-bulnesene, (E)-β-farnesene, α-amorphone, trans-γ-cadinene, cadina-1,4-diene (cadinene), farnesene, (E)-nerolidol, spathulenol, Caryophyllene oxide, globulol, viridiflorol, valerianol; for Tv, α-thujene, camphene, β-pinene, 1-octen-3-ol, 3-octanone, α-phellandrene, cis-sabinene hydrate, terpinolene, bornol, p-cymene-8-ol, α-terpinolene, cis-dihydrowere, vetverene, methylcarvacrol, isobulnesene, isolimonene, carvacrol acetate, α-humulene, (E)-β-farnesene, γ-murolene, viridiflorol, trans-γ-cadinene, δ-cadinene, Caryophyllene oxide. All these compounds were considered for calculating the total percentage of each class of constituents.
were included. The microplates were incubated at 25°C (pseudomonads) and 37°C (staphylococci) for 24 hours avoiding evaporation. The assay was made in triplicate and the mode was determined.

Results

Confirmation of identification and pigment production by *Pseudomonas fluorescens*

All isolates were phenotypically identified as *Ps. fluorescens* with an API identification percentage range of 99.7%–99.9% and a T index range of 0.87-0.97. Species-specific PCR results confirmed that all the isolates belonged to the species *Ps. fluorescens*. They were able to produce blue pigment in mozzarella cheese and in its preservation liquid, beginning from 2 days after the package opening, and more abundantly at the end of the trial. The pigmentation production was evident for all isolates except for PS63 and PS282. The lack of pigmentation was confirmed for these 2 isolates also in YEGB. All the other isolates showed a visible pigmentation at all dilutions with YEGB method beginning from 48 hours of incubation. The isolation of no-pigment producing *Ps. fluorescens* from blue mozzarella cheeses indicates that cheese samples were contaminated with different strains.

Essential oils composition

GC-MS profile revealed the presence of 7, 13, 18 and 11 different compounds with a percentage higher than 1% for Cb, Cz, Ls and Tv, respectively (Table 1). The most representative compounds mainly included in monoterpenes and sesquiterpenes were linalool acetate (34.9%) and limonene (30.8%) for Cb, leptonperoxide (22.9%) and trans-calamene (15.2%) for Ls and carvacrol (29.7%), p-cymene (20.2%) and thymol (15.3%) for Tv. Cz showed the non-terpene derivative (E)-cinnamaldehyde (55%) together with the sesquiterpene β-caryophyllene (10.2%), as main compounds.

Determination of minimal inhibitory concentration against *Pseudomonas fluorescens* and *Staphylococcus aureus*

Overall, the tested biofilm-producing *S. aureus* showed to be more sensitive than *Ps. fluorescens*. Particularly, as shown in Table 2, very promising MIC values were obtained for Ls (0.012% v/v for 4 strains out of 5, among which the MRSA, and 0.024% v/v for the remaining one), which resulted even better than Tv (0.024% v/v). Cz evidenced an intermediate effect (0.049%-0.098% v/v), while Cb was poorly effective (≥ 0.781% v/v). *Ps. fluorescens* growth was not affected by Cb and Ls, which were active only at high concentrations. Also Tv and Cz showed higher MIC values than those obtained for *S. aureus* (0.098%-0.195% and 0.195%-0.391% v/v, respectively). Ethanol solvent had negligible effect on the antimicrobial activity of Cz EO (MIC: 3.125% v/v).

Table 2. Minimal inhibitory concentration of four essential oils against the tested *Pseudomonas fluorescens* and *Staphylococcus aureus*.

|   | Cb | Cz | Ls | Tv |
|---|---|---|---|---|
| Ps. fl. PS59 | 3.125 | 0.195 | 3.125 | 0.195 |
| Ps. fl. PS60 | 3.125 | 0.195 | 3.125 | 0.195 |
| Ps. fl. PS63 | 3.125 | 0.391 | 3.125 | 0.195 |
| Ps. fl. PS70 | 3.125 | 0.391 | 3.125 | 0.098 |
| Ps. fl. PS71 | 3.125 | 0.391 | 3.125 | 0.195 |
| Ps. fl. PS231 | 3.125 | 0.391 | 3.125 | 0.195 |
| Ps. fl. PS249 | 3.125 | 0.391 | 3.125 | 0.098 |
| Ps. fl. PS251 | 3.125 | 0.391 | 3.125 | 0.098 |
| Ps. fl. PS282 | 3.125 | 0.391 | 3.125 | 0.195 |
| Ps. fl. ATCC 1325 | 3.125 | 0.391 | 3.125 | 0.098 |
| S.a. PA1 | 0.781 | 0.049 | 0.012 | 0.024 |
| S.a. PA2 | 1.563 | 0.049 | 0.012 | 0.024 |
| S.a. PI1 | 1.563 | 0.049 | 0.012 | 0.024 |
| S.a. ATCC 35556 | 1.563 | 0.098 | 0.024 | 0.024 |
| S.a. ATCC 12600 | 1.563 | 0.049 | 0.012 | 0.024 |

EOs, essential oils; Cb, *Citrus bergamia*; Cz, *Cinnamomum zeylanicum*; Ls, *Leptospermum scoparium*; Tv, *Thymus vulgaris*; S.a., *Staphylococcus aureus*; Ps.fl., *Pseudomonas fluorescens*. Results are the mode of three independent trials.
more than that of its EO (Mandal and Mandal, 2011). Generally, in our study we observed that the EO resulted to be much more active on Gram positive bacteria than on Gram negative, as reported also by Maddock-Jennings et al. (2005); in addition, van Klink et al. (2005), documented in particular the remarkable antimicrobial effect of EO major compounds, β-trichetones, on MRSA. Moreover, we recently found in another study (Fratini et al., 2017) very interesting MIC results for manuka EO, comparable to those obtained with our biofilm-forming S. aureus, on different S. aureus strains from milk (methicillin-susceptible and not biofilm-producing strains). Data about bergamot EO are scanty: a study performed by Fisher and Phillips (2006) using different Gram positive and negative microorganisms as target bacteria showed that Cb EO was more effective against the Gram-positive bacteria. S. aureus was less affected, if compared with Listeria monocytogenes and Bacillus cereus. More recently, Marotta et al. (2016) evidenced a strain-dependent activity of bergamot EO on L. monocytogenes from food samples. In our research Cb EO was effective on S. aureus only at high concentrations. Overall, it is known that Gram-negative bacteria are more resistant to the EOs effect, due to the hydrophilic lipopolysaccharides contained in the outer membrane, which create a barrier against EOs hydrophobic antimicrobial compounds (Hyldgaard et al., 2012). Generally, Ps. fluorescens sensitivity to EOs has been poorly investigated. Outtara et al. (1997) reported that cinnamon EO was more active than thyme EO on Ps. fluorescens isolated from beef, whereas we found a higher activity of Tv EO. Also, we found MIC values of about 1-2 μL/mL and 2-4 μL/mL for Tv and Cz EOs respectively, against Ps. fluorescens ATCC 13525. Similarly, Mith et al. (2014) found that Ps. fluorescens ATCC 13525 was more resistant than other Gram positive and Gram-negative reference strains to the different EOs tested and among which cinnamon and thyme, with MICs ranging from 1 to 1.5 μL/mL.

Conclusions

The need for natural alternatives to synthetic chemicals is increasing in food industry, as well as the consumers request for natural products. Our study provides interesting data about EOs sensitivity of food-related wild microorganisms, which may be useful in implementing new solutions for sanitization of food facilities surfaces and/or for direct use in food and packaging, particularly in the dairy production chain.

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Official controls according to Integrated Regional Plan in Campania Region 2011/2014

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Abstract

Due to EU law, European citizens enjoy one of the highest worldwide food safety standards. Along the entire food supply chain, mandatory controls are made to ensure that plant and animal products healthiness and food and feed safety are properly labeled and comply with the strictly EU rules. The process of review of European legislation began with the promulgation of the Regulation 178/2002/EC (Official Journal of the EC no. L31 of 01/02/2002) and was completed with the hygiene package. Regulation No. 178/2002/EC (the General Food Law) laying down the general principles and requirements of food law, establishing the European Food Safety Authority and laying down procedures in matters of food safety.

Introduction

Due to EU law, European citizens enjoy one of the highest worldwide food safety standards. Along the entire food supply chain, mandatory controls are made to ensure the plants and animals products healthiness and food and feed safety, that are properly labeled and comply with the strictly EU rules.

The process of review of European legislation began with the promulgation of the Regulation 178/2002/EC (Official Journal of the EC no. L31 of 01/02/2002; European Commission, 2002) and was completed with the hygiene package.

Regulation No. 178/2002/EC (the General Food Law) laying down the general principles and requirements of food law; establishing the European Food Safety Authority and laying down procedures in matters of food safety.

The hygiene package, initially consisting of four regulations - two related to the food production and marketing (Reg. 852/04 and Reg. 853/04/EC; European Commission, 2004a, 2004b) and two on the competent authorities modality controls by the (Reg. 854/04/EC and Reg. 882/04/EC; European Commission, 2004c, 2004d) - it was later integrated to ensure a higher level of the food chain health and hygiene assurance with the EC Regulation 183/2005 laying down the requirements for feed hygiene (European Commission, 2005a), and Regulations 2073/2005/EC, 2074/2005/EC, 2075/2005/EC and 2076/2005/EC of 5 December 2005 (European Commission, 2005b, 2005c, 2005d, 2005e) on microbiological criteria, control organizations and transitional measures.

The entry into force of the European regulations in the food safety field is a cultural revolution in official controls relating to food, feed and health and animal welfare, both in relation to the legislative radical change, previously formed by a large number of vertical standards, both for the large space reserved for some basic concepts, such as the risk assessment based controls planning to rationalize and optimize the available resources. The risk assessment is the reference tool to identify a country health priorities; it must be linked to the choices of some basic parameters for the decisions taken to make food control efficient and effective. On risk assessment, in fact, some European countries, defining the acceptable levels of certain food diseases and, consequently, the acceptable levels of contamination for certain foodstuffs specific hazards.

The official controls implementation are carried out by Competent Authority (CA), which in Italy is identified with the National Health Service in its three joints (Legislative Decree 193/2007, Art. 2): Central (Ministry of Health), Regional (Departments of Health) and territorial (Veterinary Services and the Local Health Authorities SIAN).

In the NHS, the health planning functions, including those relating to the prevention and food safety, are shared between state and regions and with the Entente no. 133/CSR of 14 June 2007, the Standing Conference for relations between the State and Regions outlined the principles for the National Integrated Plan (MANCP) implementation.

The EU Regulation 882/2004, Art. 41 provides that each Member State should develop a single Control National Multi-year integrated plan, on the food safety official controls implementation, intended to verify compliance with hygiene and health food, animal feed, welfare and animal health and plants health EU Regulations Reference. These plans should be prepared in accordance with the art. 42 and 43 Regulation no. 882/2004 criteria, supplemented by the Commission Decision n. 363 of 21 May 2007.

MANCP goal ensure a high level of consumer health by reducing biological, chemical and physical incidence, promote animal health by preventing/reducing the animal diseases incidence, improve economic growth/ cohesion/competitiveness by ensuring the goods free movement, promote animal welfare farming practices to prevent hazards animal health related, minimize environmental impacts to support the sustainable development EU strategy.

The PRI (Regional Integrated Plan) It is the Campania Region basic instrument intended to use in order that, both at regional and territorial, official controls are planned in accordance with integration and optimization principles for the Community instructions implementation in accordance with the respectively State and Regions conferred powers.

PRI describes the involved people and activities, how compliance with the required standards it is ensured and review and updating measures about food safety, animal health and welfare and plants health official controls, and the related information flows.

Materials and Methods

In Campania Region activities related to food safety and animal health are managed and recorded by a regional web platform, called GISA. This is used by the CA to insert all relevant companies information: risk-based categorization, official control...
activities and nonconformities.

From 2015 considering the provisions Law 116/14 regarding farms and DGRC 623/14 on administrative offenses, it will be also used by other Authorities such as Police Forces.

GISA, accessible from the Regional Observatory for Food Safety (ORSA, 2017) website, allows: i) all Campania establishments/companies records consultation inherent to food safety and veterinary public health; ii) identified dogs and their owners records (known as the Regional Data Bank - BDR); iii) monitoring of the operational targets and other planned activities implementation; iv) evaluate the PRI suitability to deal with the food safety, veterinary public health and plant health issues.

GISA is designed for the data exchange with other informatics systems by application cooperation mode. A Region goal is to implement this mode more and more to apply the simplification, dematerialization rules and PA efficiency respecting the principle of loyal cooperation enshrined in article 22 institutional.5 law n. 241/90.

GISA allows establishments and non-compliance georeferencing according to selected parameters. It is also equipped with a specific software intended to reporting, so as to obtain a real time data aggregation to improve the regional controls whole system efficiency. The system is accessible only to official control operators who are assigned a username and password. Access, as mentioned, is also allowed to the other Authorities who carry out complementary food safety, veterinary public health and plants health, so they can enter their inspections details. This allows all control bodies to see companies records, the carried out controls and all other relevant information.

Results and Discussion

In 2011-2014, 294.557 Official Controls were carried out and recorded. The most of checks is carried out on registered and approved establishments and on farms. The official controls are represented by 91% (270.124) by simple inspection, 8% of inspections for the risk categorization of the establishments (22.463) and 0.6% by the audits (1617); the remaining 0.4% are inspections carried out for other reasons (Figure 1). In this quadrennium were found 22.926 nonconformities in whole region, detected by 15.565 official controls at 10.563 activities. The nonconformities detection is about 99.9% during simple inspections and only 0.1% during inspections carried out for establishments’ categorization.

The devices (computers, the Internet, databases, computer files, etc.) to carry out the office work and for the network access results better than previous years although several critical remain in peripherals.

Thanks to a general strengthening of cooperation and collaboration (Art. 4, paragraph 3, Regulation (EC) n. 882/2004), the official controls regional program led to a peripheral increasing activities and obligations. The audit showed a Local Health Authorities annual growth thanks to an increase of coordination activities (activation of joint working groups, meetings, technical meetings, etc.).

In general it is registered, compared with a consolidated regional system, an increase in instructions or internal operating procedures production at peripheral level. This increase, however, is still not sufficient to ensure the staff conduct uniformity.

The regional authorities, acting on the ministerial national guidelines recommendations for the Regulations 882-854/2004, drew up and adopted starting from PNI, a multi-year Official controls Regional Plan, whose task is the manage of Community rules obligations and national legislation in the field of veterinary public health and food safety.

Regional planning, generally, has seen the involvement of other agencies or institutional bodies (eg. IIZZSS, ARPA, etc.). The program of official controls in Local Health Authorities was regular and in accordance with the instructions given at the regional level.

During the audit it was verified that the CA using the main national and regional web database for the official control collection, organization and data reporting.

The CA internal audits activities at FBO-level compared to previous years were satisfactory and increased.

Otherwise it was found diffusely growing inattention on targeted training addressed to health professionals related with food safety and veterinary public health matters. In fact, in 58% of the audits carried out in 2014, it was reported lack of theoretical/practical training initiatives in audited area.

Conclusions

During the examined four years, a gradual increase has been noticed in GISA use, both in recording, both in control and sampling activities registered. The most critical master data is borne by the feed business operators and veterinary medicinal products wholesalers. A significant decrease in the risk category in almost half of approved establishments was noticed, while for registered establishments this share amounted to 39% of the total. For the latter, it should be also noted that, compared to the number of units present in the region, controls are distributed only on a small percentage. There is particularly critical in the wholesale distribution, transportation, and public catering; oppositely there is a constant monitoring on the catering and on other activities. For approved establishments control is more uniform and constant ensuring an annual regular monitoring of all active establishments in Campania region.

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