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Significant effects of two pesticides on the bacteriostatic activity and antioxidant ability of green tea polyphenols†

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Green tea polyphenols (GTPs) are widely used in food preservation because of their strong bacteriostatic activity and antioxidant ability, and whether pesticides as common pollutants in food will affect the function of GTPs is worthy of attention. Therefore, GTPs and two pesticides, namely, acetamiprid (ACE) and diguat dibromide (DIQ) commonly used on food crops were selected as research objects and Vibrio qinghaiensis sp.-Q67 (Q67) as the test organism to explore the effects of pesticide pollutants on the bacteriostatic activity of GTPs by the time-dependent microplate toxicity analysis method (t-MTA). The binary mixture systems of GTPs and two pesticides were designed by the direct equipartition ray design method (EquRay). The effect residual ratio (ERR) method was used to quantify the toxicity interactions of binary mixture systems. Besides, the effects of these two pesticides on the antioxidant capacity of GTPs were investigated. The results indicated that the bacteriostatic activity of GTPs upon interaction with the two pesticides shows certain time characteristics. These two pesticides can affect the bacteriostatic activity of GTPs, which is enhanced or weakened with prolonged duration, i.e. time-dependent synergism or antagonism. The bacteriostatic mechanism of the three substances and their mixtures is produced by affecting the cell morphology or destroying the cell structure, and the long-term antagonism of the three substances may be due to the competition for the active site. The two pesticides can greatly reduce the antioxidant capacity of GTPs. ACE reduces the free radical scavenging ability of GTPs by 14-24% and DIQ by 39-63% in the experimental concentration ratios. In addition, the free radical scavenging ability of GTPs decreases with the increase in the concentration ratio of the two pesticides in the mixture systems.

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Introduction

As we all know, green tea polyphenols (GTPs) accounting for 20–35% of the dry weight of tea are the general name of polyphenols with catechin as the main component in tea.¹ In recent years, GTPs have been widely produced and used in food preservation²,³ industry because of their good bacteriostatic activity⁴,⁵ and antioxidant ability.⁶,⁷ According to relevant statistics, total annual production of GTPs can reach 5731 tons by 2020 in China.³ The bacteriostatic ability of GTPs can effectively inhibit the spoilage of food, and its antioxidant ability can maintain the good color of fruits and vegetables. Zhu *et al.*³ found that when the concentration of tea polyphenol was 250 mg kg⁻¹, it had a good freshkeeping effect on tomatoes, and the tomatoes still had a good sensory quality after 15 days. More importantly, the appropriate dose of GTPs is harmless to the human body and has a health

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effect for humans. ^{10,11} Therefore, GTPs as food preservatives have their unique advantages.

However, many pesticides are used on food crops to promote production. Thus, pesticides have become common residual pollutants in food, especially in fruits and vegetables. Chen *et al.*¹² carried out pesticide residue examination on fruits and vegetables in Xiamen market, and found that the highest detected cypermethrin level was 13.92 mg kg⁻¹ in a sample of pakchoi cabbage. Hjorth *et al.*¹³ found that 80.4% of samples contained pesticide residues in a total of 724 samples of 46 different fruits and vegetables from eight South American countries. Swarnam and Velmurugan¹⁴ found that pesticide residues in some fruits exceeded 2 mg kg⁻¹ in Indian market. Therefore, GTPs may coexist with pesticides in food, especially in fruits and vegetables, and the effects of pesticide pollutants on the bacteriostatic and antioxidant capacity of GTPs should be considered in order to ensure the preservation function of GTPs.

This study aims to explore the effects of pesticide pollutants on the bacteriostatic and antioxidant capacity of GTPs. To do so, two commonly used pesticides, namely, acetamiprid (ACE) and diquat dibromide (DIQ) in tea were selected as research objects. A freshwater luminous bacterium *Vibrio qinghaiensis* sp.-Q67 (Q67) which is very sensitive to toxic substances was selected as the test organism. The bacteriostatic ability data of GTPs, two pesticides and their binary mixtures to Q67 were determined by the time-dependent microplate toxicity analysis method (t-MTA).¹⁵⁻¹⁹ The binary mixture system of GTPs and two pesticides was constructed by the direct equipartition ray design (EquRay) method.²⁰ In addition, the effect residual ratio (ERR)¹⁹ method based on the concentration addition model (CA)²¹⁻²⁶ was used to evaluate the interaction between pesticides and GTPs. The bacteriostatic mechanism of the three substances and their mixtures to Q67 was preliminarily determined by observing the cellular morphology.^{5,27} Besides, the free radical scavenging ability of binary mixture rays was determined by the method of salicylic acid.²⁸

Materials and methods

Chemicals

GTPs (green tea polyphenols) were purchased from Shanghai Yuanye Biotechnology Co., Ltd (Shanghai, China). Two pesticides, namely, acetamiprid (ACE) and diquat dibromide (DIQ) were purchased from the National Pesticide Quality Supervision and Testing centre of Shenyang Chemical Research Institute (Shenyang, China). All the reagents were of analytical grade and used as received without further purification. The physical and chemical properties of the three reagents, the concentration of the stock solution, and the dilution factor (*f*) are listed in Table 1. The storage solution was prepared with Milli-Q water and stored at 4 °C.

Bacterial culture

The freeze-dried luminescent bacterium *Vibrio qinghaiensis* sp.-Q67 (Q67) was purchased from Beijing Hamamatsu Corp., Ltd. (Beijing, China). The preparation of the culture medium and culture process of Q67 are detailed in the literature.¹⁶

Table 2 Components and their concentration ratios in binary mixture rays

Ray	P_{GTPs}	$P_{ m ACE}$	Ray	P_{GTPs}	$P_{ m DIQ}$
R2 R3 R4	5.605×10^{-1} 3.378×10^{-1} 2.032×10^{-1} 1.131×10^{-1} 4.854×10^{-2}	6.622×10^{-1} 7.968×10^{-1} 8.869×10^{-1}	R2 R3 R4	9.989×10^{-1} 9.978×10^{-1} 9.955×10^{-1}	1.123×10^{-3} 2.244×10^{-3} 4.478×10^{-3}

Determination of bacteriostatic activity

The bacteriostatic data of GTPs, two pesticides and their binary mixtures to Q67 were obtained by t-MTA. Using a 96 microporous plate as an experimental carrier, 200 μ L purified water was added to the surrounding 36 holes to prevent the edge effect. In the 2nd, 6th, 7th and 11th rows of 24 holes, 100 μ L purified water was added as the blank control, and the drug was diluted to 12 different concentration gradients into the remaining 36 wells by a pre-determined dilution factor. The relative luminous unit (RLU) of each hole was measured at 0.25 h, 2 h, 4 h, 8 h and 12 h, respectively, and the luminous inhibition toxicities of single components and its mixtures to Q67 were calculated using eqn (1) to reflect the bacteriostatic activity of the pollutants to Q67 at each time point. The design of the microplate is detailed in the literature. 15,16

$$x\% = \frac{I_0 - I}{I_0} \times 100\% \tag{1}$$

where I_0 is the average RLU of the blank control group and I the average RLU at each concentration gradient.

Experimental design of mixtures

There are numerous concentration ratios and concentrations of binary mixtures. The direct equipartition ray design (EquRay) method can well select the representative mixtures to reflect the

Table 1 Some physiochemical properties, stocks, dilution factors (f), the fitted results by Weibull function and some statistics of GTPs and two pesticides

Name	Abbr.	Molecular weight	CAS-RN	Stocks (mol L ⁻¹)	f	Time (h)	α^a	β^a	RMSE	R	pEC ₅₀
Green tea polyphenols	GTPs	281.36	84650-60-2	0.005	0.6	0.25	8.53	2.81	0.022	0.9979	3.17
1 71						2	7.82	2.51	0.048	0.9908	3.26
						4	7.87	2.28	0.058	0.9896	3.61
						8	6.90	1.87	0.056	0.9874	3.89
						12	7.21	1.83	0.056	0.9868	4.14
Acetamiprid	ACE	222.67	135410-20-7	0.006	0.6	0.25	4.76	1.49	0.037	0.9930	3.44
						22	4.38	1.37	0.031	0.9944	3.47
						4	4.78	1.43	0.027	0.9963	3.60
						8	4.56	1.30	0.026	0.9961	3.46
						12	4.67	1.42	0.021	0.9975	3.55
Diquat dibromide	DIQ	344.05	85-00-7	0.0002	0.4	0.25	7.33	2.00	0.020	0.9884	0
						2	15.00	4.03	0.028	0.9710	0
						4	3.15	0.68	0.092	0.9426	5.17
						8	4.99	0.88	0.100	0.9598	6.09
						12	9.68	1.48	0.071	0.9863	6.79

 $[^]a$ α and β refer to the position and shape parameters in the Weibull function, respectively. pEC₅₀ refers to the negative logarithm of the median effective concentrations.

interaction characteristics in the whole concentration range.²⁰ Therefore, binary mixture systems of GTPs and two pesticides were constructed by the EquRay method. Each mixture system contains five rays with different concentration ratios (p_i s), and each ray was diluted to 12 different concentration gradients according to the dilution factor obtained by the pre-experiment. The p_i s of each component of binary mixtures is shown in Table 2.

Concentration-effect curve fitting

The bacteriostatic data obtained by t-MTA were fitted by logit (eqn (2)) or the Weibull (eqn (3)) functions. The calculation of effective concentrations and 95% observation-based confidence intervals (OCIs) was completed using the APTox software.²⁶ The two function formulas are as follows:

$$E = 1/(1 + \exp(-\alpha - \beta \times \log_{10}(c))) \tag{2}$$

$$E = 1 - \exp(-\exp(\alpha + \beta \times \log_{10}(c)))$$
 (3)

where *E* represents the effect $(0 \le E \le 1)$, *c* the concentration of a single compound or mixture, and α and β the model parameters.

Effects of the two pesticides on the bacteriostatic activity of GTPs

The interaction between GTPs and the two pesticides was evaluated by the ERR method¹⁹ based on the CA model. The calculation of ERR was completed using the APTox software.²⁶ Mathematically, the ERR model can be formulated as:

$$ERR_x = \frac{E_x - E_{prd}}{E_{prd}} \tag{4}$$

where E_x is the fitting effect or the effect corresponding to the upper limit or lower limit of the OCIs and $E_{\rm prd}$ the effect value predicted by CA at the same concentration. The ERR with confidence intervals is noted as ERR_{imp}. When the ERR_{imp} is significantly larger/less than 0, say the interaction is synergism/ antagonism. When 0 is located between the confidence intervals, it is additive action or has no interaction.

Bacteriostatic mechanism to Q67

In order to investigate the bacteriostatic mechanism of GTPs, two pesticides and their binary mixtures on Q67, the cell morphology of Q67 exposed to GTPs, two pesticides and their binary mixtures at a concentration of EC80 for 12 h was measured. First, the Q67 bacterial suspension of logarithmic growth period was placed in a conical flask. Then, GTPs, two pesticides and their binary mixtures were added to the conical flask, respectively, so that the concentration of drugs in the suspension was equal to the EC80 of GTPs, two pesticides and their binary mixtures at 12 h. Lastly, the conical flask was cultured in a constant temperature incubator at 22 \pm 1 $^{\circ}$ C for 12 h. After 12 h, the cell morphology of Q67 was observed using a scanning electron microscope. The specific steps for the preparation of electron microscope samples can be found in the relevant literature. 5,27

The effects of two pesticides on the antioxidant ability of GTPs

The free radical scavenging ability of GTPs and two pesticide binary mixtures under different p_i s (experimental group) were determined by the method of salicylic acid.²⁸ Then, replace the pesticides in binary mixtures with an equal volume of Milli-Q water as control (blank group) to determine the free radical scavenging ability of GTPs. The greater the difference between the free radical scavenging ability of the experimental group and the control group, the greater the effects of pesticides on the antioxidant capacity of GTPs.

Results and discussion

Bacteriostatic activity of individual components to Q67

The nonlinear least square method was used to fit the concentration–response data of GTPs and two pesticides to Q67 at 0.25 h, 2 h, 4 h, 8 h and 12 h. The results indicated that the Weibull function can effectively characterize the concentration–response relationship of GTPs, ACE and DIQ (R > 0.9, RMSE < 0.1). The specific fitting results and relevant statistic parameters are shown in Table 1. The time–concentration–response curves (t-CRCs) of single components are shown in Fig. 1.

From Fig. 1, it was found that the time characteristics of the bacteriostatic activity of GTPs and two pesticides to Q67 are different. The bacteriostatic activity of GTPs increases slowly with the prolongation of exposure time within 0.25–12 h, while the bacteriostatic activity of DIQ decreases within 0.25–2 h, but increases within 2–12 h, and the bacteriostatic activity of ACE to Q67 is not affected by exposure time. Combined with Table 1, taking the negative logarithm of EC_{50} (pEC₅₀) as the bacteriostatic activity index, the bacteriostatic activity order at 0.25 h and 2 h is as follows: ACE (pEC₅₀ = 3.44–3.47) > GTPs (pEC₅₀ = 3.17–3.26) > DIQ (pEC₅₀ = 0). The bacteriostatic activity order at 4 h, 8 h and 12 h is as follows: DIQ (pEC₅₀ = 3.81–6.79) > GTPs (pEC₅₀ = 3.61–4.14) > ACE (pEC₅₀ = 3.55–3.60).

In conclusion, the antibacterial effect of GTPs and two pesticides on Q67 varies with the exposure time.^{29,30} In addition, it was noted that besides the good bacteriostatic activity of

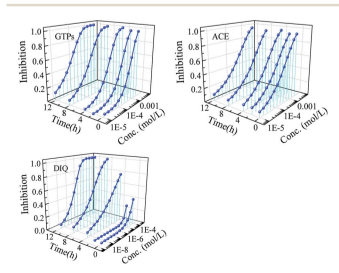
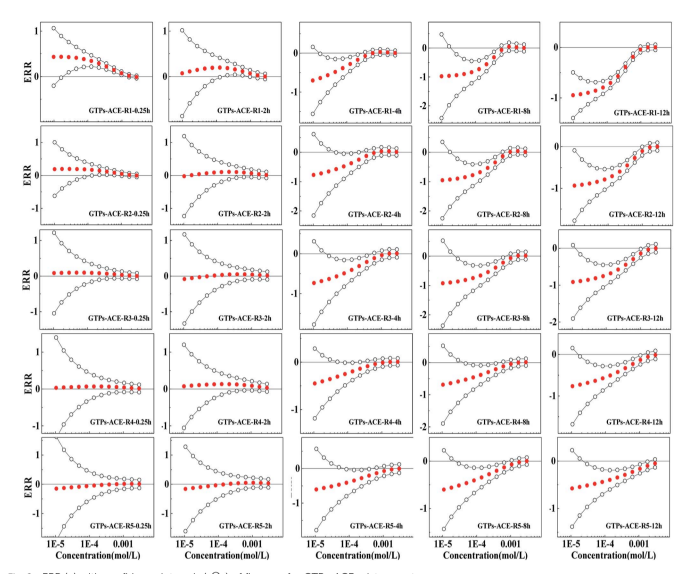


Fig. 1 t-CRCs of GTPs and two pesticides.

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ERR (•) with confidence intervals (-O-) of five rays for GTPs-ACE mixture systems

GTPs, the two pesticides, as toxic substances, have inhibitory toxicity to bacteria as they are pesticides with high toxicity to nontarget organisms.31,32

Effects of the two pesticides on the bacteriostatic activity of **GTPs**

The Weibull function can also effectively characterize the concentration-response relationship of binary mixture rays of GTPs and two pesticides (R > 0.9, RMSE < 0.1). The fitting results and relevant statistic parameters are shown in Tables S1 and S2.† The observed concentration-effect data with 95% OCIs fitted CRCs and the predicted curves by CA of representative rays are shown in Fig. S1 and S2.†

The ERR method based on the CA model can respond to the relative deviation of predicted and observed results and has the ability to quantitatively evaluate the toxicity interactions.19 Therefore, in this study, the ERR method was used to quantify the toxicity interactions between the components. The ERR_{imp} of each ray is shown in Fig. 2 and 3.

It can be observed from Fig. 2 that the five rays of the GTPs-ACE binary mixture systems show additive action or synergism in exposure times of 0.25 h and 2 h, i.e. 0 is located between the confidence intervals or ERRimp is larger than 0. However, with the prolongation of exposure time, the five mixture rays all show antagonism, and ERRimp is significantly less than 0 after 4 h. The antagonism of five rays is mainly concentrated in the middle- and low-concentration region, and the concentration region of antagonism expands with the prolonged exposure time. From Fig. 3, it can be observed that the five mixture rays of GTPs-DIQ binary mixture systems show additive action or synergism in exposure times of 0.25 h and 2 h, but with the prolongation of exposure time, the toxicity interaction changes from additive action and synergism to antagonism, which is similar to GTPs-ACE binary mixture systems. However, the antagonism of five rays of GTPs-DIQ binary mixture systems gradually shifts from the low-concentration region to the middle-concentration region with time lengthening.

Therefore, from the results of toxicity interaction analysis, it can be observed that ACE and DIQ can affect the bacteriostatic

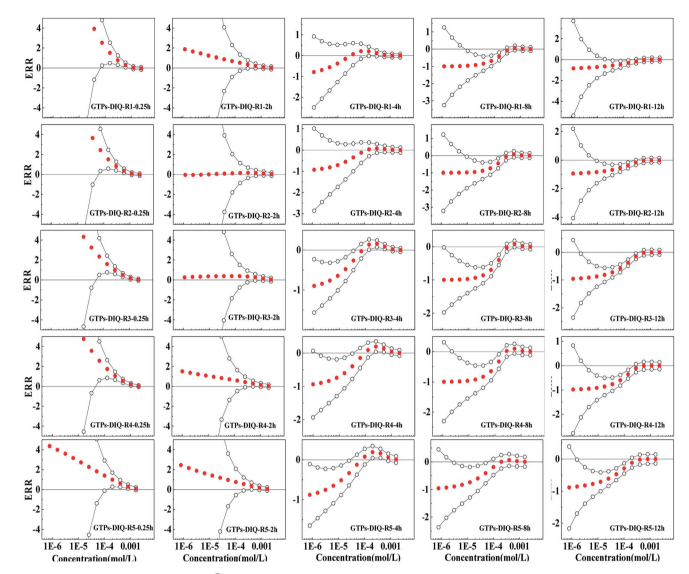


Fig. 3 ERR (•) with confidence intervals (-O-) of five rays for GTPs-DIQ mixture systems.

activity of GTPs, and the influence mode differs in different exposure times. The short-term synergism of the mixture system will strengthen the bacteriostatic ability of GTPs, while the long-term antagonism of the mixture system will lead to a decrease in the bacteriostatic ability of GTPs.

The bacteriostatic mechanism to Q67

The cell morphology of Q67 exposed to EC_{80} of GTPs, two pesticides and their representative rays for 12 h is shown in Fig. 4.

From Fig. 4, the morphology of Q67 cells is changed compared with the control group. GTPs and DIQ make Q67 cells bond with each other and destroy the membrane structure of bacteria. While ACE prolonged Q67 cells and ruptured Q67 cells. The rays with the greatest long-term bacteriostatic activity in two binary mixture systems were selected to observe their effects on the morphology of Q67 cells. It was found that mixtures also damage the morphology of Q67 cells. Besides, it was found that the amount of bacteria in the experimental

group is lower than that in the control group after centrifugation. This suggests that GTPs, two pesticides and their mixtures can inhibit the reproduction of bacteria. Ben Lagha *et al.*⁵ also found that tea polyphenols could destroy the cell structure of the bacteria. Therefore, the three substances may have similar antibacterial mechanisms, and the long-term antagonism of the three substances is due to the competition for the active site.³³

Effects of the two pesticides on the antioxidant ability of GTPs

In order to characterize the effects of two pesticides on the antioxidant ability of GTPs, the free radical scavenging ability of each binary mixture ray was determined by the salicylic acid method. The results are shown in Fig. 5. As can be observed from Fig. 5, the free radical scavenging ability of each binary mixture ray is lower than that of the control group. ACE reduces the free radical scavenging ability of GTPs by 14–24% and DIQ by 39–63% in the experimental concentration ratios. This indicates that pesticide pollutants can damage the antioxidant

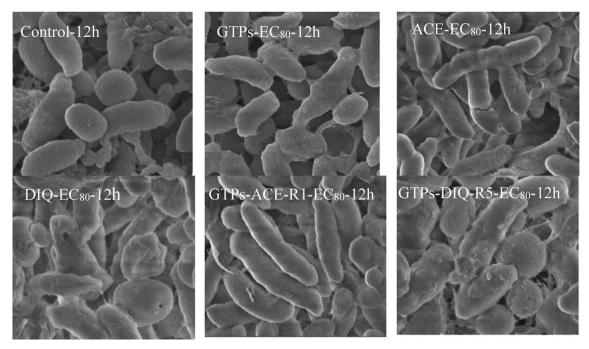


Fig. 4 Cell morphology of Q67 exposed to EC₈₀ of GTPs, two pesticides and their representative rays for 12 h.

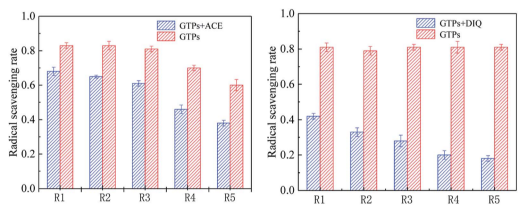


Fig. 5 Effects of ACE and DIQ on GTPs' radical scavenging ability in different concentration ratios.

capacity of GTPs, and the extent of damage is greater for DIQ than for ACE.

Besides, the reductions in the free radical scavenging ability of GTPs were linearly fitted with $P_{\rm ACE}$ and $P_{\rm DIQ}$. The results show

that there is a good linear relationship between the reductions in the free radical scavenging ability of GTPs and the concentration ratios of ACE and DIQ (Fig. 6). In addition, the free radical scavenging ability of GTPs decreases with the increase in

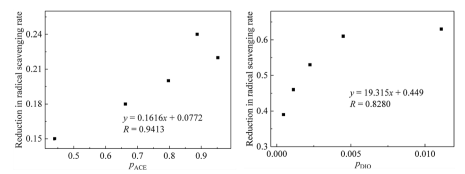


Fig. 6 Relationship between the reductions in the free radical scavenging ability of GTPs and the concentration ratio of two pesticides.

the concentration ratio of the two pesticides in the mixture systems (R > 0.8, P < 0.05).

The antioxidant capacity of GTPs plays an important role in their preservation effect.³⁴ The loss of antioxidant capacity of GTPs means that it can no longer protect food from oxidative damage. Therefore, pesticide residues in food should be avoided as far as possible in order to ensure the preservation function of GTPs, and the influence of pesticide residues in the formulation of GTP usage should also be considered.

Conclusion

GTPs and the two pesticides used all have antibacterial activity, and their bacteriostatic activity shows certain time characteristics. Both ACE and DIO can affect the bacteriostatic activity of GTPs. The bacteriostatic activity of GTPs will be enhanced when the two pesticides and GTPs coexist for a short time. However, the bacteriostatic activity of GTPs will be weakened when the two pesticides and GTPs coexist for a longer time. That is to say, the mixtures of the two pesticides and GTPs exhibit timedependent synergism or antagonism. The bacteriostatic mechanism of GTPs, two pesticides and their mixtures on Q67 involves affecting the cell morphology or destroying the cell structure. Therefore, GTPs and two pesticides may have similar antibacterial mechanisms, and the long-term antagonism of the three substances is due to the competition for the active site. The two pesticides can greatly reduce the antioxidant capacity of GTPs, and the extent of damage is greater for DIQ than for ACE. In addition, the free radical scavenging ability of GTPs decreases with the increase in the concentration ratio of the two pesticides in the mixture systems.

Conflicts of interest

There are no conflicts to declare.

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