Impact of plasma irradiation on *Tribolium castaneum*

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Abstract
Radio frequency plasma (RFP) could provide reliable, compact, cost-effective irradiation applications against insect pests of stored food and feed products. Sensitivity of red flour beetle *Tribolium castaneum* to RFP has been investigated using an irradiation applicator system with the two types of inert gases, argon (Ar) and helium (He), at 100 W for five exposure time levels of 0, 20, 40, 60 and 90 s, respectively. We demonstrated that He RFP was more efficient against *T. castaneum* than Ar RFP. In addition, a positive correlation was observed between mortality percentages of treated insect stages and exposure times for both He and Ar RFP. The adult stage showed the highest tolerance to RFP irradiation followed by larvae and pupae; however, it was more susceptible than larvae within 24 h after He RFP treatments. The optimum exposure time was 90 s with He RFP, where a full mortality at all tested stages was accomplished, while mortalities of 71.4, 65.3 and 36.7% were recorded for pupae, larvae and adult stage, respectively, after an Ar RFP treatment of 90 s. In case of treated adults, the reproduction rate was higher than treated larvae and pupae. Our findings indicated that He RFP was an effective method for inhibiting *T. castaneum* development and impacting the insect life cycle and could be considered a promising tool for pest control of stored food.

Keywords Non-thermal plasma · He and Ar gases · Stored product pest · Mortality · Reproduction rate · Red flour beetle

Key message
- Radio frequency plasma (RFP) irradiation is a promising treatment against pest insects.
- Helium (He) as a precursor gas is more effective against *T. castaneum* than argon (Ar).

Introduction
The red flour beetle *Tribolium castaneum* (Herbst) is considered a destructive pest that attacks most of stored products and economic commodities (Hagstrum 2016). It is ubiquitous in many regions around the world (Sokoloff 1977). Chemical fumigants that have been extensively used against *T. castaneum* and other stored product pests, in particular in developing countries, have detrimental effects on the ecosystem and its fauna, in particular ozone layer and human health (Zettler and Arthur 2000). Due to these environmental and health concerns, several pest control technologies have been proposed as alternatives, such as heat treatment (Arbogast 1981), essential oils (Saroukolai et al. 2010), ozone (Xinyi et al. 2017) and microwaves (Lu et al. 2010). However, drawbacks of these alternative pest control methods against *T. castaneum* include loss of nutrients and deterioration of food quality, long treatment times, temperature requirements that limit applications and high costs (Sileem et al. 2017). Over
the last few decades, ionizing radiation has been proposed to protect commodities and stored products against insect pests. It is a process, where infested stored products and commodities are irradiated in order to sterilize, kill or prevent development of insect pests (Hallman et al. 2010; Morrison 1989). Ionizing radiation is successfully and increasingly applied as a disinfestation treatment that is produced via the disintegration of radioactive atomic nuclei. Although it is a simple, robust and well-established technology, its sources are not readily available in many regions and might become more difficult to obtain in the future as well, because the irradiation sources are decaying over time. Accordingly, attempts to find more green and effective pest control strategies have focused on the application of sustainable energy sources against stored product pests. Electron beam has been suggested to be used as a source of radiation (Codex Alimentarius Commission 2003), but the high applied doses of electron beam irradiation required to kill targeted insects may have unacceptable effects on product quality. Recently, another form of radiation, cold or non-thermal plasma, has been suggested to combine properties/effects of irradiation treatment with a sustainable energy source—which is one of the major challenges in radiation processes (Kaur et al. 2020). Plasma, the fourth state of matter, is an ionized gas containing free electrons. Ionized gas is generated by the addition of energy such as intense radiation, radio frequencies, extreme heat or electrical energy (Eliezer and Eliezer 2001), resulting in a combination of charged and neutral particles such as electrons, ions, atoms and molecules (Amami et al. 2008). Depending on the process used to generate the plasma, the neutral particles, electrons and ions can have different temperatures (Chen 1984). Non-thermal plasmas are a mixture of cold ions and neutral (< 60 °C) and energetic electrons (≥1 eV). They are generated by electric discharge in a gas at vacuum condition, lower pressure or atmospheric pressure (Niemira 2012; Thirumdas et al. 2015). Radio frequency cold atmospheric plasma is generated by circulating radio frequency currents in antennas or coils, applying a radio frequency voltage across two parallel electrodes and immersed in the plasma by a dielectric window. Coupling of the electrons with electromagnetic fields facilitates energy that transferred them to sustain the plasma. The efficiency of the coupled power into the charged particles, as well as the plasma uniformity, is determined by the radio frequency excitation design (Chabert and Braithwaite 2011).

Many advantages are connected with non-thermal plasma irradiation such as low heat, short release time and relatively short process time. In addition, it involves few variables and has comparably low equipment costs. Therefore, atmospheric plasma operations have been extensively studied and its applications have gradually increased in various fields due to its appreciated economic efficiency (Tyata et al. 2012). In order to ensure its efficient adoption as a disinfestation treatment at industrial level as a safe and reliable alternative approach compared to ionizing radiation, significant research effort is still needed. It is difficult to optimize an optimal dose protocol against microorganisms and insect pests that infect food products where very varied equipment and operational environments have been used, resulting in very different plasma properties (López et al. 2019).

Recently, non-thermal plasma has been increasingly used in various industrial processes. It has been applied for medical applications such as sterilization, cancer treatment, wound healing and blood coagulation (Graves 2012). The effectiveness of non-thermal plasma as a technology for improving the shelf-life of food products by eliminating the microbial contamination from fresh and minimally processed food has been investigated, and it was observed that some species of Salmonella and bacterial spores were inactivated on dry food surfaces by non-thermal plasma (Fernández et al. 2012; Hertwig et al. 2018). It is also suggested as a promising method to protect dry food against stored product pest insects (Abd El-Aziz et al. 2014; Donohue et al. 2006; Keever et al. 2001; Mishenko et al. 2000; Mohammadi et al. 2015). The surface electrostatic excitation of membranes due to plasma may have a negative effect on nervous and neuromuscular systems of insects. This phenomenon may be attributable to the exposure to high voltage discharge and to the creation of anoxia, which may serve to anesthetize and immobilize insects (Bures et al. 2005; Donohue et al. 2006). Another effect of plasma on insects might be the breakdown of C-H bonds in the lipid layer of the insect cuticle that causes dehydration of the insect and can lead to its death (Donohue et al. 2008).

Non-thermal plasma exhibits reactive oxygen species (ROS) which includes high reactive chemicals such as peroxides, superoxide, oxygen and hydroxyl radicals, and nitric oxide radicals (Ji et al. 2019) that have a negative impact on pest insects and may intensify the secondary metabolism of plant products that could maximize the economics of post-harvest treatment (Bußler et al. 2015; Laroussi and Leipold 2004). The effectiveness of plasma against microorganisms was determined by the type of gas used to generate the plasma and therefore responsible for free radical formation. It has been reported that cold plasma had no impact on the quality of dry stored products. Mortality-responses of T. castaneum were recorded at 2500 V for 1–5 min exposure times without significant changes in the color of the refined wheat flour (Mahendran et al. 2016). Furthermore, plasma treatment of wheat grains caused effective decontamination of bacteria and fungi together with insect pests and also enhanced the shelf life, germination and initial state of growth of the grains and resulted in an increased final grain yield. Moreover, the production of qualitatively better dough was also observed (Scholtz et al. 2019; Tyata et al. 2012).
this context, it was aimed to investigate the susceptibility of *T. castaneum* to radio frequency-induced cold atmospheric plasma (RFP) as an alternative pest control method with the following main objectives, (1) Confirm the efficacy of RFP as a killing method (2) Determine the sensitivity profile of different developmental stages exposed to different RFP doses using two different precursor gases. (3) Examine the mortality of irradiated *T. castaneum* where insect development was inhibited (4) Determine the reproduction rate resulting from RFP treated larvae and pupal stages.

**Materials and methods**

**Insects**

The red flour beetle, *T. castaneum* colony was maintained for several generations under controlled laboratory conditions of 27 ± 2 °C and 70 ± 2% RH and continuous darkness. It was initially established from adults collected from infested storage wheat in Cairo governorate. The colony was fed on a rearing medium composed of wheat flour and brewer’s yeast (19:1, w/w) (Ayvaz et al. 2002). The wheat flour was disinfested at 60 °C for 10 h to eliminate possible contaminants (Tunçbilek and Kansu 1996).

**Generation of radio frequency-induced cold atmospheric plasma (RFP) irradiation**

An RFP irradiation source was designed as shown in Fig. 1. It consisted of a generator of 13.65 MHz radio frequency (RF) with an automatic tuner, a gas inlet, plasma chamber and a vacuum system. The RF antenna was fed 0–600 W.

Both the RF generator and the auto-matching network were from T&C Power Conversion Inc, USA. Around the source chamber (Bottle), two coils allowed an on-axial magnetic field which provided 0–200 Gauss. This magnetic field confined the plasma electrons which increased the plasma density. The sample holder was a Pyrex bottle (250 ml) in which indirect plasma irradiation was performed in a batch system. A pressure of 150 kPa was used. In the present experiment, helium (He) and argon (Ar) were utilized as precursor gases to create plasma, respectively.

**RFP treatment and subsequent bioassay of irradiated *T. castaneum* stages**

15- to 17-day-old larvae and one- to three-day-old pupae and 6- to 7-day-old adults were exposed to the plasma for 0 (control), 20, 40, 60 and 90 s at a power level of 100 W using helium and argon gases. For exposure, the larvae, pupae and adults were transferred separately into the Pyrex bottle, 30 larvae, 20 pupae and 30 adults, respectively, for each replicate. Three replicates were run for each dose (20, 40, 60 and 90 s) and for the control groups (0 s). The mortality of adults and larvae was recorded 6, 12, 18 and 24 h after exposure to plasma. Pupal mortality was not recorded because the dead pupae could not be distinguished during these experimental times. The remaining plasma-treated larvae, pupae and adults were then maintained in the laboratory until death to observe their mortality or development of the treated larval and pupal stages to pupae and adults, respectively. The cumulative mortality was calculated at the end of adult survival as well as larval and pupal durations.

Remaining living adults per replicate that emerged from irradiated larvae and pupae, as well as treated adults, that had survived RFP treatments, were transferred using a fine brush to glass jars containing 20 g of rearing medium. These remaining adults were observed for survival and progeny as an indication of reproductive ability, where the reproduction rate of surviving adults, defined here as “the ability of adult previously irradiated as larvae and pupae as well as irradiated adults to produce progeny,” was determined by placing those adult survivors on a rearing medium and counting the next generation (F1 progeny) produced at 43–44 days. They are expressed as number of insects per F1 progeny.

**Statistical analysis**

Data were analyzed using one way analysis of variance (ANOVA) technique and the means were analyzed using Duncan’s multiple range test, the ANOVA statistics were significant (*p* < 0.01) (Steel and Torrie 1960). The data of mortality (%) were transformed by arcsine tables, while the means and standard errors of reproduction rate were from original data. The reproduction rate averages were calculated according to Aldryhim and Adam (1999) and Régnière et al. (2012), with the methodology described by Carey (1993) as follows: R = Ne/Ns where Ns is the number of adults at the beginning of generation (P1 generation); Ne is the number of adults produced in the next generation (F1 generation).
The trendline labels were created by Excel 2010. The \( \text{LT}_{50} \) values were estimated from the total mortality-response data (defined here as the lethal plasma exposure time to kill half the population) which were expressed as a percentage transformed using probit analysis (Abbott 1925) to allow for a direct comparison of insect mortality due to differences in susceptibility to the plasma discharge by different gases and different developmental stages of the insects exposed to the plasma for differing lengths of time.

Results

Effect of RFP irradiation on \( T.\text{castaneum} \) stages

Mortality of larval, pupal and adult stages of \( T.\text{castaneum} \) at the end of each developmental stage (accumulative mortality) after exposure to RFP are shown in Fig. 2. Results indicated that both Ar and He RFPs have a lethal effect on larval, pupal and adult stages. Significant mortalities of 1.2 ± 0.00, 11.4 ± 0.00, 18.7 ± 0.00, 43.3 ± 0.02 and 65.3 ± 0.03% \( (F_{4,10} = 191.9, \ p > 0.0005) \) were obtained in the case of Ar RFP treated larvae, mortalities of 3.2 ± 1.59, 35.1 ± 1.92, 38.5 ± 0.28, 48.3 ± 1.88 and 71.4 ± 1.33% \( (F_{4,10} = 35.9, \ p > 0.0005) \) for pupae and mortalities of 4.1 ± 0.0, 8.9 ± 1.20, 11.1 ± 1.01, 33.3 ± 1.05 and 36.7 ± 1.45% \( (F_{4,10} = 48.2, \ p > 0.0005) \) for adults at 0, 20, 40, 60 and 90 s, respectively. Significant mortalities of 3.7 ± 0.00, 18.4 ± 0.02, 39.5 ± 0.05, 73.8 ± 0.04 and 100% \( (F_{4,10} = 1241.3, \ p > 0.0005) \) were also observed for He RFP treated larvae, mortalities of 6.8 ± 1.59, 47.7 ± 0.33, 47.8 ± 0.72, 89.1 ± 0.60 and 100% \( (F_{4,10} = 197.5, \ p > 0.0005) \) for pupae and mortalities of 0, 10.2 ± 0.33, 20.5 ± 0.66, 64.4 ± 0.33 and 100% \( (F_{4,10} = 191.9, \ p > 0.0005) \) for adults, respectively. The results indicated that the mortalities increased with increasing treatment time for both gases. Moreover, it was observed that He RFP was more effective than Ar RFP, where full mortality was obtained for larvae, pupae and adults after 90 s with He RFP, while relative mortalities were recorded in these stages at the same exposure times treated with Ar RFP.

Evaluation of the reproduction rates of insect stages irradiated with RFP

Figures 3 and 4 are depicting the effect of Ar and He RFP treatments, respectively, on the ability of irradiated larvae, pupae and adults to procreate a next generation compared to untreated insects as expressed by reproduction rates. The reproduction rates of larvae amounted to 4.4 ± 0.5, 3.4 ± 0.3, 1.6 ± 0.17 and 1.2 ± 0.16 \( (F_{4,10} = 10.9; \ p = 0.0013) \) upon Ar RFP treatment and 3.7 ± 0.76, 2.9 ± 0.65, 1.6 ± 0.57 and 0.0 \( (F_{4,10} = 19.9; \ p = 0.0001) \) upon He RFP treatment for 20, 40, 60 and 90 s, respectively. They were significantly reduced in comparison with the larvae reproduction rate of the control treatment (5.1). Similarly, the reduction of the reproduction rate was significant when the adult stage was treated with He RFP (7.5 ± 0.60, 4.4 ± 0.58, 4.1 ± 0.10 and 0.0; \( F_{4,10} = 27.4, \ p < 0.0005) \) and Ar RFP (4.9 ± 0.34, 4.9 ± 0.05, 4.3 ± 0.63 and 3.4 ± 0.10; \( F_{4,10} = 10.8; \ p = 0.0012 \)) for 20, 40, 60 and 90 s, respectively, as compared to control treatment (8.8). Moreover, the reproduction rate was significantly decreased in pupae treated with He RFP (3.1 ± 0.10,
Fig. 4 Reproduction rates (Avg.) of larval, pupae and adult stages treated with different exposure times of RFP using He gas

2.8 ± 0.33, 2.4 ± 0.29 and 0.0; \( F_{4,10} = 10.6, p = 0.0032 \)
and was also decreased for Ar RFP (4 ± 0.57, 3.3 ± 0.35, 2.6 ± 0.32 and 2.5 ± 0.28; \( F_{4,10} = 5.2, p = 0.0153 \)) for 20, 40, 60 and 90 s, respectively, as compared to the control treatment (4.5 ± 0.29). The results indicated that the reproduction rates of treated adults were the highest in comparison with treated larvae and pupae for both Ar and He RFP treatments. Moreover, for Ar RFP, the reproduction rates of treated larvae were higher than for pupae at 20 and 40 s, while the rates were higher in case of treated pupae than larvae at 60 and 90 s. Similarly, the reproduction rates were lower for treated pupae than for treated larvae at exposure times of 20 and 40 s with He RFP, while it was lower for He RFP treated larvae than pupae after 60 s.

**Determination of the lethal exposure time (LT) of He RFP treatment on *T. castaneum* stages**

The median lethal exposure time (Lt50) in linear trend was calculated to obtain clear information about the susceptibility of insect stages to He RFP (Fig. 5) Lt50 of pupae (26.3 s) was drastically lower than of larvae and adults (39.0 s and 53.8 s), respectively. Results indicated that the pupal stage was susceptible by twofold compared to adults and by 1.5-fold compared to larvae.

**Mortality assay of *T. castaneum* stages 24 h after He RFP treatments**

The stored product pest management efficiency depends on instantly getting rid of larvae and adults. In this context, the mortality within 24 h upon He RFP treatment was determined (Fig. 6). It was observed that the mortality not only increased with increasing the exposure times of RFP but also varied with time up to 24 h after treatments. 6 h after treatment for 90 s with He RFP, intermediate mortalities of 42.3 ± 1.17 and 38.9 ± 1.31% were observed for larvae and adults, respectively, while no lethal effects were recorded at exposure times of 20 and 40 s (Fig. 6a). This trend continued 12 h after RFP treatment, where the mortality was drastically increased to 96 ± 2.08 and 98 ± 1.15% for larvae and adults, respectively, for 90 s treatment with He RFP as compared to mortalities of 11.2 ± 0.91 and 7.5 ± 1.04% for 60 s (Fig. 6b). Also, high mortalities of 98.5 ± 1.04 and 100.0% were obtained 18 h after He RFP treatment of larvae and adults, respectively, for 90 s compared to mortalities of 25.0 ± 1.76 and 27.8 ± 1.85% for 60 s. In the same time frame (18 h after treatment), no mortality (0%) was recorded for larvae treated for 20 and 40 s, however, the mortality percentages were 8.4 ± 0.64 and 9.3 ± 0.38% for treated adults, respectively (Fig. 6c). As shown in Fig. 6d, full mortality (100%) was obtained for both larvae and adults treated for 90 s 24 h after treatments compared to mortalities of 53.4 ± 0.65 and 64.4 ± 6.41% for larvae and adults, respectively, treated for 60 s. In the same time frame (24 h after treatment), the mortality of adults exposed for 20 and 40 s was 10.2 ± 0.63 and 20.5 ± 0.63%, respectively, while still no mortality was observed for the larval stage. These obtained data revealed that the mortality of the adult stage was higher than that of the larval stage of *T. castaneum*, both 18 and 24 h after plasma treatment. The obtained results for mortalities of *T. castaneum* larvae and adults within 24 h after treatment with He RFP indicated that adults were less...
tolerant than larvae at sub-lethal doses (Fig. 6). These differences in tolerance within 24 h after treatment were converted in the accumulated mortality-responses that were recorded at the end of the larval duration and adult survival (Fig. 1).

Discussion

Results depicted in Fig. 2 are in accordance with other studies. For example, an inactivation of *T. castaneum* adults induced by Ar RFP was shown by Sileem et al. (2020) and also Carpen et al. (2019) who found a varied mortality induced by Ar/oxygen and Ar/nitrogen plasmas, where the nitrogen-containing plasma resulted in a higher mortality than argon/oxygen plasmas, while the He RFP was effective for the control of green peach aphids (Bures et al. 2005) and *T. castaneum* adults (Sileem et al. 2020). Evidently, plasma intensity and exposure times have different impacts on different insect stages and species. The power of 2500 V for 15 min at a 3.7 cm distance between electrodes induced full mortality of larvae and adult *T. castaneum* stages (Ramanan et al. 2018). Abd El-Aziz et al. (2014) found that a decrease in the distance of the plasma jet electrodes to the sample and increasing pulse numbers lead to increased mortality of *P. interpunctella* pupae. Nevertheless, at the maximum number of pulses and minimum distance, a larvae mortality of 86% was not exceeded. The impact of non-thermal plasma on *Blattella germanica* has been investigated by Donohue et al. (2008). They observed slightly opened wings, flaccid antennae, twitched legs and broken abdomen at high plasma doses, while lower doses caused limited responses such as lack of normal movement of treated insects. Malformation of *T. castaneum* larvae and partial and whole internal melanization of hemolymph has also been reported due to non-thermal plasma treatment (Ramanan et al. 2018). It was also suggested that the oxidative stress of plasma may damage the larval cuticle and epidermis and attract hemocytes to the affected area resulted from damage signals. The insect defense response by disintegration of fat bodies to heal the damage might cause the clotting and melanization. Similar responses were observed by Ferreira et al. (2016) on *D. melanogaster* larvae who reported that the effect of plasma...
on larval death might be extreme oxidative stress due to ROS, as well as the melanization caused by the activation of the immune system.

The various responses to radiation treatment of *T. castaneum* stages observed in our study were also obtained by Tunçbilek et al. (2003) in response to gamma rays, who found that the larvae were more susceptible than the adult stage. Also, Sileem et al. (2017) observed that the adult stage of *Oryzaephilus surinamensis* exposed to low doses of gamma radiation resulted in less sterility than those previously irradiated as larvae, while the irradiated pupae showed the highest sterility. The same trend was recorded by Aldryhim and Adam (1999) who found that the adults of *Sitophilus granarius* previously treated as pupae with cold plasma had a lower reproduction rate than the adults originating from treated larvae.

Comparing data after Ar and He RFP treatments revealed that there are differences in the reproduction rates of treated adults and also the adults that produced from treated larvae and pupae. Our results indicated that a reduction of the reproduction of treated insects with sub-lethal doses may be due to physiological changes in particular DNA damage due to the impact of active ingredients created by plasma such as ROS and charged particles and could be referred as indirect effect of plasma treatment. These differences of reproduction rates might due to differences in reactive species of He and Ar plasmas. For example, Kwon et al. (2019) found a different distribution of different ROS in two atmospheric plasmas. However, they could not confirm the role of ROS that resulted in insect death after plasma treatments. Furthermore, Kawasaki et al. (2015) found that the ROS amount varied depending on the equipment used for plasma generation. It was observed that in case of cold plasma treatments of *T. castaneum*, the differences of applied voltage, electrode distances and exposure time caused differences in mortality (Mahendran et al. 2016). The insect mortality, reproduction rate and insect sterility after ionizing radiation treatments have been investigated (Hallman et al. 2010). These responses were attributed to the direct effect of ionizing radiation on DNA and their indirect effect due to released free radicals attacking DNA chains (Hutchinson 1985). These effects on DNA were also investigated with UV radiation (Ravanat et al. 2001) and non-ionizing electromagnetic fields (Saliev et al. 2019). In this context, the mechanism of cold plasma which creates many active ingredients may contribute to DNA damage of insect cells. More research will be required to examine whether non-thermal plasma is inducing this damage. Our results presented a reduction of F₁ generation (reproduction rate), which indicates latent effects of plasma on treated insect expressed as negative impact on insect fertility rather than direct mortality. This effect may be attributed to the reactive species in particular the reactive oxygen species (ROS) mediated by plasma that cause indirect effects on DNA, immune system and circulatory system of insects. This hypothesis is going in line with the impact of ROS produced by chemical treatment (Kumar et al. 2015), thermal treatment (Lubawy et al. 2019) and low doses of gamma radiation (Zhikrevetskaya et al. 2015) on DNA damage of *P. interpunctella*, *Gromphadorinha coquereliana* and *Drosophila melanogaster*, respectively. Another possible explanation is the direct impact of plasma on the nervous system (Donohue et al. 2006). Moreover, in order to maintain homeostasis, the antioxidant enzymes of the defense system were significantly changed in *P. interpunctella* exposed to non-thermal plasma due to the free radicals (Abd El-Aziz et al. 2014). However, to confirm these mechanisms further, biochemical and molecular genetic studies are needed.

Our results regarding lethal exposure times are contradictory to Keever et al. (2001) who suggested that pupal and adult stages were more tolerant to plasma because they have a hard exterior and puparium for protection against plasma radiation. Their statement only refers to the direct effect of plasma radiation that was thought to be effective only at the target’s surface, but this hypothesis does not take into account the indirect effect due to ROS. It has been shown that ROS of cold plasma can penetrate into deeper layers of model bacteria biofilms resulting in DNA damage (Govaert et al. 2020). This finding supports our hypothesis of inflicted damage via plasma penetration of pupal cells similar to the penetration through meat and crevices and cracks of seeds (Misra et al. 2019). This indirect effect of ROS could explain the finding by Nasr et al. (2020) who recorded that the adults of *T. castaneum* and *Sitophilus granaries* were the most susceptible to direct cold plasma treatments of up to 25 min. Although these results contradict our findings, their study on the tolerance of larval and pupal stages goes in line with the results of our study.

It is clear that the adult stage was more susceptible for the direct effect of RFP that was visible within 24 h after treatment than the larval stage, while it was more tolerant than the larval stage to the indirect effect of RFP caused by reactive species that could cause free radicals with oxidizing effects in treated insect cells. These radicals attack the cell molecules, which may have time-delayed lethal effects on the insect after the plasma treatment. This effect has been demonstrated in case of gamma radiation treatments which caused different mortality-responses during the life span of treated insects. However, it was observed that the capacity of total antioxidants of *Drosophila melanogaster* stages was nearly constant (Patil et al. 2017). Oxidizing effects in *P. interpunctella* larvae treated by cold plasma have been reported that caused a significant reduction of protein and glutathione contents, as well as high levels of lipid peroxide. In comparison with the other insect stages, the plasma treatment resulted in a high sensitive reaction of the larval stage.
(Abd El-Aziz et al. 2014). The response of larvae, pupae and adults to that time-delayed effect of plasma treatment will be different by its variance in the intrinsic physiological characteristics and the immunity system. To confirm our hypothesis, further biochemical and molecular research is required to identify the repair enzymes involved in the different insect stages in relation to the induced DNA damage.

Conclusion

Radio frequency-induced cold atmospheric plasma (RFP) can be considered a promising alternative for irradiation processes against pest insects. The sensitivity of T. castaneum to the two types He and Ar RFP was investigated and compared. He RFP is more effective in inactivating T. castaneum at all three developmental stage than Ar RFP. However, both He and Ar had a toxic effect on larvae, pupae and adults. The mortality gradually increased with increasing exposure time for both types of plasma. Moreover, both types Ar and He RFP had latent effects on insect stages, where they were effective in reducing the reproduction rate of treated adults and the adults produced from treated larvae and pupae. The optimum exposure time of RFP for preventing development and procreation was 90 s applying He RFP. The susceptibility of insect stages to He RFP treatment differed. The pupal stage was the most susceptible followed by larval and adult stages. However, the adult stage was more susceptible than larvae within 24 h after treatments. Our results indicated that RFP could successfully be used for managing the T. castaneum pest due to both toxic and latent effects. But further research is required to evaluate the ability of non-thermal plasma to penetrate the various food products for using as a phytosanitary treatment rather than surface treatment. In addition, the effects of cold plasma treatments on insects and their fertility need to be studied in more detail on a molecular and a species-specific level in order to design targeted plasma treatments with the desired effects on specific insects while conserving the stored goods. Research is also required on plasma treatment of infested food under typical storage conditions and volumes. Treatment conditions need to be determined where the respective treated food remains unaltered while the targeted pest insects are inactivated. In addition, upscaling and economic as well as environmental aspects need to be considered.

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Authors’ contributions WAS, TMS and RSH conceived and designed research. TMS and RSH conducted experiments. WAS, TMS and RSH analyzed data. WAS and BAR wrote the manuscript. All authors read and approved the manuscript.

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Compliance with ethical standards

Conflicts of interest The authors declare that they have no conflict of interest.

Ethics approval This article does not contain any studies with human participants or vertebrates performed by any of the authors.

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