INTRODUCTION

The Delphinium L. genus (Ranunculaceae), with almost 300 species worldwide, consists of 29 species in Turkey (17 endemic to Anatolia). Their usages have a long history of medicinal use. Dioscorides mentioned their use against lice and scorpions. The powders of Delphinium staphisagria L. and Delphinium peregrinum L. were used against lice by the British army in the battle of Waterloo. The dried and mature seeds of Delphinium staphisagria, known as “kokarot, kokarotu, müzüdek, mevezek and mevzek” (Baytop 1999) were used as emetics, purgatives and sedatives (in the treatment of rabies, tetanus, and epilepsy) in traditional Turkish medicine. The usage was left aside due to their high toxicity. Despite their toxicity, their external applications (as ointment, powder form or infusion (1-
was collected from Köprübaşı-Trabzon (DW1, DW2) and D. consolida L. (syn: Consolida orientalis (Gay) Schröd.) and D. regalis (L. syn: Consolida regalis S.F. Gray subsp. regalis) are antiparasitic (Ulubelen et al. 2001; Hiller and Melzig 2006).

Vector-borne diseases cause epidemics, leading to serious human health problems. Aedes aegypti (Culicidae), one of these vectors, transmits viruses like Dengue, Yellow fever, Zika, and Chikungunya. According to WHO, yellow fever is an acute viral haemorrhagic disease transmitted by infected mosquitoes, and Dengue is a mosquito-borne viral infection, which causes a flu-like illness, and evolves on occasion into lethal complications. WHO also emphasized that, the symptoms of the infection of Zika virus, transmitted primarily by Aedes mosquitoes, are skin rash, mild fever, muscle and joint pain, headache, conjunctivitis or malaise, and is also a cause of Guillian-Barré syndrome and microcephaly (Masi et al. 2017; WHO February 2018; WHO May 2018; WHO April 2019). Many interventions are conducted to prevent these diseases, and one of them is integrated mosquito control, including the implementation of personal protection, destruction and reduction of its habitats, and insecticidal treatment regimens against adult and larval mosquitoes for the reduction of the virus spread. Reducing the amount of synthetic insecticides or pesticides applied, is preferred because of their undesirable and toxic effects. Frequent chemical interventions can also cause the development of insecticide resistance (Tabanca et al. 2013a; Masi et al. 2016). Therefore, an increasing number of plant-based extracts, essential oils or phytochemicals are being investigated with the aim of determining an effective agent against A. aegypti (Kamar et al. 2010; Pitarokili et al. 2011; Maheswaran and Ignacimuthu 2012; Liu et al. 2012; Kumar et al. 2012; Tabanca et al. 2013a,b; Reegan et al. 2015; Cantrell et al. 2016; Masi et al. 2016; Dias et al. 2017; Carroll et al. 2017; Chantawee and Soonwera 2018; Stappen et al. 2018; Tabanca et al. 2018, 2016a, 2016b).

The alkaloids, one of the most remarkable groups of natural products in these plants, have a wide variety of biological activities. The structures of over a hundred alkaloids have been identified; many of them have elicited an expected anticancer activity (Lu et al. 2012). In addition to the studies on cytotoxic and anticancer effects (Liu et al. 2017; Nugroho et al. 2015; Chanakul et al. 2011), several investigations into insecticidal activities have been published (Bandara et al. 2000; Garcez et al. 2009; Liu et al. 2012; Masi et al. 2017). The diterpene and norditerpene alkaloids have also shown insecticidal activity (Ulubelen et al. 2001; Kukel and Jennings 1994). Also based on previous studies in literature, the cytotoxicity and insecticidal activities are related to each other, and diterpene alkaloids can be protective agents for parasite control strategies (Gonzalez-Coloma et al. 2004; Reina and Gonzalez-Coloma 2007).

This preliminary study was presented to assess the cytotoxic and insecticidal potentials of extracts, obtained from Delphinium formosum an endemic species in Trabzon (Turkey), which has a role in traditional medicine and causes serious poisoning. The different extracts of its aerial parts, roots, and flowers were investigated for their cytotoxic activity against the human liver carcinoma cell line (HepG2) and primary human umbilical vein endothelial cells (HUVEC). The mosquitocidal activity of the cytotoxic extract against Ae. aegypti was also determined for the possibility to use its lethal potential as an insect repellent.

MATERIALS AND METHODS

Plant material

Delphinium formosum was collected from Köprübaşı-Trabzon and Macka-Trabzon, in August 2011 and July 2012. The voucher specimen has been deposited in the Herbarium of the Faculty of Pharmacy, Istanbul University (ISTE 102747; 98086).

Extraction procedure

Dried and powdered roots, collected in the first year, were percolated with EtOH in four portions (10 L, 3 L, 2 L, 5 L) and each portion was evaporated to dryness at 40°C in vacuo (DW1, DW2, DW3 and DW4). The residues of the portions (DW1 and DW2) were acidified to pH1 by 200-300 mL 0.5 N H₂SO₄ and extracted with CHCl₃. These CHCl₃ extracts were evaporated to dryness (DW1A1K and DW2A1K). The acid solutions were basified with 5% NaOH to pH10 and extracted a second time with CHCl₃. The extracts were evaporated to dryness, and named as DW1A2K and DW2A2K (Figure 1).

Dried and powdered aerial parts, flowers and roots, collected next year, were percolated with EtOH in one portion and evaporated to dryness at 40°C in vacuo. The same process was applied. Two different chloroform extracts of each part (HA1K, HA2K, FA1K, FA2K, 2DWA1K, and 2DWA2K) were acquired (Figure 2).

Biological Assays

Cytotoxicity Assays

Cytotoxicity test, cell lines, culture conditions and treatments

Roswell Park Memorial Institute Medium 1640 (without glutamine; RPMI-1640) with glutamine, 10% FCS and penicillin / streptomycin mixture was used for the human liver carcinoma cells (HepG2). Primary human umbilical vein endothelial cells (HUVEC) were isolated from umbilical cords by digestion with 0.01% collagenase A solution (Roche) and grown in Endothel-
niales Cell Growth Medium with a supplement mix (Promocell; C-39215) containing 10% FCS, penicillin (100 U/mL) / streptomycin (100 mg/mL) mixture, and kanamycin (50 mg/mL). Umbilical cords were obtained with the consent of patients (permission by the local ethics committee). All cells were grown at 37°C and 5% CO₂ in a humidified cell incubator. The culture medium was changed every 2 days. Monolayer cells grown to 75-85% confluence were detached with trypsin-ethylenediamine tetraacetic acid to make single cell suspensions, and the viable cells were determined using the trypan blue exclusion test and diluted with medium to give a final density of 10⁵ cells/mL. The passage number range for HepG2 cell lines was maintained between 21 and 26. The passage number for primary HUVEC cells was maintained between 3 and 5. The extracts were initially dissolved in dimethyl sulfoxide (DMSO) and the stock solution of each extract was prepared at a concentration of 20 mg/mL. Next, the stock solutions were diluted in the medium to obtain final concentrations of 10-400 μg/mL. 100 μL of cell suspension per well were seeded into 96-well plates at a plating density of 10⁵ cells/well for the HepG2 cells and 2x10⁴ cells/well for HUVEC, and incubated to allow for cell attachment at 37°C and 5% CO₂ for 24 h. After 24 h, the medium was aspirated, and the cells were treated with 100 μL of each dose. Permethrin mixture of 46.1% cis and 53.2 trans isomers (Chemservice, West Chester, PA) was used as a positive control of ethanol or DMSO was included. Assays were repeated at least three times on separate days using different hatches of eggs.

**Cytotoxicity studies**

For MTT tests, the cells were then incubated with 150 μL MTT (0.5 mg/mL in medium) solution for 3 h. After the removal of the MTT solutions, the formed formazan crystals were solubilized in 80 μL of DMSO and then, the absorbance was measured at 550 nm and at 690 nm (as control wavelength) using a microplate reader.

The cell viability in treated cells compared to that of negative controls was calculated. Then, the half maximal inhibitory concentration (IC₅₀) was expressed as the sample concentration that caused an inhibition of 50% in cytotoxicity in the cells calculated by extrapolation. The percentage of cell viability was calculated with respect to solvent control as follows:

% Cell viability = \( \frac{{\text{Abs}_{\text{Compound}}}}{\text{Abs}_{\text{Solvent Control}}} \times 100 \)

The results were expressed as cell death (%) compared to the negative control.

**Insecticidal Activity**

**Mosquito Colony**

_Aedes aegypti_ used for testing were pesticide susceptible, and provided by the CMAVE insectary. The “Orlando1952” strain was collected near Orlando, Florida, USA in 1952, and has been in continuous laboratory colony for 64 years. Rearing procedures are standardized and have been described previously (Tabanca et al. 2016b).

**Larvicidal Activity**

Larvicidal activity testing was performed essentially as described previously (Pridgeon et al. 2008), but the assay was modified for 96-well plates to conserve limited amount of test samples by the use of smaller volumes (Masi et al. 2017). The HA1K extract was diluted in dimethyl sulfoxide (DMSO) to make 100 mg/mL. Mortality was determined in the larval assays at four different concentrations (1.0, 0.5, 0.25, and 0.1 μg/μL) in a final volume of 200 μL of larval rearing media. For each assay, a positive control of permethrin stock and a negative control of ethanol or DMSO was included. Assays were repeated at least three times on separate days using different hatches of eggs.

**Adulticidal Activity**

The toxicity of HA1K extract was tested in assays against adult _Aedes aegypti_ using cohorts of 3-6-day post-emergence females as described previously (Pridgeon et al. 2008). Mosquitoes were cold anesthetized on ice, and groups of 10 females sorted into individual plastic cups. An application of 0.5 μL of the appropriate dilution of the test chemical was made by repeater pipettor (Hamilton PB600) with a 25 μL blunt tip glass syringe (Hamilton 7100 series) to at least twenty females at each dose. Permethrin mixture of 46.1% cis and 53.2% trans isomers (Chemservice, West Chester, PA) was used as a positive control, and acetone was used as negative control. After treatment, the mosquitoes were kept in plastic cups at 24-26°C and 80% humidity, and supplied with 10% sucrose in water for 24 h prior to recording mortality.

**RESULT AND DISCUSSION**

Due to the diterpenoid alkaloid content of _Delphinium_ species, we expected distinct cytotoxic actions when testing their effect on HepG2 cells. Only one extract (HA1K) was active among them, in the studied concentration ranges. The same extract was also examined against the HUVEC cells. The IC₅₀ values of HA1K against two cell types (244.9 μg/mL for HepG2 and 144.4 μg/mL for HUVEC) were similar.

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**Figure 2.** Representative scheme of extraction procedure of the aerial parts and flowers, collected in the first year, and the roots, collected in the second year.
The MTT assay demonstrated the effect of the extracts on the cell viability of HepG2 and HUVEC cell lines as shown in Figure 3.

The HA1K extract, found cytotoxic against both cell lines, was further investigated for its insecticidal activity (Table 1). The extract was evaluated at the dose of 5 µg/mosquito and had 97% mortality against adult Aedes aegypti. The positive control permethrin at 6.33 µg/mL resulted in 53±11% mortality, while the negative control acetone resulted in 0% mortality. In larval activity, the HA1K extract showed 80% mortality at the highest dose of 1 µg/mL against 1st instar Aedes aegypti; however, the mortality tittered off quickly at the lower doses. Negative control mortality in larvicidal assays was 0% for DMSO, and positive control permethrin resulted in 100% at the 47.4 µg/mL. Since the negative control acetone resulted in 0% mortality. In larval activity, the HA1K extract showed 97% mortality against adult Aedes aegypti, isolation of the active compound/s through bioassay-guided fractionation and its/their characterization may be promising.

This is the first investigation on the cytotoxicity and insecticidal activity of D. formosum. On the other hand, several studies on the cytotoxic effect of the isolated compounds from other species exist in the literature. De Ines et al. isolated 43 norditerpenoid alkaloids from Aconitum, Delphinium and Consolida species for their cytotoxic effects on the tumor cell lines SkMel28 (human malignant melanoma), HeLa (human cervical adenocarcinoma), SkMel25 (human melanoma), CT26 (murine colon adenocarcinoma), and SW480 (human colon adenocarcinoma) and the non-tumor cell line CHO (Chinese hamster ovary cells). Browniine, ajadelphinine, 8-O-methylcolumbianine, dehydrotakosamine, lycoctonine, 14-deacetylajadine, pubescine, 14-deacetylpubescine, 1,14-diactetylcardiopetaline, 18-CHO (Chinese hamster ovary cells). Browniine, ajadelphinine, (human colon adenocarcinoma) and the non-tumor cell line (human cervical adenocarcinoma), SkMel25 (human melanoma), in a previous study (Tanker and Ozden 1975). Durust et al. also isolated certain norditerpenoids and norditerpenoid alkaloids from D. formosum, Tribolium castaneum (Herbst.) to evaluate the repellent effects. Twenty-one alkaloids exhibited promising insect repellent activity (Ulubelen et al. 2001). Hetsine, a diterpen alkaloid, was found to have the most active repellency (59.12%). The compounds, delsemin-B, lycoctonine and browniine, showed also an activity against Tribolium castaneum (Ulubelen et al. 2001).

In a previous study from the Black Sea area (Trabzon) in Turkey, D. formosum was collected and, delsemine A/B, 14-demethylajadine (N-acetyldelectine), lycoctonine, antranoyllycoctonine, delcosine and delectine were identified from the aerial parts (Merici et al. 1996). Additionally, lycoctonine and delcosine were isolated from D. formosum roots, collected in Trabzon, in a previous study (Tanker and Ozden 1975). Durust et al. (1999) also isolated some phenolic acids such as p-coumaric, p-hydroxy benzoic, caffeic, protocatechic and vanillic acids in light of the cytotoxic activity. The difference of the cytotoxicity of D. formosum (aerial parts and roots) can be explained with the diversity of secondary metabolites in the root-extracts. In light of it all, it gives the impression that the extracts may contain different percentages of the active substances.

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**Figure 3.** The % cell viability values of the HA1K against HepG2 and HUVEC cell lines.

**Table 1.** The mortalities of the HA1K against adult female mosquito Aedes aegypti and 1st instar Aedes aegypti larvae

<table>
<thead>
<tr>
<th>Adult female mosquito Aedes aegypti</th>
<th>1st instar Aedes aegypti larvae (% mortality)</th>
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<tbody>
<tr>
<td>5 µg/mosquito (% mortality)</td>
<td>1 µg/µL 0.5 µg/µL 0.25 µg/µL 0.1 µg/µL</td>
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<tr>
<td>96.7±5.8</td>
<td>80 53.3±23.1 13.3±11.5 0</td>
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</tbody>
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Sen-Utsukarci et al. The cytotoxicity and insecticidal activity of extracts from Delphinium formosum Boiss. & Huet
stem bark. (Diptera: Culicidae).

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