Time and dose-dependent effects of *Viscum album quercus* on rabbit spermatozoa motility and viability *in vitro*

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26 Short title: Viscum album quercus affects spermatozoa motility

28 Summary

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The target of this study was to evaluate the effect of extract of the European mistletoe –
 Viscum album quercus L. on spermatozoa motility and viability *in vitro*. The CASA

32 system was used to determine the spermatozoa motility parameters at different time $\frac{1}{2}$ integrals (0, 1, 2, and 2, b) and an arrestored with ility range determined in first different

intervals (0; 1; 2 and 3 h) and spermatozoa viability was determined in five different
doses of *Viscum album quercus* L [10 (QA); 6.6 (QB); 3.3 (QC); 2.5 (QD) and 2 (QE)

mg/mL]. Results in experimental groups detected a significant deterioration on rabbit

36 spermatozoa after 1, 2 and 3 hours, compared to the control. The initial total

37 spermatozoa motility showed increased value for all doses of *Viscum album quercus* in

- 38 comparison to control. After *in vitro* culture a dose–dependent decrease (QA: reduction
- of 69.7%; QB: reduction of 40.9%) was found. For the progressive spermatozoa most
- 40 significant decrease (86.8% for QA vs. 48.5% for QB) was detected compared to the
- 41 control after 3 hours of culture. Spermatozoa viability (MTT test) was decreased in all
- 42 experiment groups at the end of experiment, but the differences were not significant.
- 43 Significant alterations of membrane integrity were found in groups with the highest $V_{integrity}$ alternative concentration (OA OB) but concerns integrity should be
- *Viscum album quercus* concentration (QA, QB), but acrosome integrity showed no
 significant changes. Results suggest negative dose- and time-dependent effect of
- 46 *Viscum album quercus* at higher doses on spermatozoa motility and viability parameters
- 40 *viscum album quer* 47 *in vitro*.
- 48

- 49 Key words: Viscum album; spermatozoa; CASA; viability; in vitro
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51 Introduction

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53 Anticancer preparations made from plants have been an object of scientific interest for

- many years. It is worth noting that as many as 25% of cytostatics used in the anticancerchemotherapy are obtained from plants.
- 56 Extracts of the European mistletoe (*Viscum album* L.) have been widely used for
- 57 decades as alternative, complementary treatment (Kovacs *et al.* 2006, Felenda *et al.*
- 58 2019, Suveren *et al.* 2017). In clinical practice mistletoe therapy is often given
- 59 concomitantly to conventional chemotherapy. Mistletoe plants are generally growing on
- 60 different host trees, like apple, oak, or pine. Cytotoxic glycoproteins, the mistletoe
- 61 lectins, are active component of mistletoe extracts and can stimulate effector cells of the
- 62 innate and adaptive immune system (Stein *et a*l. 2002, Braedel-Ruoff 2010, Gren and Earmichti 2012, Gren and Massanyi 2016)
- 63 Formicki 2013, Gren and Massanyi 2016).
- Experiments also indicate a statistically significant increase in albumin fraction level
- and lymphocyte count. Moreover, decrease of the total protein content, protein fractions
- globulins alpha2, beta, gamma and neutrophil, monocyte count in mouse serum wasobserved (Gren 2009).
- 68 The reproductive ability and the semen quality of animal species can be affected by
- 69 many environmental sources, as well as age, stress, hormonal status, nutrition and
- toxins (Mangelsdorf et al. 2003, Lukac et al. 2011, Mousa-Balabel and Mohamed 2011,
- Fallas-López *et al.* 2011, Tirpak *et al.* 2017, Saha *et al.* 2019). In the many years,
- 72 following the increased success rate of cancer treatments, great efforts have been made
- to improve quality of life in survivors, including fertility preservation in young patients
- 74 (Masopotova *et al.* 2018). Because of their gonadotoxic effects, chemo– and
- radiotherapy can temporarily or permanently compromise fertility (Di Bisceglie *et al.*
- 76 2013). Oncological treatments present severe gonadotoxic effects on both germ and
- T7 Leydig cells. Of note, in a significant percentage of patients (20 50%)
- spermatogenesis is impaired even before cancer treatments, probably due to the
- 79 malignancy itself. The recovery of normal spermatogenesis after treatment may require
- several years, and mainly depends on various factors initial spermatozoa count, type
- 81 and dose of specific oncological treatments and patient age. Disturbance of homeostasis
- of reproductive system in some diseases can be seen from the decline in physical and
- chemical parameters of spermatozoa, such as pH, semen volume, concentration,
- 84 motility, and the percentage of spermatozoa viability. These data justify the increasing
- 85 efforts in identifying prevention and treatment strategies to preserve reproductive
- functions in young men with malignancies (Colpi *et al.* 2004, Maltaris *et al.* 2006;
- Vitku *et al.* 2015; Heráček *et al.* 2018). The mechanisms underlying the male infertility
 of *Viscum album* extracts have not been investigated.
- 89 The objective of this *in vitro* study was to determine the effect of various concentrations

90 of *Viscum album quercus* during various time periods (0 - 3 h) on the selected

- 91 parameters of rabbit spermatozoa motility and viability.
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96 Materials and methods

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98 Animals, semen samples and *in vitro* culture

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100 Male rabbits (n=10; New Zealand White) kept under standard conditions at the 101 Experimental Station of the Animal Production Research Centre Nitra, Slovak Republic 102 were selected on the basis of age normally associated with reproduction (12 - 14)103 months). Animals were housed in a partially air-conditioned rabbit house (Animal 104 Production Research Centre, Nitra) under a photoperiod 16L : 8D (minimum light 105 intensity of 80 lux). Animals were kept in individual cages and fed with a commercial 106 diet and were provided water *ad libitum*. An air temperature of $20 \pm 2^{\circ}$ C and relative

humidity of $70 \pm 5\%$ was maintained in the rabbit house. Conditions of their care,

manipulations and use corresponded to the instruction of EC no. 178/2002 and related
 EC documents, and they were approved by local ethics committee.

110 Semen samples (n = 5) in five replicates were collected on a single day (early in the

111 morning) with the help of artificial vagina (Krockova *et al.* 2012, Parkanyi *et al.* 2015).

112 Immediately after collection the individual doses of semen exhibiting a white color

113 without presence of any gel and artificial particles, were mixed together to obtain

114 pooled sample. The spermatozoa concentration in semen was $0.40 - 0.63 \times 10^9$ per mL.

The obtained semen samples were diluted according to routine methods (Chrenek *et al.*2007, Roychoudhury and Massanyi 2008).

Later the spermatozoa were incubated in thermostat $(37 \pm 0.5^{\circ}C)$ with various

118 concentrations of Viscum album quercus (Iscador Qu 10 mg; Weleda, Verein für

119 Krebsforschung Institute Hiscia – Arlesheim, Switzerland) dissolved in physiological

solution. Concentrations of the Iscador Qu were based on manufacturer's values and

121 later diluted with physiological solution to reach decreased experimental concentration 122 (6.6 - 2.0 mg). The control (QK) group was cultured only with physiological solution.

122 (6.6 - 2.0 mg). The control (QK) group was cultured only with physiologic 123 The scheme of experiments is presented in Table 1.

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125 **2.2.** Computer-assisted semen analysis

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Spermatozoa motility was used as an indicator of cell quality. The motility analysis was 127 carried out using a CASA (Computer Assisted Semen Analysis) system - Sperm Vision 128 TM program (MiniTub, Tiefenbach, Germany) with the Olympus BX 51 microscope 129 (Olympus, Tokyo, Japan) at cultivation times 0, 1, 2 and 3 hours (Time 0 - 3). Each 130 sample was placed into the Makler Counting Chamber (depth 10 µm, Sefi-Medical 131 132 Instruments, Haifa, Israel (Massanyi et al. 2008). This study was performed in five replicates at each concentration. At least 1000 spermatozoa were analyzed in each 133 sample (Lukac et al. 2013, Halo et al. 2018). Using the rabbit specific set up the 134 following parameters have been evaluated – total motility (MOT; %), progressive 135 motility (PRO; %), distance average path (DAP; μ m), distance curved line (DCL; μ m), 136 distance straight line (DSL; µm), average path velocity (VAP; µm/s), velocity curved 137 line (VCL; µm/s), velocity straight line (VSL; µm/s), straightness (STR), linearity 138 (LIN), wobble (WOB), amplitude of lateral head displacement (ALH; µm) and beat-139 cross frequency (BCF; Hz) as described previously (Roychoudhury et al. 2010, Tvrda et 140 141 al. 2015 Adamkovicova et al. 2016).

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144 Viability analysis – MTT test

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Viability of rabbit spermatozoa cultured with Viscum album quercus was evaluated by 146 the metabolic activity (MTT) assay after 3 hours of culture. This colorimetric assay 147 148 measures the conversion of 3-(4.5-dimetylthiazol-2-yl)-2.5-diphenyltetrazolium bromide (MTT; Sigma-Aldrich, St. Louis, USA) to purple formazan particles by 149 mitochondrial succinate dehydrogenase of intact mitochondria of living cells. Formazan 150 was measured spectrophotometrically by a microplate ELISA reader (Multiskan FC, 151 ThermoFisher Scientific, Finland). The data are expressed in percentage of control. 152 Results from the analysis were collected during four repeated experiments for each 153 concentration (Slanina et al. 2016). 154

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156 Viability – membrane integrity – Eosin-nigrosin

157 158 The spermatozoa viability was evaluated using eosin-nigrosin staining methods (Slanina et al. 2018). From all the samples smears were prepared after 3 hours of culture. 159 Experimental samples and the control sample were diluted in the ratio 1:2:2 with 5% 160 161 eosin (Eosin Y) and 10% nigrosin (Nigrosin) solution (both Sigma-Aldrich, St. Louis, USA). For each slide 300 cells were counted under a light microscope (1000×, Leica 162 163 DMIL LED; Leica Microsystems CMS GmbH, Germany) and classified as viable 164 (intact membrane) and dead (damaged membrane). The experiment was realized in six replicates. The results of viability evaluation were expressed as the percentage of viable 165 and dead spermatozoa (in %). 166 167

168 Acrosomal integrity

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The acrosomal status was assessed after 3 hours of culture following the fast green-rose 170 Bengal staining protocol designed by Pope et al. (1991). This single-step staining 171 method applies a mixture consisting of 1% fast green (Sigma-Aldrich, St. Louis, USA), 172 1% rose bengal (Sigma-Aldrich, St. Louis, USA) and 40% ethyl alcohol (Centralchem, 173 174 Bratislava, Slovak Republic) in 0.1 M citric acid - 0.2 M disodium phosphate buffer (Sigma-Aldrich, St. Louis, USA). Twenty microliters of the sample were mixed with 20 175 µl of the staining solution and incubated for 70 s at room temperature. Ten microliters 176 177 of the mixture were smeared on a tempered glass slide and air-dried at 37°C. Acrosomal integrity was evaluated using bright field microscopy at 1000x using oil immersion. At 178 least 200 cells per slide were evaluated for the presence or absence of acrosome, and 179 expressed as a percentage rate (Tvrda et al. 2017). 180

182 Statistical analysis

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184 The control group (medium without *Viscum album quercus*) was compared to the 185 experimental groups. Statistical analysis was carried out using the GraphPad Prism

program (version 3.02 for Windows; GraphPad Software, La Jolla California USA).

187 Descriptive statistical characteristics (mean, standard deviation) were evaluated at first.

- 188 One-way ANOVA with Dunnett's post-test was used for statistical evaluations. The
- 189 level of significance was set at *** (p < 0.001), ** (p < 0.01) and * (p < 0.05). For
- 190 individual measurements average value (x), minimum (min) and maximum (max) value,
- standard deviation (SD) and coefficient of variation (CV) were recorded.

192 **Results**

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194 Spermatozoa motility

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196 The initial spermatozoa motility (Time 0) showed increased value for all doses of Viscum album quercus in comparison to the control group. Statistically significant 197 increase was observed (p < 0.05) in the sample QA (77.23 \pm 8.64%), and also (p < 198 0.001) for QB ($81.17 \pm 8.08\%$) and QC ($80.87 \pm 8.83\%$). After 1 h of the culture the 199 average spermatozoa motility in control group was $72.17 \pm 11.94\%$. At the same time a 200 statistically significant (p < 0.001) decrease (23.8%) was observed in group QA (54.98 201 \pm 15.53%). After 2 hours of *in vitro* cultivation significantly decreased (p < 0.001 and 202 0.01) total spermatozoa motility in comparison to the control group was observed in 203 groups QA and QB (doses 10 and 6.6 mg/mL). Also, after 3 hours of in vitro cultivation 204 a dose-dependent decrease (QA: $19.39 \pm 7.7\%$; reduction of 69.7%; QB: $37.84 \pm$ 205 18.29%; reduction of 40.9%) was found (Figure 1). 206

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208 Progressive spermatozoa motility

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At Time 0 the average progressive spermatozoa motility was higher in all experimental 210 211 groups with Viscum album quercus in comparison with the control ($44.67 \pm 13.03\%$). These changes were statistically significant. After 1 hour of culture the progressive 212 spermatozoa motility was the highest in the control group QK ($60.97 \pm 13.98\%$) and a 213 statistically significant decrease was detected in groups QA and QB. This decrease was 214 compared to control 42.4% for QA and 24.2% for group QB. After 2 hours of *in vitro* 215 culture, the progressive spermatozoa motility was significantly reduced in groups QA 216 and QB with a decrease up to 53% in QA ($22.43 \pm 16.34\%$) and 29.5% in group QB 217 218 $(33.63 \pm 13.31\%)$. Even more dramatic reduction (86.8% for QA vs. 48.5% for QB) was detected compared to the control $(45.12 \pm 19.61\%)$ after 3 hours of culture (Figure 2). 219

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221 Distance parameters

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Spermatozoa distance path parameters confirm the negative dose and time-dependent
effect of *Viscum album quercus*. Spermatozoa distance average path significantly
decreased at Time 1 in experimental groups QA, QB and QC. Later (Time 2) a
significant decrease was detected in almost all experimental groups (except the lowest
concentration) and at Time 3 only in groups with the highest *Viscum album quercus*concentrations (Table 2).

229

Spermatozoa distance curvilinear line showed similar trends as spermatozoa distance
path with various significant decrease at all time periods in experimental groups
compared to control (Table 3).

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Spermatozoa distance straight line significantly decreased at all time periods in groups
QA - QC. Only at Time 2 group QD showed significant decrease (Table 4).

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240 Velocity parameters

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Also the spermatozoa velocity parameters were affected by the addition of *Viscum album quercus* (Iscador Qu) to the culture medium. The spermatozoa velocity curved

line at all time periods in control group was $97.58 - 131.47 \,\mu$ m/s. After a significant

increase at Time 0 (groups QB, QD) a signifacnt decrease was found in all experient

groups. Later the most significant decrese was detected in group QA compared to control $(131.47 \pm 16.38 \text{ vs. } 93.25 \pm 10.19 \text{ at Time } 1:108.93 \pm 17.72 \text{ vs. } 70.77 \pm 19.69$

control $(131.47 \pm 16.38 \text{ vs. } 93.25 \pm 10.19 \text{ at Time 1}; 108.93 \pm 17.72 \text{ vs. } 70.77 \pm 19.69$ at Time 2 and 97.58 ± 21.79 vs. 47.86 ± 21.66 at Time 3). Also in group QB a

significant decrease was detected after 1, 2 and 3 hours of culture (Figure 3).

250

The spermatozoa velocity average path clearly confirm the data for spermatozoa velocity curved line with similar significant decrease (Table 5).

253

Also the spermatozoa velocity straight line was negative affected by the *Viscum album quercus* addition. In group with the highest concentration this parameter decreased at Time 1 to 55.64%, at Time 2 to 54.00% and at Time 3 to 47.77% compared to control (Table 6).

258

259 Other fine motility parameters

260

The initial amplitude of lateral head displacement (ALH) was $3.78 \pm 0.63 \mu m$. At Time 261 0 a statistically significant increase was detected in all experiment groups. After 1 hour 262 of cultivation the ALH was in control group $4.70 \pm 0.44 \ \mu m$ and the values decreased 263 by 15.1% for group QA and 15.9% for QB. After 2 hours the ALH was the highest in 264 control groups $(4.16 \pm 0.44 \,\mu\text{m})$ and the most significant decrease was noted in group 265 QA $(3.24 \pm 0.88 \ \mu\text{m}, \text{p} < 0.001)$ with a decrease of 22.1%. After 3 h of *in vitro* 266 cultivation, the lowest value was found in the sample QA ($2.72 \pm 1.16 \mu m$) compared to 267 control $(3.98 \pm 0.79 \,\mu\text{m})$ and was statistically significant (Figure 4). 268

269

270 The average value of the initial beat cross frequency (BCF) was 29.79 ± 2.98 Hz. After 271 1 h of *in vitro* cultivation the BCF in QK group increased to 31.91 ± 2.60 Hz. In all experimental samples a statistically significant decreased was found (time-dependent). 272 273 However, after 2 hours the BCF in control groups (QK) reached 29.55 ± 2.18 Hz. The most significant decrease (25.2%) was found for group QA (22.11 \pm 8.10 Hz). After 3 274 hours the BCF in the control sample was 28.46 ± 2.75 Hz. In the group QA a 275 statistically significant decrease (34.6%; 18.62 ± 8.49 Hz), and also in the group OB 276 $(17.9\%; 23.36 \pm 6.03 \text{ Hz})$, compared to the control group was detected (Figure 5). 277 278

The straightness of spermatozoa movement was in all time periods significantlydecreased only in group QA (Table 7).

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282 Very similar trends were found for spermatozoa linearity (Table 8).

283

The spermatozoa wobble was significantly affected only in groups QA and QB (Time 1 and 2) with the highest difference detected at Time 1 (0.51 ± 0.03 in group QK vs. 0.44 ± 0.03 in group QA (Table 9).

288 Viability, membrane integrity, acrosomal integrity

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After 3 hours of incubation, viability of rabbit spermatozoa showed decreased values in
all doses of *Viscum album quercus* in comparison to the control group, but the
difference were not significant (Figure 6).

293

Significant decrease of membrane integrity (intact) was found in groups with the highest *Viscum album quercus* concentrations QB ($64.00\pm2.65\%$; p < 0.05) and QA ($60.33\pm5.51\%$; p < 0.01) compared to control group (QK) ($79.33\pm1.53\%$). Also, in groups QE, QD, QC lower values in comparison to control group were detected (p < 0.05; Figure 7).

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The values of acrosomal integrity showed very similar tendency (94.00 - 97.33%) in all
experimental groups with no significant difference (Figure 8).

303 **Discussion**

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305 Researchers are unanimous about the negative impact of various factors on male reproduction and cell/spermatozoa quality (Burton 2013, Lukac et al. 2013, Petrovova 306 307 et al. 2014, Lukacova et al. 2015, Kolarova et al. 2017). The mistletoe (Viscum album L.) extract has been shown to be an effective complementary drug in the treatment of 308 cancer patients after surgical removal of the primary tumor. It improved survival, 309 recovery from damage caused by irradiation or cytostatic therapy, and quality of life. In 310 animal tests, clear anti-carcinogenic effects of Iscador were demonstrated. Mainly 311 immune stimulation but also direct cytotoxic activity is believed to be responsible for 312 the anti-carcinogenic activity of Iscador. Tests examining the acute toxicity, genotoxic 313 314 effects as well as effects on reproduction showed no adverse effects of Iscador preparations. Genotoxic effects and effects on reproduction, which cannot be evaluated 315 in clinical use, have been cleared up in animal tests. Iscador was shown to be clearly 316 317 non-genotoxic and free of relevant toxic effects on reproduction in vivo (Maldacker 318 2006, Gardin 2009).

319 On the other hand, it has been stated that anticancer drugs could cause harmful effects 320 on the spermatozoa quality and spermatogenic cell arrangement of male. Therefore, this

study was designed to evaluate the *in vitro* effects of *Viscum album quercus*

322 (Iscador) on rabbit spermatozoa in order to assess possible beneficial and/or toxic

effects of this compound. Rabbits were chosen as the experimental animal in this

research for their well-defined reproductive systems (Paal *el al.* 2014, Rafay and

Parkanyi 2016). Some herbal extracts have been proven to have effects on male

326 infertility, for example, gossypol, papaya seed, neem oil, neem seaweed and

verbascoside (Mosher and Pratt 1991, Dehghan *et al.* 2005, Roychoudhury *et al.* 2009,
Okab *et al.* 2013, Vizzarri *et al.* 2019).

- 329 Since decades, *Viscum album* preparations have been used in Europe in oncology. They
- show multi-facetted anti-tumor *in-vitro* activities, which include inhibition of tumor cell

331 proliferation, induction of apoptosis, inhibition of angiogenesis, modulation of immune

- 332 competence and gene signature expression. Recently, it was demonstrated *in vitro* that
- 333 *Viscum album* exerts an anti-inflammatory effect, mostly directed to chronic

334 inflammation by selectively inhibiting cytokine-induced expression of cyclooxygenase-

335 22 (Seibert *et al.* 1989, Bussing 2006, Bussing *et al.* 2008, Hegde *et al.* 2011, Hajto *et*

- al. 2011). Iscador as an anticancer drug used in the treatment of a variety of neoplastic
- 337 lesions. On the other hand, treatment with *Viscum album quercus* (Iscador) is
- accompanied by different toxic effects on different body organs.
- An *in vitro* study investigated the effect of a standardized mistletoe preparation on the
- action of Trastuzumab, a drug used for the treatment of Her-2 positive breast cancer. A
- 341 dose dependent anti-proliferative effect of *Viscum album* exctract (VAE) was observed
- at concentrations $\geq 10 \ \mu g/mL$ after 3 days of incubation. After 7 days a significant
- 343 growth inhibition of 60% with the clinically relevant concentration 1 μ g/mL was
- detected and no proliferating cells were left at VAE concentrations of 10 and 100 μ g/ml.
- 345 With 10 μ g/ml VAE after 7days the proportion of early apoptotic cells raised from
- 9.0 % in the control to 17.4 % and that of late apoptotic/necrotic cells from 18.7 % in
 control to 78.7 %, respectively (Weissenstein *et al.* 2016).
- 348 Effects of Viscum album quercus (VA Qu) extract in various doses were also
- 349 investigated in an *in vitro* model with tumor cells: three multiple myeloma (MM) cell
- lines (OPM-2, RPMI-8226, U-266) and three B cell lymphoma cell lines (U-698,
- 351 DOHH-2, WSU-1). *Viscum album*/Qu extract markedly downregulated the membrane
- expression of IL-6R and gp130 in RPMI-8226 (down to 29% and 32%) and the
- expression of gp130 in WSU-1 (down to 22%). There was a marked reduction of viable
- cells of RPMI-8226 (down to 28%) and WSU-1 (down to 8%) at 100 μ g/10⁶ cells /ml.
- There was a clear relationship between the inhibition of proliferation and viability. VA Qu was more effective in cells having a high proliferation rate than in those with a low proliferation rate (Kovacs *et al.* 2006).
- 358 Our observations indicate that Iscador in higher doses decreases the spermatozoa
- motility, progressive spermatozoa motility and some other fine spermatozoa motility
 parameters (velocity curved line, amplitude of lateral head displacement and beat cross
 frequency). Thus, Iscador (in high doses *in vitro*) can have long-term effects on
- 362 testicular function and can be a potent gonadal toxic drug.
- 363 Spermatozoa motility is one of the most important effective factors in male fertility. The
 364 mechanism by which Iscador affects the spermatozoa motility has not been clearly
 365 elucidated.
- 366 A decline in fructose level due to alteration in carbohydrate metabolism after Iscador
- 367 Qu treatment can be suggested (Rigau *et al.* 2001, Gren *et al.* 2011). Reason for
- decreased spermatozoa motility, can be also a decreased level of androgen carrier
- 369 proteins involved in spermatozoa motility. Mistletoe lectins have been identified as
- 370 main active components and exhibit cytotoxic effects. Therefore, probably lectins have
- a negative effect on the spermatozoa motility found in this study. Also a relationship
- between diminished spermatozoa quality and anticancer treatment may a result of a
- series of cascade events that cause a fall in intracellular ATP levels (Luria *et al.* 2002,
- Turner 2006, Storey 2008), release of different apoptogenic factors (as pro-caspases,
- 375 cytochrome C, and apoptosis inducing factor) (Casao *et al.* 2015) from mitochondria
- into the cytosol through disruption of mitochondrial membrane, inactivation of some
- biochemical pathways, enzyme dysfunction, disturbed axonemal protein
- 378 phosphorylation, increased membrane permeability and generation of spermicidal
- products, which have adverse effects on the spermatozoa functions. Similarly, the
- 380 spermatozoa motility is also reduced by ergonovine (Tash and Means 1982, Gallagher
- and Senger 1989), and the ergot alkaloids induce motility changes in bovine sperm cells
- by interacting with alpha-adrenergic receptors (Wang *et al.* 2009).
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384 385	Conclusion
386 387 388 389	The results of this study indicate that spermatozoa are a useful <i>in vitro</i> model for the toxicological evaluation of chemicals providing quantitative as well as qualitative data. In conclusion, the present study shows reduction of spermatozoa quality after <i>Viscum album quercus</i> addition <i>in vitro</i> .
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396 397	Conflict of Interest
398 399	Authors have no conflict of interest to declare.
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- 584

585 Table 1 – Concentrations of *Viscum album quercus* used in the study.

Group	Semen (µL)	Iscador Qu	Physiological	Concentration
		10 mg (µL)	solution (µL)	of Iscador Qu in
				samples
				(mg/mL)
QK – control	100	0	300	0
QA	100	300	0	10
QB	100	200	100	6.6
QC	100	100	200	3.3
QD	100	100	300	2.5
QE	100	100	400	2

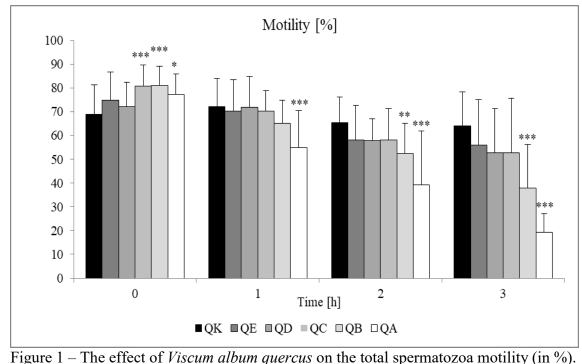




Figure 1 – The effect of *Viscum album quercus* on the total spermatozoa motility (in %) QK – 0; QE – 2; QD – 2.5; QC – 3.3; QB – 6.6; QA – 10 mg/mL of Iscador Qu. The level of significance was set at *** (p < 0.001), ** (p < 0.01) and * (p < 0.05).

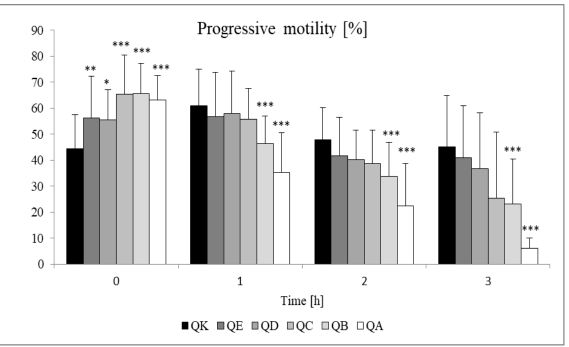


Figure 2 – The effect of Viscum album quercus on the progressive spermatozoa motility (in %). QK – 0; QE – 2; QD – 2.5; QC – 3.3; QB – 6.6; QA – 10 mg/mL of Iscador Qu. 593 The level of significance was set at *** (p < 0.001), ** (p < 0.01) and * (p < 0.05). 594

Table 2 – Spermatozoa distance average path (DAP; µm) in experimental groups and 596 time periods. 597

GroupMeanS.D.C.V.MinimumMaximumTime 0QK22.904.4319.3616.0232.75QA24.563.7915.4316.8231.60QB25.693.9215.2518.3532.61QC25.415.2120.5211.9733.70QD26.145.7622.0214.0538.53QE22.973.9917.3716.4031.49Time 1QK28.913.69QA18.05***2.2212.3014.1422.83QB20.24***2.2511.1416.4126.01QC22.92***2.3810.3818.7027.44QD26.232.429.2121.1530.61QE26.493.4412.9921.5432.76Time 2QK22.513.3414.8415.2931.10QA13.27***3.3625.340.0016.81QB17.26***2.4113.9712.6821.90QC17.45***1.8810.8013.8121.56QD19.02**2.7214.3014.4527.30QE20.552.8013.6013.6426.45Time 3	time periods	ume periods.							
$\begin{array}{c c c c c c c c c c c c c c c c c c c $	Group	Mean	S.D.	C.V.	Minimum	Maximum			
$\begin{array}{c c c c c c c c c c c c c c c c c c c $	Time 0								
$\begin{array}{c c c c c c c c c c c c c c c c c c c $	QK	22.90	4.43	19.36	16.02	32.75			
$\begin{array}{c c c c c c c c c c c c c c c c c c c $	QA	24.56	3.79	15.43	16.82	31.60			
$\begin{array}{c c c c c c c c c c c c c c c c c c c $	QB	25.69	3.92	15.25	18.35	32.61			
$\begin{array}{c c c c c c c c c c c c c c c c c c c $	QC	25.41	5.21	20.52	11.97	33.70			
Time 1QK28.91 3.69 12.75 21.19 36.69 QA 18.05^{***} 2.22 12.30 14.14 22.83 QB 20.24^{***} 2.25 11.14 16.41 26.01 QC 22.92^{***} 2.38 10.38 18.70 27.44 QD 26.23 2.42 9.21 21.15 30.61 QE 26.49 3.44 12.99 21.54 32.76 Time 2QK 22.51 3.34 14.84 15.29 31.10 QA 13.27^{***} 3.36 25.34 0.00 16.81 QB 17.26^{***} 2.41 13.97 12.68 21.90 QC 17.45^{***} 1.88 10.80 13.81 21.56 QD 19.02^{**} 2.72 14.30 14.45 27.30 QE 20.55 2.80 13.60 13.64 26.45	QD	26.14	5.76	22.02	14.05	38.53			
$\begin{array}{c c c c c c c c c c c c c c c c c c c $	QE	22.97	3.99	17.37	16.40	31.49			
$\begin{array}{c c c c c c c c c c c c c c c c c c c $			Tir	ne 1	•	•			
QB 20.24*** 2.25 11.14 16.41 26.01 QC 22.92*** 2.38 10.38 18.70 27.44 QD 26.23 2.42 9.21 21.15 30.61 QE 26.49 3.44 12.99 21.54 32.76 Time 2 QK 22.51 3.34 14.84 15.29 31.10 QA 13.27*** 3.36 25.34 0.00 16.81 QB 17.26*** 2.41 13.97 12.68 21.90 QC 17.45*** 1.88 10.80 13.81 21.56 QD 19.02** 2.72 14.30 14.45 27.30 QE 20.55 2.80 13.60 13.64 26.45	QK		3.69	12.75	21.19	36.69			
QC22.92***2.3810.3818.7027.44QD26.232.429.2121.1530.61QE26.493.4412.9921.5432.76Time 2QK22.513.3414.8415.2931.10QA13.27***3.3625.340.0016.81QB17.26***2.4113.9712.6821.90QC17.45***1.8810.8013.8121.56QD19.02**2.7214.3014.4527.30QE20.552.8013.6013.6426.45	QA	18.05***	2.22	12.30	14.14	22.83			
QC22.92***2.3810.3818.7027.44QD26.232.429.2121.1530.61QE26.493.4412.9921.5432.76Time 2QK22.513.3414.8415.2931.10QA13.27***3.3625.340.0016.81QB17.26***2.4113.9712.6821.90QC17.45***1.8810.8013.8121.56QD19.02**2.7214.3014.4527.30QE20.552.8013.6013.6426.45	QB	20.24***	2.25	11.14	16.41	26.01			
QD26.232.429.2121.1530.61QE26.493.4412.9921.5432.76Time 2QK22.513.3414.8415.2931.10QA13.27***3.3625.340.0016.81QB17.26***2.4113.9712.6821.90QC17.45***1.8810.8013.8121.56QD19.02**2.7214.3014.4527.30QE20.552.8013.6013.6426.45	QC	22.92***	2.38	10.38	18.70	27.44			
Time 2QK22.513.3414.8415.2931.10QA13.27***3.3625.340.0016.81QB17.26***2.4113.9712.6821.90QC17.45***1.8810.8013.8121.56QD19.02**2.7214.3014.4527.30QE20.552.8013.6013.6426.45	QD		2.42	9.21	21.15	30.61			
QK22.513.3414.8415.2931.10QA13.27***3.3625.340.0016.81QB17.26***2.4113.9712.6821.90QC17.45***1.8810.8013.8121.56QD19.02**2.7214.3014.4527.30QE20.552.8013.6013.6426.45	QE	26.49	3.44	12.99	21.54	32.76			
QA13.27***3.3625.340.0016.81QB17.26***2.4113.9712.6821.90QC17.45***1.8810.8013.8121.56QD19.02**2.7214.3014.4527.30QE20.552.8013.6013.6426.45			Tir	me 2	·	•			
QB17.26***2.4113.9712.6821.90QC17.45***1.8810.8013.8121.56QD19.02**2.7214.3014.4527.30QE20.552.8013.6013.6426.45	QK		3.34	14.84	15.29	31.10			
QB17.26***2.4113.9712.6821.90QC17.45***1.8810.8013.8121.56QD19.02**2.7214.3014.4527.30QE20.552.8013.6013.6426.45	QA	13.27***	3.36	25.34	0.00	16.81			
QC17.45***1.8810.8013.8121.56QD19.02**2.7214.3014.4527.30QE20.552.8013.6013.6426.45		17.26***	2.41	13.97	12.68	21.90			
QD19.02**2.7214.3014.4527.30QE20.552.8013.6013.6426.45		17.45***	1.88	10.80	13.81	21.56			
QE 20.55 2.80 13.60 13.64 26.45		19.02**	2.72	14.30	14.45	27.30			
			2.80	13.60	13.64	26.45			
			Tir	me 3					

QK	19.54	4.31	22.06	11.80	27.42
QA	9.40***	3.99	42.45	0.00	16.90
QB	14.17***	4.06	28.65	0.00	22.14
QC	16.70	4.42	26.48	8.95	22.17
QD	18.56	4.40	23.72	11.09	27.61
QE	18.44	3.18	17.25	12.06	22.85

Legend: S.D. – standard deviation; C.V. – coefficient of variation; *** (p < 0.001), ** (p < 0.01) (experimental group vs. control). QK – 0; QE – 2; QD – 2.5; QC – 3.3; QB –

6.6; QA – 10 mg/mL of Iscador Qu.

Table 3 – Spermatozoa distance curvilinear line (DCL; μ m) in experimental groups and time periods.

Crease		SD	C.V.	Minimum	Maximum			
Group	Mean	S.D.		Iviinimum	Maximum			
Time 0								
QK	47.84	7.80	16.30	35.09	67.59			
QA	52.25	6.81	13.03	36.55	65.73			
QB	52.76	6.52	12.35	38.31	65.55			
QC	50.01	7.19	14.37	34.42	63.38			
QD	56.03**	14.35	25.61	35.85	92.81			
QE	47.66	6.48	13.59	33.89	62.09			
		Tii	ne 1					
QK	56.01	7.39	13.19	44.27	77.16			
QA	41.19***	4.74	11.52	33.71	53.96			
QB	43.79***	4.70	10.74	36.12	53.41			
QC	45.98***	4.07	8.86	40.28	54.68			
QD	52.54	5.33	10.14	40.52	61.28			
QE	53.61	7.27	13.56	41.29	66.05			
		Tii	me 2					
QK	46.87	7.61	16.24	33.81	68.31			
QA	31.38***	9.01	28.71	0.00	44.65			
QB	40.40***	4.93	12.21	31.29	50.00			
QC	38.71***	5.07	13.09	29.61	50.94			
QD	41.62*	5.56	13.35	28.80	59.57			
QE	45.42	5.46	12.03	31.92	58.97			
		Tiı	ne 3					
QK	42.31	9.20	21.76	27.39	60.82			
QA	20.69***	9.53	46.07	0.00	37.80			
QB	34.22**	10.26	29.97	0.00	52.86			
QC	37.83	9.40	24.85	21.49	49.89			
QD	43.07	8.81	20.45	25.50	58.38			
QE	41.58	7.61	18.31	25.81	54.31			
	11.50							

Legend: S.D. – standard deviation; C.V. – coefficient of variation; *** (p < 0.001), ** (p < 0.01) and * (p < 0.05) (experimental group vs. control). QK – 0; QE – 2; QD – 2.5;

QC - 3.3; QB - 6.6; QA - 10 mg/mL of Iscador Qu.

Group	Mean	S.D.	C.V.	Minimum	Maximum			
Time 0								
QK	17.63	3.87	21.97	10.02	26.30			
QA	16.62	3.16	19.00	12.07	23.31			
QB	18.15	3.57	15.25	11.38	24.51			
QC	19.18	4.75	20.52	8.00	26.47			
QD	17.84	3.59	20.11	9.68	24.21			
QE	16.91	3.24	19.17	11.06	23.93			
		Tir	ne 1					
QK	22.28	3.37	15.13	16.09	29.73			
QA	12.84***	1.52	11.88	9.67	16.74			
QB	15.33***	1.81	11.78	12.13	19.33			
QC	17.72**	1.94	10.92	13.87	22.38			
QD	19.85	1.74	8.75	15.93	23.06			
QE	19.88	2.18	10.95	16.33	23.88			
		Tir	ne 2					
QK	17.38	2.66	15.29	12.61	24.09			
QA	9.60***	2.48	25.82	0.00	13.09			
QB	12.73***	2.04	16.05	8.87	17.83			
QC	13.53***	1.57	11.62	10.57	17.16			
QD	14.82**	2.58	17.40	11.06	22.86			
QE	15.58	1.95	12.52	10.86	19.14			
		Tir	ne 3					
QK	14.86	3.12	20.97	8.69	20.65			
QA	6.72***	2.81	41.89	0.00	10.76			
QB	10.08***	2.75	27.26	0.00	15.48			
QC	12.12**	3.16	26.10	5.58	16.52			
QD	13.31	3.37	25.33	7.81	22.37			
QE	13.60	2.12	15.59	9.66	17.87			

Table 4 – Spermatozoa distance straight line (DSL; µm) in experimental groups and 608 time periods. 609

Legend: S.D. – standard deviation; C.V. – coefficient of variation; *** (p < 0.001) (experimental group vs. control). QK – 0; QE – 2; QD – 2.5; QC – 3.3; QB – 611

6.6; QA – 10 mg/mL of Iscador Qu. 612

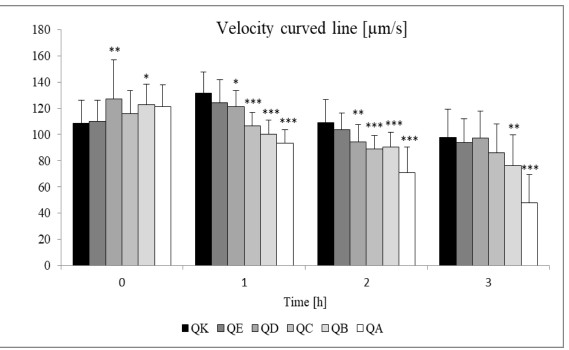


Figure 3 – The effect of *Viscum album quercus* on the spermatozoa velocity curved line

615 $(\mu m/s)$. QK - 0; QE - 2; QD - 2.5; QC - 3.3; QB - 6.6; QA - 10 mg/mL of Iscador Qu.

- 616 The level of significance was set at *** (p < 0.001), ** (p < 0.01) and * (p < 0.05).
- 617

 $\begin{array}{ll} \text{618} & \text{Table 5-Spermatozoa velocity average path (VAP; <math>\mu$ m/s) in experimental groups and time periods. \end{array}

Crease		C D	CV	Minimum	Mariana			
Group	Mean	S.D.	C.V.	Minimum	Maximum			
Time 0								
QK	52.47	10.24	19.51	33.87	73.56			
QA	57.45	9.21	16.03	39.74	73.75			
QB	60.27	9.37	15.55	43.24	79.03			
QC	59.22	12.53	21.16	27.74	76.57			
QD	60.01	12.74	21.22	30.63	82.13			
QE	53.54	9.79	18.28	37.27	73.56			
		Tiı	ne 1					
QK	68.12	8.16	11.99	50.96	86.26			
QA	41.16***	5.22	12.68	31.48	51.78			
QB	46.62***	5.25	11.26	36.14	59.89			
QC	53.53***	6.09	11.37	43.38	64.99			
QD	60.81***	5.87	9.65	48.00	73.53			
QE	61.79**	8.34	13.50	50.35	78.56			
		Tiı	me 2					
QK	52.56	7.84	14.91	35.50	72.26			
QA	30.32***	7.56	24.93	0.00	38.99			
QB	38.92***	5.39	13.84	29.51	49.88			
QC	40.41***	4.07	10.07	32.51	48.94			
QD	43.58***	6.71	15.41	31.77	60.93			
QE	47.11 [*]	6.80	14.43	33.09	64.36			
	· .	Tiı	me 3	•	•			

QK	45.30	10.31	22.75	27.50	64.75
QA	22.61***	10.05	44.43	0.00	40.14
QB	31.86***	9.39	29.48	0.00	49.84
QC	38.17	10.45	27.38	19.32	52.12
QD	42.18	10.34	24.51	25.95	61.73
QE	41.81	7.56	18.08	28.83	52.11

620 Legend: S.D. – standard deviation; C.V. – coefficient of variation; *** (p < 0.001), **

621 (p < 0.01) and * (p < 0.05) (experimental group vs. control). QK - 0; QE - 2; QD - 2.5;

 $622 \qquad QC - 3.3; \, QB - 6.6; \, QA - 10 \text{ mg/mL of Iscador Qu}.$

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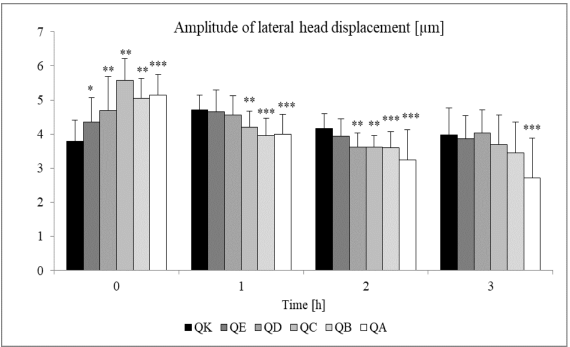
Table 6 – Spermatozoa velocity straight line (VSL; μ m/s) in experimental groups and time periods.

Group	Mean	S.D.	C.V.	Minimum	Maximum			
Time 0								
QK	40.53	8.81	21.73	21.06	60.28			
QA	39.07	7.71	19.72	27.11	54.48			
QB	42.93	8.68	20.22	26.91	59.89			
QC	44.94	11.41	25.39	18.85	60.35			
QD	41.47	9.28	22.36	21.05	58.10			
QE	39.72	7.99	20.12	25.20	56.26			
		Tir	ne 1					
QK	52.75	7.60	14.41	38.78	70.71			
QA	29.35***	3.70	12.60	21.52	37.00			
QB	35.44***	4.17	11.78	27.66	44.88			
QC	41.50***	5.04	12.14	31.08	52.56			
QD	46.26***	4.27	9.24	36.38	55.44			
QE	46.64***	5.25	11.25	39.21	57.29			
		Tir	me 2					
QK	40.72	6.28	15.41	29.37	56.76			
QA	21.99***	5.56	25.60	0.00	30.47			
QB	28.76***	4.53	15.76	20.79	40.55			
QC	31.40***	3.55	11.31	24.63	39.57			
QD	34.09***	6.39	18.75	23.67	51.32			
QE	35.68**	4.51	12.63	26.71	44.68			
		Tir	ne 3					
QK	34.52	7.51	21.77	20.31	49.01			
QA	16.49***	7.56	45.86	0.00	31.82			
QB	22.65***	6.40	28.23	0.00	34.89			
QC	27.81**	7.45	26.80	12.09	38.72			
QD	30.22	7.86	23.01	18.52	50.15			
QE	30.80	4.97	16.14	21.21	40.57			

626 Legend: S.D. – standard deviation; C.V. – coefficient of variation; *** (p < 0.001), **

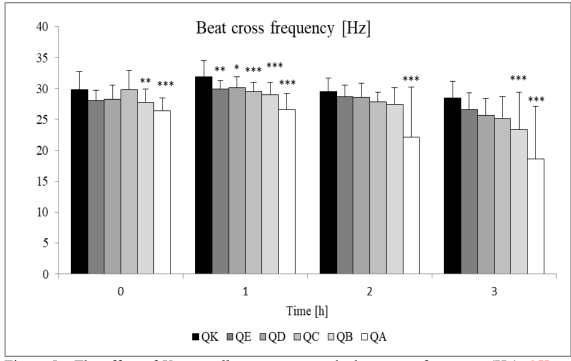
627 (p < 0.01) (experimental group vs. control). QK - 0; QE - 2; QD - 2.5; QC - 3.3; QB - 2

628 6.6; QA – 10 mg/mL of Iscador Qu.



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Figure 4 – The effect of *Viscum album quercus* on the amplitude of lateral head displacement (μ m). QK – 0; QE – 2; QD – 2.5; QC – 3.3; QB – 6.6; QA – 10 mg/mL of Iscador Qu. The level of significance was set at *** (p < 0.001), ** (p < 0.01) and * (p < 0.05).



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Figure 5 – The effect of *Viscum album quercus* on the beat cross frequency (Hz). QK – 0; QE – 2; QD – 2.5; QC – 3.3; QB – 6.6; QA – 10 mg/mL of Iscador Qu. The level of significance was set at *** (p < 0.001), ** (p < 0.01) and * (p < 0.05).

640 Table 7 – Spermatozoa straightness (STR) in experimental groups and time periods.

Group	Mean	S.D.	C.V.	Minimum	Maximum				
	Time 0								
QK	0.77	0.06	7.88	0.61	0.89				
QA	0.67***	0.06	9.28	0.57	0.77				
QB	0.70**	0.06	9.17	0.59	0.80				
QC	0.75	0.06	8.50	0.63	0.87				
QD	0.69***	0.11	15.98	0.45	0.84				
QE	0.73	0.04	5.16	0.67	0.82				
		Tir	ne 1						
QK	0.77	0.04	5.57	0.62	0.86				
QA	0.71***	0.05	6.91	0.58	0.79				
QB	0.76	0.04	5.45	0.69	0.86				
QC	0.77	0.03	3.59	0.71	0.81				
QD	0.76	0.05	6.26	0.64	0.85				
QE	0.75	0.03	4.51	0.68	0.82				
		Tir	ne 2						
QK	0.77	0.04	5.42	0.62	0.85				
QA	0.70***	0.14	19.46	0.00	0.89				
QB	0.73	0.05	6.56	0.65	0.84				
QC	0.77	0.02	3.10	0.72	0.82				
QD	0.77	0.05	5.89	0.69	0.85				
QE	0.75	0.04	5.50	0.66	0.84				
		Tir	ne 3						
QK	0.76	0.05	6.45	0.66	0.91				
QA	0.64 ^C	0.26	40.91	0.00	31.82				
QB	0.69	0.14	20.23	0.00	0.92				
QC	0.73	0.05	6.66	0.61	0.80				
QD	0.71	0.06	8.58	0.61	0.81				
QE	0.74	0.04	5.89	0.66	0.81				

Legend: S.D. – standard deviation; C.V. – coefficient of variation; *** (p < 0.001), ** (p < 0.01) (experimental group vs. control). QK – 0; QE – 2; QD – 2.5; QC – 3.3; QB –

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Table 8 – Spermatozoa linearity (LIN) in experimental groups and time periods.

Tuble 0 Bpe	Table 6 Spermatozoa miearry (Ent) in experimental groups and time periods.							
Group	Mean	S.D.	C.V.	Minimum	Maximum			
		Tir	ne 0					
QK	0.37	0.05	13.31	0.23	0.45			
QA	0.32**	0.03	10.81	0.26	0.37			
QB	0.34	0.05	14.89	0.27	0.44			
QC	0.38	0.07	17.39	0.24	0.48			
QD	0.33	0.09	25.70	0.18	0.47			
QE	0.35	0.04	12.00	0.27	0.42			
		Tir	ne 1					
QK	0.40	0.04	10.28	0.29	0.50			
QA	0.31***	0.03	9.77	0.24	0.36			
QB	0.35***	0.03	8.88	0.29	0.46			

 $[\]begin{array}{ll} \text{642} & (p < 0.01) \text{ (experimental group vs. co} \\ \text{643} & \text{6.6; } QA - 10 \text{ mg/mL of Iscador Qu.} \end{array}$

QC	0.38	0.03	7.33	0.30	0.44
QD	0.38	0.04	9.56	0.30	0.47
QE	0.37*	0.02	6.42	0.32	0.41
		Tir	ne 2		
QK	0.37	0.03	8.10	0.27	0.42
QA	0.31***	0.12	38.07	0.00	0.86
QB	0.31***	0.03	8.75	0.26	0.38
QC	0.35	0.03	7.42	0.30	0.40
QD	0.35	0.04	10.76	0.28	0.42
QE	0.34	0.02	6.99	0.29	0.39
		Tir	ne 3		
QK	0.35	0.03	9.49	0.30	0.44
QA	0.32	0.18	56.85	0.00	0.74
QB	0.29	0.11	38.09	0.00	0.81
QC	0.32	0.04	12.17	0.24	0.40
QD	0.30	0.04	12.96	0.25	0.39
QE	0.33	0.03	10.36	0.27	0.40

Legend: S.D. – standard deviation; C.V – coefficient of variation; *** (p < 0.001), ** (p < 0.01) and * (p < 0.05) (experimental group vs. control). QK – 0; QE – 2; QD – 2.5; QC – 3.3; QB – 6.6; QA – 10 mg/mL of Iscador Qu.

Table 9 – Spermatozoa wobble (WOB) in experimental groups and time periods.

Group	Mean	S.D.	C.V.	Minimum	Maximum			
Time 0								
QK	0.48	0.04	8.73	0.38	0.55			
QA	0.47	0.02	3.71	0.43	0.50			
QB	0.48	0.03	7.21	0.44	0.55			
QC	0.50	0.05	9.99	0.36	0.57			
QD	0.47	0.06	11.90	0.39	0.56			
QE	0.48	0.04	7.73	0.40	0.53			
Time 1								
QK	0.51	0.03	6.17	0.44	0.60			
QA	0.44***	0.03	6.29	0.38	0.49			
QB	0.46***	0.02	4.89	0.42	0.53			
QC	0.50	0.03	5.35	0.43	0.56			
QD	0.50	0.02	4.84	0.46	0.57			
QE	0.49*	0.02	4.81	0.43	0.53			
Time 2								
QK	0.48	0.02	4.70	0.43	0.52			
QA	0.42**	0.13	30.50	0.00	0.86			
QB	0.42**	0.02	5.45	0.39	0.47			
QC	0.45	0.03	6.08	0.30	0.40			
QD	0.46	0.03	6.83	0.39	0.51			
QE	0.45	0.02	4.99	0.40	0.50			
Time 3								
QK	0.46	0.03	6.50	0.39	0.57			

QA	0.42	0.20	46.81	0.00	0.80
QB	0.41	0.12	28.25	0.00	0.87
QC	0.44	0.03	7.70	0.34	0.50
QD	0.43	0.03	6.84	0.38	0.49
QE	0.44	0.03	6.16	0.38	0.52

Legend: S.D. – standard deviation; C.V. – coefficient of variation; *** (p < 0.001), **

(p < 0.01) and * (p < 0.05) (experimental group vs. control). QK - 0; QE - 2; QD - 2.5;

 $\label{eq:QC-3.3} \textbf{QC-3.3}; \textbf{QB-6.6}; \textbf{QA-10} \text{ mg/mL of Iscador Qu}.$

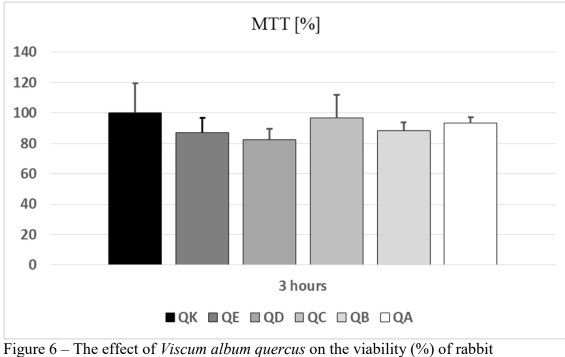


Figure 6 – The effect of *Viscum album quercus* on the viability (%) of rabbit
spermatozoa after 3 hours of incubation. QK – 0; QE – 2; QD – 2.5; QC – 3.3; QB –
6.6; QA – 10 mg/mL of Iscador Qu.

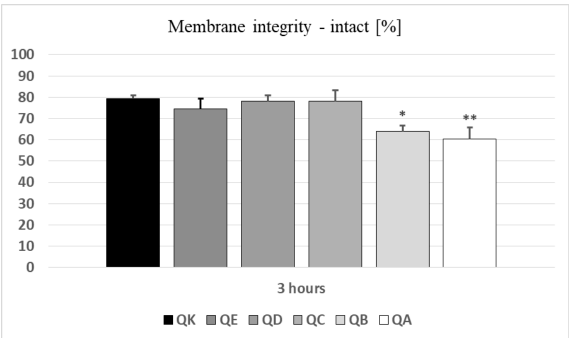


Figure 7 – The effect of *Viscum album quercus* on the membrane integrity (%) of rabbit spermatozoa assessed after 3 hours of incubation. QK - 0; QE - 2; QD - 2.5; QC - 3.3; QB - 6.6; QA - 10 mg/mL of Iscador Qu. The level of significance was set at ** (p < (0.01) and * (p < 0.05).

