RESUMO: O receptor de ecdisona, naturalmente ativado por hormônios esteroidais, é uma proteína-chave nos processos de muda e reprodução de insetos. A ativação artificial desse receptor por meio de pesticidas específicos induz um processo de ecdise anômala, levando o inseto à morte por dessecção e inanição. Neste trabalho, foi estabelecido um protocolo para a triagem de moléculas agonistas em relação ao receptor de ecdisona nas linhagens celulares responsivas S2 (Diptera) e Sf9 (Lepidoptera), transfectadas com o plasmídeo repórter ere.b.act.luc. Para tanto, curvas de dose-resposta foram estabelecidas com o ecdisteroide 20-hidroxiecdisona, o fitoecdisteroide ponasterona-A e tebufenozida, um pesticida pertencente à classe das diacilhidrazinas. Em ambas linhagens celulares, os valores médios de concentração efetiva para indução gênica (EC50) ponasterona-A foram menores, significando que este é o agonista mais potente. Em células Sf9, a tebufenozida apresentou EC50 menor que a 20-hidroxiecdisona, indicando uma alta atividade agonista e especificidade deste inseticida a lepidópteros. O protocolo estabelecido neste trabalho pode ser utilizado para uma rápida triagem e busca racional de pesticidas de alvo bioquímico específico.

PALAVRAS-CHAVE: plasmídeo; tebufenozida; 20-hidroxiecdisona; ponasterona-A.
The ecdysone receptor is a protein that belongs to the family of Nuclear Receptors that are involved in the processes of insects molting, metamorphosis and reproduction. Knowledge of the function and structure of ecdysteroids and their receptor allowed the development of non-steroidal compounds with agonist activity, mainly represented by the insecticide class of diacylhydrazines (WING et al., 1988). Although these compounds are used as pesticides, especially for controlling harmful lepidopterans and some mites in crops, there are few diacylhydrazines registered for commercial use.

In addition to diacylhydrazines, different chemical groups of non-steroidal compounds, such as oligobestenes (MENG et al., 2001), alkaoids (DINAN et al., 2001), α-acilaminoketones (TICE et al., 2003), acylaminoketones, tetrahydroquinolines (SOIN et al., 2010) or unclassified molecules (HARADA et al., 2011; HU, et al., 2018) exhibit agonist or antagonist activity at insect ecdysone receptor. Thus, ecdysone receptor is a poorly explored, but interesting target for continuous search of a broad range of safer and selective molecules.

Cell lines secrete all necessary components for activating and transactivating the ecdysone receptor (ZOTTI et al., 2013). Thus, cell-based bioassays associated with screening of chemical libraries may be used for the prospection of selective insecticides. Therefore, the objective of this study was to propose a rapid and specific protocol for screening molecules based on the activation of ecdysone receptor in cell lines. For this, based on dose response curves, we estimated activity patterns of different agonist molecules towards dipteran (S2) and lepidopteran (Sf9) cell lines.

The Sf9 cell line, derived from embryonic cells of Spodoptera frugiperda (J. E. Smith, 1797) (Lepidoptera: Noctuidae), was maintained at 27°C in SF900 ™ culture medium (Gibco®). The S2 cell line, derived from embryonic cells of Drosophila melanogaster (Meigen, 1830) (Diptera: Drosophilidae), was maintained at 27°C in InsectXpress ™ (Lonza®) culture medium.

Cells were transiently transfected with the plasmid ere.bact. luc., a reporter constructed specifically for the detection of ecdysone receptor activity. The plasmid was constructed with seven copies of the Drosophila ecdysone response element hsp27 (ere.), the basal actin promoter (b. act.), the firefly luciferase enzyme as reporter gene (luc.) and a termination signal (SWEVERS et al., 2004). Thus, molecules were assayed for their ability to promote ecdysone receptor activation, which is measurable according to the light emitted by the luciferase enzyme in transfected cells with the reporter plasmid.

For preparing the transfection medium for S2 cells, 497 μL of culture medium was added to an Eppendorf tube followed by 3 μL of transfection reagent Escort IV™ (Sigma-Aldrich®) and 100 ng reporter plasmid. The reaction was incubated at room temperature (RT) for 30 minutes. For Sf9 cells, transfection medium was prepared with 496 μL culture medium, 4 μL of transfection reagent and 300 ng reporter plasmid were mixed into an Eppendorf tube, and also incubated at RT for 30 minutes.

Prior to the transfection itself, the 24-well plates were filled with 5 × 10³ and 3 × 10³ cells of S2 and Sf9, respectively. Cell counting was performed with a Neubauer Chamber and Trypan Blue (Sigma-Aldrich) as a cell viability reagent. A period of 1 hour was expected for cells to adhere to the bottom of the plate. Subsequently, the culture medium was withdrawn and replaced by the transfection medium. Both cell lines were incubated for 5 hours with the transfection medium, previously described. After this time, the transfection medium was withdrawn and replaced with fresh culture medium.

To establish this screening system, in both cell lines we estimated the EC₅₀ of two steroidal agonists, 20-hydroxyecdysone and ponasterone-A (Sigma-Aldrich), and one non-steroidal compound of the diacylhydrazine group, the tebufenozide (Sigma-Aldrich) (Fig. 1). The carrier solvent used was ethanol, in which serial dilutions of the molecules were performed to obtain a robust dose response curve and as negative control, we used pure ethanol. For each concentration, 1 and 1.5 μL of the diluted molecules were added to the S2 and Sf9 cell plates, respectively.

Luminescence was measured 24 hours after the molecules were added to the cells. Thus, cells were resuspended and 100 μL of them were transferred to wells of a white 96-well plate. Thereafter, 100 μL of the luciferase substrate (Steady-Glo Luciferase Assay System Kit) was also added to the 96 well plate. The model of the luminometer used was VitorTM X5 and readings started 5 minutes after the addition of luciferase substrate into the 96-well plates. Each treatment was performed in quadruplicate, and the experiments were performed 4 times. EC₅₀ values (mean effective concentration values for induction of Luciferase enzyme) were calculated with 95% reliability and transformed into LogX using GraphPad Prism v.4 software (GraphPad Software Inc. La Jolla, Ca) and the accuracy of the data evaluated based on R² values.

The EC₅₀ (LogX) of tebufenozide in S2 was -3.4, whereas in Sf9 it was -5.1, i.e., the activity of this molecule in Lepidoptera cells was almost 100 times higher than in Diptera (Fig. 1 and Table 1). The concentration of tebufenozide required to activate the ecdysone receptor of S2 to an 80% equivalent observed in Hemiptera (TOHIDI-ESFAHANI et al., 2011).

Thus, ecdysone receptor is a poorly explored, but interesting target for continuous search of a broad range of safer and selective molecules.
Compared activity of agonist molecules towards ecdysone receptor in insect cell-based screening system

Giles, 1926 (Diptera: Culicidae) found that some diacylhydrazines, such as methoxifenozide and KU-106, cause similar symptoms to those seen in lepidopterans, such as earlier ecdysis and malformations (BECKAGE et al., 2004; MOROU et al., 2013). Our results show that tebufenozide can activate ecdysone receptor of S2 cells. It points out that the ecdysone receptor of Diptera presents a flexibility, that allows binding of tebufenozide, which makes the search for novel ecdysone receptor agonist pesticides available.

The 20-hydroxyecdysone is the most representative ecdysteroid of the insects; however, in this study, we show that phytoecdysteroids may have a higher activation power (Fig. 1). Phytoecdysteroids are found in approximately 6% of the plant species and more than 500 ecdysteroids, including from fungi, have been identified in the nature (ECDYBASE, 2019).

The ability of plants to produce active phytoecdysteroids may be an evolutionary adaptation, and the activity of these compounds may vary among insect orders (DINAN et al., 2001). In this paper, we show the high activity of Ponasterone-A as an ecdysone receptor agonist (Fig. 1). This phytoecdysteroid can cause similar symptoms to insecticides, such as diacylhydrazines, when absorbed in the insect mesenteric, representing a significant barrier against herbivory (BOWERS, 2012; CHAUBEY, 2018).

Phytoecdysteroids such as castasterone, may have a higher affinity for dipteran rather than the lepidopteran ecdysone receptor, unlike diacylhydrazines (ZOTTI et al., 2013). Furthermore, phytoecdysteroids are also capable of antagonizing the 20-hydroxyecdysone hormone, as already documented in Phormia terranovae Robineau-Desvoidy, 1830 (Diptera: Calliphoridae) and Thrips tabaci Lindeman, 1889 (Thysanoptera: Thripidae) (HETRU et al., 1986; RICHTER; KOOLMAN, 1991; MIYAJI et al., 2014).

In the present study, we established a protocol for the search of ecdysone receptor agonist compounds by verifying different activation patterns of molecules from different origins.

Bars represent means ± standard errors. MW: molecular weight; MF: molecular formula.

**Figure 1.** Dose-response sigmoid curves referring to the activity of the ecdysteroid 20-hydroxyecdysone, the phytoecdysteroid ponasterone-A, and the non-steroidal agonist tebufenozide in lepidopteran (Sf9) and dipteran (S2) cells.

**Table 1.** Agonistic activity of 20-hydroxyecdysone, ponasterone-A and tebufenozide in S2 and Sf9 cell lines.

|          | S2       |          | S9       |          |
|----------|----------|----------|----------|----------|
|          | -logEC_{50} | 95%IC    | R^2      | -logEC_{50} | 95%IC    | R^2      |
| 20H      | 4.90     | 4.98 - 4.78 | 0.964    | 4.21     | 4.28 - 4.12 | 0.889    |
| PonA     | 5.57     | 5.85 - 5.30 | 0.920    | 5.27     | 5.31 - 5.24 | 0.984    |
| Tebu     | 3.4      | 3.41 - 3.38 | 0.999    | 5.01     | 5.03 - 4.99 | 0.994    |

Data are given as median response values together with the 95% confidence interval (both in -logEC_{50}). R^2 represents the accuracy of data fitting to the sigmoid curve.
(Fig. 1). Besides that, another possibility is the search for antagonistic compounds to the ecdysone receptor, expanding the possibilities of discovery (HU et al., 2018). For example, compounds derived from Cyperaceae, such as resveratrol, are capable of competing for the active site with ponasterone-A and 20-hydroxyecdysone, generating antagonistic activity in dipteran cells (MENG et al., 2016).

Ecdysone receptor cell-based assays are useful to find the pattern of activation of compounds or to identify new bioactive molecules. Once the workflow is established, this kind of assay allows testing several molecules with a relatively low cost of time, space and financial resources, especially because there is no need for insect rearing. Even though, if research focuses on finding new bioactive molecules, subsequent to cell-based assays, the ideal is testing promising compounds on whole organisms (SWEVERS; SMAGGHE, 2016). Several molecules capable of activating the ecdysone receptor of Si2 cell line, derived from Spodoptera littoralis (Boisduval, 1833) and 20-hydroxyecdysone, generating antagonistic activity in vivo (SOIN et al., 2010). In addition, virtual substituted diacylhydrazine radicals may vary in toxicity in vivo, unlike predicted by virtual screening (DENG et al., 2016).

The insect cell-based screening system, highly specific for ecdysone receptor, established in this work is an agile, sustainable and relatively inexpensive methodology for testing a wide range of different molecules. The 20-hydroxyecdysone, Ponasterone-A and tebufenozide can be used as a standard and a comparative for other compounds.

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