Evaluation of Receptor Affinity, Analgesic Activity and Cytotoxicity of a Hybrid Peptide, AWL3020

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Abstract
In the present contribution we report design, synthesis and evaluation of receptor affinity, analgesic activity and cytotoxicity of a hybrid peptide, AWL3020. The peptide includes two pharmacophores, one of δ-opioid receptor (δOR) agonists and one of neurokinin-1 receptor (NK1R) antagonists. The design was motivated by the desire to obtain a compound with strong analgesic action and potential additional antiproliferative action. The compound displays high δOR affinity (IC50 = 29.5 nM). On the other hand, it has only poor affinity for the NK1R (IC50 = 70.28 μM). The substance shows good analgesic action which is however weaker than that of morphine. Regarding the effect on proliferation, the compound exhibits no pro-proliferative action in the assayed range. In higher concentrations, it has also cytotoxic activity. This effect is however not selective. The strongest effect of AWL3020 was found for melanoma MeW164 cell line (EC50 = 46.27 μM in reduction of cell numbers after a few days of incubation; EC50 = 37.78 μM in MTT assay).

Keywords Cancer pain · Multitarget drugs · Neurokinin-1 receptor · Delta opioid receptor

Introduction
Various types of cancers are among the most important problems of modern medicine. Cancer is often difficult to treat and its progression is frequently accompanied by pain (Portenoy and Ahmed 2018). In particular, pain is most problematic in advanced stages of the disease. Pain seems to be a key factor contributing to the lowering of patients’ quality of lives.

Current treatment of cancer pain relies mostly on opioids like morphine, fentanyl, buprenorphine, tramadol and others (Money and Garber 2018). Many ways of administrations are available for the opioids including oral, transdermal, transmucosal or via enteral tubes. Unfortunately, these therapies are far from ideal. In many cases, they lack enough efficacy in relieving pain and their prolonged use causes tolerance. The opioids can also have unpleasant or even life-threatening side-effects (Benyamin et al. 2008), which include: nausea, vomiting, constipations and respiratory depression. Recently, there are also some reports that opioids can stimulate growth of cancer cells and induce progression of cancers (Lazarczyk et al. 2010; Lennon et al. 2012; Nguyen et al. 2014; Bimonte et al. 2015; Aich et al. 2016). Other papers bring contradicting
conclusions (Chen et al. 2017). The matter is subject of ongoing research summarized in several recent reviews (Zhang et al. 2018; Brinkman et al. 2018; Chen et al. 2019a; Tuerxun and Cui 2019).

Nevertheless, an ideal antipain drug for the treatment of cancer pain should be a strong analgesic, without significant side-effects and without stimulatory effects on the development of cancer. Optimally it could also have some anticancer activity on its own. In this way, it could positively cooperate with the main therapy. With these assumptions in mind, we have endeavoured to find analgesic peptides with auxiliary anticancer properties. To this aim, we have designed a hybrid peptide (named AWL3020, Fig. 1) consisting of two pharmacophores, one of δ-opioid receptor (δOR) agonists and one of neurokinin-1 receptor (NK1R) antagonists. The opioid sequence (Tyr-D-Ala-Phe-His-Leu-Leu-Asp…) was taken from a well-known δOR agonist, [D-Ala², Leu⁶]-deltorphin (Janecka et al. 2004). The neurokinin-1 antagonist sequence (…D-Phe-Phe-D-Phe-Leu-Leu-NH₂) was that of a peptidic NK1R antagonist, spantide I (Gether et al. 1993), with two conservative exchanges of D-Trp for D-Phe. The design of AWL3020 was based on the following rationale. First, δOR agonists are strong analgesic agents (Gavériaux-Ruff and Kieffer 2011). Second, hybridization of opioid agonist and NK1R antagonist pharmacophores produces analgesic molecules with reduced tolerance and side-effects (Largent-Milnes et al. 2010). Third, NK1R antagonists have been shown to have anticancer activity in vitro and in vivo (Harford-Wright et al. 2013; Garnier et al. 2015; Muñoz and Coveñas 2016).

In this paper, we report the pharmacological effects of AWL3020 in several in vitro and in vivo experiments that were aimed at validating the design rationale. First, we assayed AWL3020 for receptor affinities. Having found high affinity for the δOR, we performed molecular modelling of the compound bound to this receptor and assessed in vivo its analgesic activity. Finally, we comprehensively assessed the cytotoxicity of the compound against several cancer and normal cell lines.

Materials and Methods

AWL 3020 Synthesis

Compound AWL3020 was synthesized by the use of a modified method described previously (Laskowska et al. 2017). The synthesis was performed on solid support (SPPS), following a typical protocol, on the Rink amide resin and with HOBT/TBTU coupling reagents. The cleavage of peptide from the resin was conducted by using trifluoroacetic acid. Its mass was confirmed by mass spectrometry (C₈₂H₁₀₉N₁₅O₁₅, MW: 1544.834 u, monoisotopic mass 1543.82 u, ESI–MS ion found m/z [M+2H]²⁺ 772.90, calculated [M+2H]²⁺ 772.82). The purity of the preparation was found to be greater than 98% by using HPLC with UV/VIS detector (tᵣ = 11.7 min, linear gradient 3–97% of phase B in 20 min, phase A: 0.05% aq. FA, phase B: ACN + 0.05% FA, total flow rate: 1.2 ml/min). The chromatograms and the mass-spectrum are given in the Supplementary Materials in Figures ESM-CHEM-1 to ESM-CHEM-3.

Receptor Binding Assays

The binding affinity of compound AWL3020 for μOR, δOR and NK1R was determined in competitive radioligand binding assays using rat brain homogenates. The membrane fractions containing the receptors were obtained from rats euthanized by decapitation after which an immediate brain removal, separation of cerebellum and placement of the remaining brain in an ice-cold 50 nM Tris–HCl (pH 7.4) was conducted. Brain tissue was subject to homogenization with up-down strokes in a hand glass homogenizer and suspended in 20 Tris–HCl volumes. A
The mean IC50 values were determined with SEMs (GraphPad Prism version 5.0, San Diego CA). Displacement curves were drawn and measured in a scintillation counter MicroBeta LS, Trilux (PerkinElmer, USA) was added to each well. Radioactivity was measured in the presence of either 10 μM naloxone (opioid receptors) or pre-soaked for 3 h (in the case of NK1R) so as to minimize the extent of non-specific binding. Filter discs were placed separately in 24-well plates soing. Filter discs were placed separately in 24-well plates containing the peptide and in the receptor) were set as expected at 4RWD (Fenalti et al. 2015)) by using AutoDock 4.2 (Morris et al. 2009). The protonation states of the residues (in the peptide and in the receptor) were set as expected at physiological pH. Other preparatory steps were performed in AutoDock Tools 4 and AutoGrid 4 (Morris et al. 2009). The docking box was set around the ligand found in the structure (H-Dmt-Tic-Phe-Phe-NH2), but extended so that it encompassed the cavity up to the receptor outlet and slightly further.

Molecular Modelling

AWL3020 was docked to the δOR (PDB accession code: 4RWD (Fenalti et al. 2015)) by using AutoDock 4.2 (Morris et al. 2009). The protonation states of the residues (in the peptide and in the receptor) were set as expected at physiological pH. Other preparatory steps were performed in AutoDock Tools 4 and AutoGrid 4 (Morris et al. 2009). The docking box was set around the ligand found in the structure (H-Dmt-Tic-Phe-Phe-NH2), but extended so that it encompassed the cavity up to the receptor outlet and slightly further.

Animals

The experiments were performed in duplicate.

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The experimental design encompassed the cavity up to the receptor outlet, the cavity extended up to the cleft between the receptor outlet and the adjacent receptor side, but it did not encompass the cavity up to the receptor outlet and slightly further.

The rats (of 200–250 g weight, 10 animals for each test group and for control group) were prepared for the i.t. administration by a method described by Yaksh and Rudy (1976). The animals were anaesthetised with isoflurane (Forane, USP, Baxter) and placed in a stereotaxic device. Into their intrathecal space implanted were silicone tubing catheters (ID = 0.30 mm and OD = 0.64 mm, length = 12 cm, Silastic Laboratory Tubing, Dow Corning Corporation) with the tip located at L1 spinal level. At about 2/3 of its length it contains a silicone limiter in a form of a ball which prevents undesired displacements of the device. After the surgery and waking up from anaesthesia, the rats were given 2–3 days for recovery. They were housed in individual cages in a temperature- and light-controlled environment (12 h/12 h day/night cycle), with food and water ad libitum. Rats that suffered with plegia after the catheterisation were excluded from further experimental procedures.

All housing and experiments were conducted in accordance with the Polish Act of 21 January 2005 on Experiments on Animals (Journal of Laws No 33 of the Republic of Poland, item 289 as amended). All animal experiments were approved by the IV Local Ethics Committee for Experiments on Animals in Warsaw, Poland (permissions no.: 46/2013). All efforts were made to minimize the number of animals used and their suffering.

Tail-Flick Test After Intrathecal Drug Administration

The analgesic activity of compound AWL3020 was measured in vivo in adult male Wistar Rats by a tail-flick test after intrathecal (i.t.) administration.

The compounds under consideration (AWL3020 and morphine) were dissolved in a small amount of DMSO and then the saline was added so to obtain a solution with DMSO concentration not greater than 10%. They were administered via the above mentioned catheters in a total volume not exceeding 10 μl. Control animals received saline solution (Polfa). For the sake of comparison with a ‘gold standard’ treatment, another control group received a solution of morphine. Both the test and the control groups consisted of at least five animals.

Analgesia was quantified by the means of the Plantar Test and Tail Flick Analgesia Meter apparatus (IITC Life Science Inc.). The rats were gently restrained and a radiant heat beam was projected onto the dorsal side of the tail at cca 1/3rd of its length. The latency time before the tail withdrawal was measured by a built-in timer. The thermal stimulus was reapplied three times at 10 s intervals. The temperature of the beam was adjusted so that the baseline response was approximately 2 s. If a rat (not having been administered a test compound) exhibited the baseline response of more than 4 s, it was excluded from further experiments. For the purpose of avoiding tail burns the apparatus was not allowed to emit the beam for more than 7 s. Rats’ struggling or ambulation were considered as ambiguous responses and if present, the measurements were repeated.

The measurements were performed before the compound administration, as well as 5, 15, 30, 60 and 120 min after the
administration. Each measurement was repeated three times with some minor changes to the place where the beam fell on the tail.

The analgesia level was calculated for each time point and expressed as a maximum possible effect (% MPE) according to the Brady and Holtzmann formula (Brady and Holtzman 1982):

\[
%\text{MPE} = 100\% \times \frac{\text{postdrug latency} - \text{predrug latency}}{\text{maximum latency (7s)} - \text{predrug latency}}
\]

Cell Cultures

Human melanoma cell lines MeW151, MeW155, MeW164, human lung cancer E14, urinary bladder carcinoma cell line T24, and also normal adult human fibroblast cell lines Fib9 and FIW180 and human foetal fibroblast cell line FIWp95 were obtained from the institutional cell bank at the Maria Skłodowska-Curie Memorial Institute and Oncology in Warsaw. The T24 line was generously gifted by Prof. Georg Klein from the Karolinska Institute in Stockholm (Bubeník et al. 1973).

All the lines were cultured in Eagle’s 1959 MEM medium (Biomed, Lublin, Poland) with addition of 10% foetal calf serum (Invitrogen), 50 µg/ml penicillin G, 50 µg/ml streptomycin, and 0.1% glutamine. All the cell lines were kept at 37 °C, under humidified atmosphere containing 5% CO₂.

Cell Proliferation Assay

The cells were plated in 24-well plates (5000 cells per well). After two hours (with the cells adhered to the well bottoms), the compound AWL3020 was added to the wells in the range of concentrations. These were: 100 µM, 50 µM, 25 µM for all cell lines; and additionally for MeW151 and MeW164: 10 µM, 5 µM and 2.5 µM. The control consisted of wells where the cells were untreated with AWL3020. After 24-h incubation (in the presence of AWL3020 or without it), the tetrazolium salt (MTT) was added to the wells. The plates were then kept at 37 °C. Four hours later, the detergent was added (according to the ATCC protocol), and the plates were stored in the dark for two hours at room temperature. Then, the absorbance was read at 570 nm by using HR 7000 spectrophotometer (Hamilton). The absorbance values are proportional to the number of viable cells. Two independent experiments (each data point in three repetitions) were performed.

MTT assay

Cell viability was assessed by the MTT ((3-(4,5-dimethylthiazolyl-2)-2,5-diphenyltetrazolium bromide) colorimetric assay which is based on the direct measure of mitochondrial activity. The MTT assay was performed following the method proposed by the American Type Culture Collection (ATCC), catalogue number 30-1010K (Collection ATC MTT Cell Proliferation Assay Instruction Guide 2016). The cells were seeded in 96-well plates (5000 cells per well). The culture medium was RPMI-1640 (Gibco) with 10% foetal calf serum (Invitrogen). After two hours (with the cells adhered to the well bottoms), the compound AWL3020 was added to the wells in the range of concentrations. These were: 100 µM, 50 µM, 25 µM for all cell lines; and additionally for MeW151 and MeW164: 10 µM, 5 µM and 2.5 µM. The control consisted of wells where the cells were untreated with AWL3020. After a 24-h incubation (in the presence of AWL3020 or without it), the tetrazolium salt (MTT) was added to the wells. The plates were then kept at 37 °C. Four hours later, the detergent was added (according to the ATCC protocol), and the plates were stored in the dark for two hours at room temperature. Then, the absorbance was read at 570 nm by using HR 7000 spectrophotometer (Hamilton). The absorbance values are proportional to the number of viable cells. Two independent experiments (each data point in three repetitions) were performed.

Colony Formation Assay

Cancer cells (100 cells per dish) were seeded on Petri dish (diameter: 2.5 cm). After two hours, AWL3020 was added to the dishes in certain concentrations. These were: 100 µM, 50 µM and 25 µM. The cells were incubated for 7 days. Then, the cultures were rinsed with PBS and dyed with crystal violet. After 20 min, the dishes with stained cells were rinsed with distilled water. The number of colonies formed were counted under the microscope. The experiment was repeated twice with three replicates. The control consisted in cells without the addition of AWL3020.

Evaluation of Ki-67 Expression

Expression of the Ki-67 protein was evaluated because this protein is a marker for proliferating cells, as it is present in the cell during all phases of the cell cycle (G₁, S, G₂, M), but it is undetectable with the resting state (G₀ phase). The cells (5000 cells per well) were seeded on coverslips (12 mm, Life Sciences) located in 24-well plates. After two hours, the compound AWL3020 was added in a few concentrations. These were: 100 µM, 50 µM and 25 µM. The plates were incubated for 4 days. After that time, the cells were rinsed three times with PBS and then fixed with 4% paraformaldehyde (PFA, 15 min, at room temperature). After another rinsing with
PBS (3 × 5 min), a permeabilising mixture (0.25% Triton in PBS) was added to the cells and left for 15 min at room temperature. For the sake of minimising unspecific binding, a blocking mixture (10% GS Goat Serum in PBS) was added. After an hour, the cells were treated with the primary Ki-67 antibody (dilution: 1:500, IgG1 Novocastra/Leica). The cells were left overnight at 4 °C. After rinsing the cells with PBS (3 × 5 min), the secondary antibody Alexa Fluor 546 (dilution: 1:1000, Anti-Mouse IgG Invitrogen) was added and the cells were left in the dark for another hour. Then, another rinsing with PBS was performed and Hoechst 33,258 staining dye (dilution: 1:150, in PBS, 5 min, in the dark) was added. The dyed and fixed cells were counted under inverted microscope Eclipse Ni-U (Nikon). The proliferation index was the percent of cells exhibiting Ki-67 expression in randomly selected populations. Two independent experiments (each data point in three repetitions) were performed. In each experiment, the counting was performed on 10 views.

Statistical Analysis

Quantitative data from the experiments are expressed as the mean ± standard deviation (S.D., cellular assays) or standard error of the mean (SEM, binding affinity and in vivo analgesic activity). Statistical analyses were performed by analysis of variance (ANOVA) with post-hoc Dunnett test when comparing against the control values or with Tukey’s multiple comparisons test when comparing groups against each other (at significance level α = 0.05). Statistical tests and curve fitting were performed by using GraphPad Prism version 7.0.2 for Windows, GraphPad Software, San Diego, California USA, www.graphpad.com.

Results

Receptor Binding Affinity

In order to validate the design rationale for AWL3020, we have first evaluated receptor binding affinity of the compound for μ- and δORs, as well as for the NK1R. The affinity was determined in a radioligand displacement binding assay. The displacement curves are given in Fig. 2. Half-maximal inhibitory concentration (IC50) is provided in Table 1. AWL3020 binds very well δOR with IC50 being 29.5 nM. On the contrary, it has a significantly lower affinity for the μ-opioid receptor, for which its IC50 reads 1070 nM. Even poorer IC50 value is found in the case of the NK-1 receptor (IC50 = 70,280 nM).

![Fig. 2](image-url) Radioligand displacement curves for AWL3020 binding to a δ-opioid receptors b μ-opioid receptors c neurokinin-1 receptors

| Tested compound | Log IC50 ± SEM | IC50 | Log IC50 ± SEM | IC50 | Log IC50 ± SEM | IC50 |
|-----------------|----------------|------|----------------|------|----------------|------|
| AWL3020         | −5.97 ± 0.10   | 1070 | −7.53 ± 0.10   | 29.5 | −4.15 ± 0.25   | 70,280|
| Reference opioids |               |      |                |      |                |      |
| Morphine        | −8.46 ± 0.07²  | 3.5² |                |      |                |      |
| Biphain         | −8.82 ± 0.12²  | 1.5² | −7.54 ± 0.13²  | 29.0² | ndb            |      |
| Reference NK1R ligands |            |      |                |      |                |      |
| Substance P     | ndb            | ndb  | ndb            |      | −8.94 ± 0.13²  | 1.1² |
| Aprepitant      | ndb            | ndb  | −6.63 ± 0.13²  | 236.5² |              |      |

n/d not determined, ndb no detectable binding at 10⁻⁴ M

¹IC50 values are given in nM

²Unpublished results
Thus AWL3020 is a relatively strong δOR ligand, of similar affinity as a reference peptidic biphelin whose IC₅₀ is 29.0 nM. In the case of μOR binding, the affinity of AWL3020 is low, compared to both the peptidic biphelin as well as prototypical alkaloid morphine. This opioid profile is in fact not unexpected as the sequence of AWL3020 in the opioid part stems from a selective δOR peptide, [D-Ala², Leu⁶]-deltaorphin (Janecka et al. 2004).

What is surprising, AWL3020 exhibits only very poor NK1R affinity. It is more than 4 orders of magnitude lower than that of the reference peptidic Substance P (IC₅₀ = 1.1 nM) and more than two orders of magnitude lower than that of a small molecular antagonist, aprepitant (IC₅₀ = 236.5 nM).

Molecular Modelling

A probable structural basis for high affinity of AWL3020 for δOR is provided by molecular modelling. The highest ranked binding mode is presented in Figs. 3 and 4. The first of the Figures contains a detailed depiction of the contacting receptor residues.

According to the modelling analysis, AWL3020 enters deeply into the binding pocket with the opioid sequence (the N-terminal part) towards the receptor interior. The antitachykinin part (from the 7th position to the C-terminus) locates near the receptor outlet. Let us note here that as the receptor structure used for docking does not contain the N-terminus of the receptor, it is impossible to discuss potential interactions of the tachykinin sequence with the opioid receptor.

Regarding the opioid part of the ligand, it is anchored in the receptor by the canonical charge-assisted hydrogen bond formed by the protonated amine of Tyr¹ with D128. The tyrosine’s side chain contacts a few hydrophobic amino acids of the binding pocket bottom, including e.g. I277, M132, W274, V281, V217 and F218. Relatively closely located is also H278. The Phe³ is surrounded by side-chains of K214, F202 and L200. As to His⁴, it forms hydrophobic interactions with W284. The contacts of Leu⁵ and Leu⁶ include K108, Y109 and L48, V297, H301, respectively. The last amino acid of the opioid sequence, the charged Asp⁷ side chain is predicted to be situated closely to R291 (although our docking results do not show a charge-charge interaction between these two). Nevertheless, given that the receptor structure was rigid in our study, such contact cannot be excluded if receptor side-chains would be considered.

Comparing the obtained binding mode of AWL3020 with the one found experimentally for DIPP-NH₂, (H-Dmit-Tic-Phe-Phe-NH₂, 4RWD PDB structure (Fenalti et al. 2015, Fig. 4c, d), it is found that both have the above-mentioned canonical interaction, amine…D128. However, in AWL3020, the Tyr¹ is displaced ‘down’ towards TM5
compared to Dmt\(^1\) of DIPP-NH\(_2\). Phe\(^3\) of both compounds are located in completely different subpockets. In the case of AWL3020, it is directed to TM5, while in the DIPP-NH\(_2\) the aromatic ring points to TM3 of the receptor. Interestingly, a partial overlap is predicted for Tic\(^2\) residue of DIPP-NH\(_2\) with His\(^4\) of AWL3020.

**In Vivo Analgesic Effects**

The compound AWL3020 displayed high affinity for δOR, so it was tested for analgesic activity in the tail-flick test following intrathecal (i.t.) administration in rats. NaCl solution was used as a negative control, and a positive control consisted in morphine (the ‘gold standard’, 12 nmol/kg dose). The results are presented in graphical form in Fig. 5, and numerical data are given in Table ESM-ANALG-1. Consistent with its receptor affinity, the compound was found to have a strong time- and dose-dependent analgesic activity.

The compound was administered at doses of 10, 40 and 60 nmol/kg. The first of them was not able to produce analgesic effect greater than 45% of the Maximal Possible Effect (MPE) at any of the measured time points. At 40 nmol/kg, the compound gave 53% MPE after 5 min post injection. The peak effect (78.7% MPE) was observed after 15 min and it lasted at a similar level also at 30 min post injection, with a minor drop (to 57.5% MPE) at 60 min post injection. This
level of analgesia persisted even 120 min after the injection. Increasing the dose (60 nmol/kg) did not have a statistically significant influence on the analgesic effect.

Comparing with the analgesic ‘gold standard’, morphine, compound AWL3020 seems a weaker analgesic. Analgesia equipotent to that obtained with i.t. administration of 12 nmol/kg morphine, requires dosing as much as 40 nmol/kg AWL3020. This result is in agreement with receptor affinities. AWL3020 is a relatively strong δOR ligand, however it has low affinity for μOR. Therefore, morphine, a nanomolar μOR ligand, gives stronger analgesia compared to that compound.

Cellular Pharmacological Effects in Cancer and Normal Cells

The effects that AWL3020 has on cancer and normal cells were determined with regard to four aspects, that is with regard to:

a. direct effect on the number of cells following a few days of incubation,
b. cytotoxic effects as found in the MTT test,
c. effects on colony formation,
d. the impact on the cells proliferation as measured by determination of Ki-67 protein expression (proliferation index).

Effect on the Number of Cells Following a Few Days of Incubation

Eight cell lines (5 cancer and 3 normal ones) were incubated in the presence of 25–100 μM of AWL3020. For two melanoma cell lines, additional concentration set of the range 1–10 μM was used. The results are graphically presented in Fig. 6 and the numerical data are given in Table ESM-CYTOX-1.

Cytotoxic Effects in the MTT Test

Cell viability of the AWL3020 compound was measured according to the MTT test. The assay readout is proportional to the number of metabolically active (alive) cells in the tested population. Here again, eight cell lines (5 cancer and 3 normal ones) were incubated in the presence of 25–100 μM of AWL3020. For two melanoma cell lines, additional concentration set of the range 2.5 μM–10 μM was used. The
results are graphically presented in Fig. 7 and the numerical data are given in Table ESM-CYTOX-2.

In the MTT assay, all cell lines (except for MeW155) are significantly sensitive to AWL3020 even at 25 μM concentration. For MeW164 the effect is found also at 10 μM concentration (EC50 = 37.78 μM, Figure ESM-CYTOX-2), and for MeW151 at 2.5 μM. Comparing the values obtained at 100 μM (that is a concentration at which all cell lines are significantly sensitive), we found an order of sensitivity: T24 ~ MeW164 > FIW180 ~ FIWp95 ~ Fib9 ~ E14 ~ MeW151 > MeW155 (see Listing ESM-CYTOX-2 for the results of statistical analysis, including the statistical significance of this comparison). Thus we observe no clear pattern with respect to type of cells (normal vs cancer ones) and the sensitivity in MTT assay. There is also no correlation between results of MTT and the number of cells after the multiday incubation (coefficient of determination $R^2 < 0.30$, Figure ESM-CYTOX-3).

**Effects of the AWL3020 Compound on Colony Formation**

Further, we evaluated the influence of AWL3020 on the ability of cells to form colonies. In this test, only cancer cells (5 cell lines, concentrations 25–100 μM) were subject to analysis. This is because normal cells did not form colonies in the particular testing conditions (seeding density, time) that we used for the test. The results are graphically presented in Fig. 8 and the numerical data are given in Table ESM-CYTOX-3.
Here, we found statistically significant differences in colony formation only at 100 μM concentration. It was however not observed for MeW155 for which none of the results is different than the control (at α = 0.05). Leaving aside this cell line, we see that the results for remaining cell lines are essentially identical (no difference at α = 0.05, Listing ESM-CYTOX-3), being 85 ± 5% of the control on average.

Impact on Proliferation Index

As the last element of cytotoxicity testing we determined the impact of AWL3020 on the Ki-67 index. The protein is a marker for proliferating cells, as it is present in the cell during all phases of the cell cycle (G1, S, G2, M), but it is undetectable with the resting state (G0 phase). In this test three cell lines (2 cancer and 1 normal one) were incubated in the presence of 25–100 μM of AWL3020. The results are graphically presented in Fig. 9 and the numerical data are given in Table ESM-CYTOX-4.

For MeW151 cell line, a statistically significant influence of AWL3020 on proliferation index is found for as low as 25 μM concentration of the compound. In the highest concentration, AWL3020 reduces the proliferation index to 38 ± 3% (control value: 72 ± 3%). Another cancer cell line, T24, is less sensitive with regard to the proliferation index. In our tests it displayed a significant reduction of the index just at 50 μM concentration of AWL3020. With 100 μM, the index was 59 ± 10% (control value: 88 ± 10%). The normal
cell line, FlWp95, was found significantly sensitive only at 100 μM, with the index value of 59 ± 13% (control value: 85 ± 12%).

Discussion

The compound presented here, AWL3020, is a hybrid peptide created by simple joining of δOR and NK1R pharmacophoric sequences. Consistently with the design, the compound displays high δOR affinity and a rather low μOR binding. On the other hand, contrary to the design assumptions, AWL3020 has only very poor affinity for NK1R. We have also found that AWL3020 produces a relatively strong analgesic effect in vivo. Regarding the effects on cells, as desired, our compound has no pro-proliferative action. In fact, it displays some antiproliferative effects in higher concentrations which is however not selective with respect to normal or cancer cells.

Regarding the opioid affinity and analgesic action, the reported results are not surprising. The sequence of δOR agonist produces an opioid affinity within reasonable boundaries (IC_{50} = 29.5 nM) compared to the parent opioid compound. This in turn results in a relatively strong analgesic effect. What must not be unnoticed, the effect is significantly less pronounced than in the case of a gold standard analgesic, morphine. This is consistent with the receptor affinity profile of our derivative (very low μOR affinity and reasonable δOR binding). Notably, AWL3020 is weaker than our previously reported hybrid JZ031, which is associated with the presence of the opioid part. What must not be unnoticed, the effect is significantly less pronounced than in the case of a gold standard analgesic, morphine. This is consistent with the receptor affinity profile of our derivative (very low μOR affinity and reasonable δOR binding). Notably, AWL3020 is weaker than our previously reported hybrid JZ031, which was characterized by a fast onset of analgesic activity (Dyniewicz et al. 2017).

That our derivative does not have appreciable NK1R affinity is somehow unexpected given that AWL3020 is created based on a slightly modified sequence of the established NK1R antagonist, spantide I. It might mean that either this slight modification decreases NK1R affinity or this decrease is associated with the presence of the opioid part.

Considering the first of these options, let us recall here that AWL3020 contains C-terminal fragment ‘…D-Phe-Phe-D-Phe-Leu-Leu-NH$_2$’, which is based on the spantide I sequence whose counterpart fragment is ‘…D-Trp-Phe-D-Trp-Leu-Leu-NH$_2$’. The change in sequence, D-Phe$^7$ instead of D-Trp$^7$ and D-Phe$^9$ instead of D-Trp$^9$, is rather conservative in nature (bulky hydrophobic side-chains) and it was not expected to produce a significant change in affinity. To the best of our knowledge, peptides with exact ‘…D-Phe-Phe-D-Phe-Leu-Leu-NH$_2$’ sequence have not been reported so far except for a report of AA3052 by Kowalczyk et al. (Kowalczyk et al. 2016, 2018) who however did not evaluate the receptor affinity but inferred that the compound acts via NK1R based on functional studies. On the other hand, early works on structure–activity relationships of Substance P analogues revealed that various derivatives with D-Phe$^7$ and/or D-Phe$^9$ modifications were able to maintain reasonable NK1R affinity and antagonism (Folkers et al. 1981; Bailey and Jordan 1984; Dutta 1993). Regarding the second option, that the low NK1R affinity is due to the presence of the opioid part, previous reports on the opioid-(anti)tachykinin hybrids provide useful material for comparisons. In the past many papers dealing with such compounds were reported, including a few with our contributions (Matalinska et al. 2013; Dyniewicz et al. 2017). Yamamoto and co-workers (Yamamoto et al. 2007) considered a series of compounds, where an enkephalin fragment was joined with mixed organic/peptide antitachikinin pharmacophore, H-Tyr-D-Ala-Gly-Phe-Xxx-Pro-Leu-Trp-O-3,5-Bzl(CF$_3$)$_2$ (Bzl—benzyl, Xxx—various amino acids). The NK1R inhibition constant (K$_i$) values varied only slightly, between 0.2 and 3.0 nM. A much more pronounced variation was observed in another paper by these authors (Yamamoto et al. 2008), where they considered derivatives of H-Tyr-D-Ala-Gly-Phe-Pro-Leu-Trp—… with modifications of the C-terminal part directly involved in the interactions with the NK1R. Here, K$_i$ values spanned from as low as 0.0046 nM to 100 nM. Further analogues in which the opioid part was cyclized were reported (Yamamoto et al. 2010) and here, the derivatives in which -Pro-Leu-Trp-NH-3,5-Bzl(CF$_3$)$_2$ fragment was left outside the cycle had subnanomolar K$_i$ values. If however, only the Trp-NH-3,5-Bzl(CF$_3$)$_2$ fragment was exocyclic, a significant affinity drop was observed. In another contribution by this group (Yamamoto et al. 2011), modifications of the C-terminal part (NK1R pharmacophore) resulted in variations of three orders of magnitude range, still however the weakest derivatives had nanomolar K$_i$ values. In another series of hybrids (Giri et al. 2015), the focus was on modifying the 1st and 4th position of the enkephalin sequence. These substitutions did not affect the NK1R affinity and the K$_i$ values for all derivatives in the paper were of a few nanomoles. The structure of the enkephalin-(anti)tachykinin hybrid was skilfully truncated (Nair et al. 2013, 2015) which allowed for finding a very short compound of structure, H-Tyr-D-Ala-Trp-O-3,5-Bzl(CF$_3$)$_2$ (NK1R K$_i$ = 0.004 nM) in which opioid and NK1R pharmacophores significantly overlapped. A series of structural changes that led to this derivative, shortening of the sequence and substitutions, resulted in NK1R affinity fluctuations of two to three orders of magnitude, but still all reported K$_i$ values were subnanomolar. Another group of chimeras found by fusing both pharmacophores so that they partially overlapped, included Dmt-DALDA (Riba et al. 2002) sequence as the opioid pharmacophore, with H-Dmt-D-Arg-Aba-Gly-N-3,5-Bzl(CF$_3$)$_2$ being the first example (Ballet et al. 2011). This compound showed very good NK1R affinity (K$_i$ = 0.5 nM). Modifications of the N-terminal part resulted in much variation of affinity even up to micromolar values (Guillemyn et al. 2015).
ties and NK1R affinity with \( K_i = 180 \) nM (Matalinska et al. 2017). It is also 3-arylpiperidine-based NK1R pharmacophores that were successfully introduced into the discussed type of hybrids (Betti et al. 2015). In that study, low nanomolar \( K_i \) values for NK1R affinity were observed, notwithstanding changes in the opioid part.

Overall, in the above summarized literature data, NK1R affinity significant drops were associated with changes of the C-terminal, neurokinin-1 pharmacophoric part and not with the modifications in the opioid part. Whether this is the case for AWL3020 cannot be stated as certain without further studies. A probable answer to this problem might come from molecular modelling using recently published structures of the receptor (Yin et al. 2018; Schöppe et al. 2019; Chen et al. 2019b). We did not attempt such modelling, as the structures were unavailable at the time our study was being performed. Further, the structures are solved in the present of small molecular ligands, so proper modelling will certainly require a careful validation of the binding site for a peptide ligand, in particular against experimental hints from the literature (Pellegrini et al. 2001; Valentín-Hansen et al. 2014).

Regarding the influence of AWL3020 on cancer and normal cells, the compound displays no pro-proliferative action in any of the tests used. In fact, for several combinations cell line/assay, some antiproliferative activity could be found. For instance, incubation with AWL3020 reduced the number of MeW164 cancer cells (compared to control) with \( EC_{50} = 46.27 \) μM. This cell line was also found very sensitive to action of AWL3020 in MTT assays (\( EC_{50} = 37.78 \) μM). However, such activity levels were not found with other cell lines and excluding MeW164 from comparisons, we found no strong, general selectivity in the action of AWL3020 on cancer and normal cells.

So far, the only reports on the influence that hybrids made of opioid and NK1R pharmacophores exert on the proliferation of cancer or normal cells came only from our group. In our previous contribution, we have disclosed compound AA3266 that showed nanomolar opioid affinities and hNK1R affinity with \( K_i = 180 \) nM (Matalinska et al. 2013). At the same time it showed significant selectivity in influencing the proliferation of cancer and normal cell lines (similar set to the one studied here).

Other hybrid opioid peptides assessed as to their action on cancer cells were made of \( \mu \)-opioid tripeptide sequence and trans-1-cinnamylpiperazine moiety (Laskowska et al. 2017). A significant antiproliferative effect on cells was reported. The discussed high affinity \( \mu \)OR ligands exhibited much greater impact on cell viability in 2D and 3D cell cultures of pancreatic cancer than a reference drug gemcitabine. The effect was however not selective and normal cells were also affected to a high degree. The authors did not investigate a putative mechanism of the antiproliferative action of their compounds.

Our working assumption was that introduction of NK1R antagonistic component would result in some antiproliferative (cytotoxic) activity against cancer cells. AWL3020 however turned out to have very low NK1R affinity. The question arises of whether the observed antiproliferative action could be associated with this only low affinity. The reports of antitumour activity of NK1R antagonists dealt with high affinity antagonists, including aprepitant, L-733,060 or L-732,138. These compounds, which are nanomolar binders of the NK1R, were shown to exert antiproliferative influence on various cancer cells lines with IC\(_{50}\) values of several to several dozen micromoles, depending on the cell line type (Muñoz and Rosso 2010; Muñoz et al. 2010, 2015; Rosso et al. 2012; Munoz and Covenas 2012; 2016, 2019). Further, this activity was found to occur via the receptor and by the induction of cell apoptosis. Our low-affinity hybrid achieves similar effective concentration only with a melanoma cell line, MeW164. In this case, we have a distinct proportion of NK1R affinity and toxicity: nanomolar affinity/micromolar cytostaticity in the previous reports, and micromolar affinity / micromolar cytostaticity for AWL3020. This could be indicative of some other mechanism being at work here, e.g. a one via classical or non-classical opioid receptors.

Even though opioids were often found to have pro-proliferative effects on cancer cells, other reports brought opposing findings (Tegeder et al. 2003; Tegeder and Geisslinger 2004; Chen et al. 2008; Lazarczyk et al. 2010; Lennon et al. 2014; Gonzalez-Nunez et al. 2014; Kim et al. 2016). This discrepancy in results stems from different opioid types, cell lines, modes and scheme of administration and probably other factors in a particular study. They also reflect complexity of the effects opioids have on cell growth, survival and death (Tegeder and Geisslinger 2004). There were different suggestions as to whether these actions might be associated with opioid receptors of GPCR family or with other non-classical opioid receptors. A thoroughly explored phenomenon is the anti-proliferative effect of [Met\(^5\)]-enkephalin (called otherwise opioid growth factor, OGF) exerted via OGF receptors (OGFr) (McLaughlin and Zagon 2012). These intracellular binding sites do not appreciably bind many typical opioids, like DAMGO, DPDPE, Dynorphin A or morphine, while binding [Met\(^5\)]-enkephalin. Much evidence has been reported that this OGF–OGFr interaction inhibits proliferation in both normal and cancer cells (McLaughlin and Zagon 2012). The mechanism behind this effect involves regulation of cell cycle (G\(_2\)/S phases) and induction of cyclin dependent inhibitory kinases p16 and/or p21 expression. Further research on AWL3020 must certainly address involvement of the classical and non-classical opioid receptor pathways in its effects on cells.

Interestingly, there are some reports on -D-Phe-Phe-D-Phe-hexapeptides (or derivatives) exhibiting relatively strong antifungal activity (Kundu et al. 2000, 2002). Whether the
toxicity of AWL3020 could have anything in common with this activity is hard to even speculate on, as the mechanisms behind microbial and mammalian cytotoxicities are often utterly different. However, for the sake of comprehensiveness let us note it at this place.

In general, the described antiproliferative action of AWL3020 is not very high. The effective antiproliferative concentrations seem unlikely to be reached with normal administration routines, in particular when one considers metabolic instability of peptides. It means that this antiproliferative action of AWL3020 cannot be expected to be either a problem in analgesic treatment or an advantage from the point of view of auxiliary anticancer properties. The latter could be achieved only after further modifications of AWL3020 structure that could bring more NK1R affinity so that selective anticancer activity would follow. Else, if a mechanism different than via NK1R could be elucidated, by which AWL3020 exerts its antiproliferative action, one could attempt to optimize the compounds’ structure to obtain anticancer effects via this putative mechanism. Anywise, it is clear that AWL3020 structure requires more work, as the design assumptions were only partially realized. That the search of analgesic compounds with adjuvant anticancer properties (Matalinska et al. 2013) is still worth pursuing emerges from a recent study (Ge et al. 2019) where the authors were able to demonstrate that a NK1R antagonist SR140333 exerts anticancer action in mice xenografted with human myeloid leukemia cells and further produces a potent antinociceptive effect in leukemia-induced bone pain model.

Conclusions

In the present paper we have reported the design, synthesis and in vitro evaluation of AWL3020, a hybrid peptide made of delta opioid agonist and neurokinin-1 antagonist sequences. The compound displayed high δOR binding, but contrary to the design assumptions only a very poor affinity for the NK1R. As strived for, the novel hybrid has no pro-proliferative action on either cancer or normal cells. In larger concentrations, it displays antiproliferative activity which is however not much selective.

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Compliance with Ethical Standards

Conflict of interest The authors declare no conflict of interest.

Ethical Approval All animal experiments were approved by the IV Local Ethics Committee for Experiments on Animals in Warsaw, Poland.

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