Cytotoxicity of *Quillaja saponaria* Saponins towards Lung Cells Is Higher for Cholesterol-Rich Cells

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Abstract: The purpose of the study was to compare cytotoxicity of two *Quillaja saponaria* bark saponin (QBS) mixtures against two lung cell lines: normal MRC-5 fibroblast cell line and tumor A-549 epithelial cells of lungs’ alveoli. The study, performed both at a macro-scale and in a dedicated microfluidic device, showed that QBS was more toxic to the cell line more abundant in cholesterol (MRC-5). The QBS mixture with higher saponin fraction was found to be more cytotoxic towards both cell lines. The results may help to better understand the cytotoxicity of saponin-rich herbal medicines towards normal and tumor cells depending on their cholesterol content.

Keywords: MRC-5; A549; cytotoxicity; saponins; microfluidics

1. Introduction

Saponins are secondary metabolites found mostly in plants [1]. Their biological role has not been fully elucidated yet, but is most likely related to the plant organisms’ defense against predators [2]. Certain saponins are toxic in their native form, while others become biologically active only upon chemical transformation. For example, the normally non-toxic saponins avenacosides A and B, stored in healthy leaves of oat, can become hydrolyzed by the plant’s enzyme avenacosidase to release antifungal membranolytic products in response to a fungal infection [3]. Saponins are by no means the sole biologically active molecules produced by plants, yet the cooperative activity of different biomolecules is often ignored. One known example of such activity is the saponin–saporin cooperativity. Saponins are ribosome-inactivating protein (RIP) toxins that require assistance of saponins to successfully enter the cytosolic compartment of the attacked cell [4,5].

Many saponins, including *Quillaja saponaria* bark saponin (QBS), display pronounced amphiphilic character [6–8] and have been traditionally used mostly as foaming and cleaning agents; hence the name “saponins”, which derives its origin from a Latin word for soap—“sapo” [9,10]. Purifying saponin-rich plant extracts such as QBS, containing dozens of saponins and several other components (polyphenols, tannins, sugars, calcium oxalate, etc.) [11] is still a complicated and time-consuming task [10,12]. Hence, in most cases, mixtures of saponins are used, except for special applications, such as an adjuvant for vaccines QS-21 [13]. The hydrophobic parts of saponin molecules present in QBS comprise several triterpenoid aglycones, including quillaic acid (Figure 1). Their hydrophilic sugar chains (glycones) consist of typically 2–5 frequently branched sugar units attached to the C-3 and C-28 carbon atoms of the aglycone. The exact composition of a given QBS product depends on a number of factors, including the botanical conditions and extraction procedure (the use of highly alkaline or acidic conditions, temperature, etc.), but also on the sample history (e.g., storage conditions) [13–16].
It has shown a great promise in cancer research [46], point of care diagnostics [47], and other benefits of the microfluidic approach in biology come from a comparable scale of DNA analysis [48] thanks to a rapid sample processing and low reagent consumption [49].

The microfluidic approach offers numerous advantages for testing cytotoxicity [45]. It has shown a great promise in cancer research [46], point of care diagnostics [47], and DNA analysis [48] thanks to a rapid sample processing and low reagent consumption [49]. Other benefits of the microfluidic approach in biology come from a comparable scale of the device microstructure and of the cells. Furthermore, flow conditions, high surface area to volume ratio, fast diffusive heat, and mass transfer enable the process to more closely mimic the in vivo cell–cell or cell–extracellular matrix signals [50–52]. Despite the high potential for high-throughput screening, the microfluidic approach is rarely employed in cytotoxicity studies of saponins. To the best of our knowledge, only one study has thus far been devoted to the analysis of saponin cytotoxicity using a microfluidic setup [53].

In this contribution, we used a macroscopic cytotoxicity test (MTT) and a dedicated microfluidic setup (with dead cells stained with propidium iodide and the living ones with calcein AM) to compare cytotoxicity of two commercially available QBS mixtures ("Sigma” from Sigma-Aldrich and “SuperSap” from Desert King Int). In our previous report, the effect of both mixtures on model lipid monolayers was compared, pointing to significant differences resulting from the differences in saponin profiles and total saponin content [16]. The question that we want to answer in this contribution is to what extent the total amount of cholesterol in two lung cell lines and the saponin profile of two QBS mixtures affect the cytotoxic activity.

Figure 1. General structure of Quillaja bark saponins (QBS); R1 and R2 are different sugar groups.
2. Materials and Methods

2.1. Chemicals

Two commercially available *Quillaja* bark saponin mixtures (QBS) were used: Saponin (Sigma-Aldrich, 84510) and Super Sap (Desert King Int). They will be referred hereafter as “Sigma” and “SuperSap”, respectively. Their saponin profiles were acquired using a reverse phase high performance liquid chromatography (RP-HPLC), as described in [16]. The chromatograms and the relative peak areas of the identified individual saponin for both mixtures are shown in Figure S1 (Supplementary Data). Trypsin (Sigma: T4799), phosphate-buffered saline (Sigma: P5493), Eagle’s minimum essential medium (Sigma: M4655), and 70% ethyl alcohol (POCH Poland) were used for cell culturing. Propidium iodide (Life Science: P4170), Calcein AM (Fluka: 17783), and Trypan Blue (Invitrogen: T10282) were used for cell staining. MTT assay consisted of a MTT Cell Proliferation Assay Kit (Vybrant: V-13154) and dimethyl sulfoxide (Sigma: D4540). Silicone elastomer curing agent and a polydimethylsiloxane(PDMS) silicone elastomer (Sylgard 184, DOW Corning), A1518 Developer (MicroChemicals GmbH), Photoresist S1818 (MicroChemicals), Capillary film, and Pro Cap 50 (Chromaline) were used for the microfluidic setup. Other chemicals—hydrofluoric acid, ammonium fluoride, acetone, methanol, and isopropanol—were of analytical purity (p.a.) and were purchased from POCh Poland.

2.2. Cell Lines

Human cell line A-549—lung cancer epithelial cells (ATCC: CCL-185) and human cell line MRC-5 (ATCC: CCL-171) were used as lung cancer and normal cells, respectively.

2.3. Cholesterol Content

The whole-cell cholesterol content in dead cells from A-549 and MRC-5 lines was determined spectrophotometrically and fluorimetrically using a MAK043 kit from Sigma Aldrich. Cholesterol from the dead cells was extracted with a 7:11:0.1 mixture of chloroform, isopropanol, and Triton X-100 under sonication for 20 min. The extract was centrifuged for 20 min at 4500 rpm and dried using dry air and by storing in a desiccator under vacuum for 30 min. The lipids were then suspended in a buffer and placed in a 96-well plate for quantitative analysis following the instructions of the MAK043 kit.

2.4. Microfluidic Setup

To fabricate microcavities for cell culture, we coated clean sodium glass slides (75 × 25 × 1 mm) under clean-room conditions with a thin layer of a photoresist using spin coating technique (1 min, 2000 rpm). They were subsequently irradiated with UV light for 3 min with the applied mask and washed with the developer and Milli-Q water, dried with nitrogen, and digested for 25 min with a mixture of NH₄F and HF (6:1). The PDMS part was fabricated with soft lithography method. First, a capillary film was placed on the sodium glass plates (75 × 25 × 1 mm) using a double-sided adhesive tape. The plates were then irradiated with UV light for 3 min through the mask with the concentration gradient generator pattern [54]. This stamp with a mapped design was covered with the liquid mixture of the prepolymer and cross-linking agent and heated (70 °C, 2 h) until curing. In order to equip the PDMS layer with the inlets and outlet, we cooled the polymer with a liquid nitrogen prior to drilling the holes (1.3 mm). The PDMS part was finally bonded with the glass part in order to obtain the finished microfluidic system chip (Figure 2).
2.5. Cytotoxicity Measurement in the Micro Scale

A pure culture medium and the “Sigma” or “SuperSap” QBS solution of 200 μg/mL concentration were introduced simultaneously into the chip at the rate of 1.5 mL/min during 30 min with a peristaltic pump. The concentration gradient generator (as described in more detail in [54]) was employed to fill 5 lines of chambers with the solutions of 5 different QBS concentrations (200, 150, 100, 50, 0 μg/mL). The cell viability was determined with a differential staining method. For this purpose, a mixture of propidium iodide (red color fluorescence, staining dead cells) and calcein AM (green color fluorescence, staining live cells) was introduced into the chip at the rate of 1.5 μL/min for 10 min. Fluorescence images of each chamber were taken using an inverted fluorescent microscope (Olympus IX71).

2.6. Cytotoxicity Measurement in the Macro Scale Using MTT Test

The 200 μg/mL QBS solutions (“Sigma” or “SuperSap”), prepared as above, were diluted with pure culture medium in order to achieve the concentrations of 200, 150, 100, and 50 μg/mL. The cell line passaging was performed on a sterile 96-well plate in the amount sufficient to obtain the cell density of 10^5 cells per well and incubated for 24 h in 37 °C until the cells adhered to the plate surface. Next, the medium from above the cells was removed and the respective QBS solutions were added (200 μL per well). The plates were then incubated for 24 h at 37 °C. The test was performed according to the instructions of the Vybrant MTT Cell Proliferation Assay Kit. The medium from above the cells was replaced with the MTT reagent (60 μL per well), followed by incubation at 37 °C until the conversion of MTT to formazan and consequent precipitation of the purple formazan crystals from the yellow MTT solution. After around 4 h, the solution was carefully removed from above of the cells and 200 μL of DMSO was added to each well to dissolve the formazan crystals. After 10 min of incubation at 37 °C, the absorbance at λ = 570 nm was measured on a previously calibrated multiwell plate reader (BIOTEK, Cytation 3).

3. Results

Saponins are known for their high affinity to membrane lipids, especially to sterols [14, 16,22,23,25,29,55,56]. Consequently, numerous postulated mechanisms of membranolytic activity of saponins assume that the primary site of saponin attack is cholesterol [57–61]. To support this hypothesis, in the present study, we compared the effect of *Quillaja* saponins on cell lines differing in total cholesterol content. For this purpose, we chose two lung cell lines. A-549 is a lung cancer cell line, first developed by J. Giard et al. in 1972 from the explanted tumor of a 58-year-old Caucasian male. A-549 cells are basal epithelial cells of lungs’ alveolus. The cells are adherent in in vitro environment, forming a monolayer [62]. The second cell line, MRC-5, are fibroblasts derived from normal lung tissue of a 14-week-
old Caucasian male in 1966. In in vitro environment, the adherent cells of MRC-5 line also form a monolayer.

In the first step, the total cholesterol content in the cell lysates of A-549 and MRC-5 was determined using the spectrophotometric and fluorimetric assays. The two methods provided comparable results: 0.070 ± 0.001 µg/mL and 0.101 ± 0.007 µg/mL, respectively, for MRC-5, and 0.030 ± 0.001 µg/mL and 0.047 ± 0.016 µg/mL for A-549, respectively. Interestingly, despite some discrepancy between the two methods, the ratio between the determined cholesterol content in MRC-5 and A-549 lysates was very close: 2.3 (for the spectrophotometric) and 2.2 (for the fluorimetric). Thus, for the purpose of this study, the normal lung fibroblast cells (MRC-5) were used as a model high-cholesterol cell line, while their tumor counterparts (A-549) were used as a low-cholesterol cell line. All subsequent experiments were performed in parallel for two commercially available QBS mixtures in order to enable discussion of possible effects of the differences in their composition. The mixtures are the same as those used in our previous study on the effect of QBS on model lipid monolayers and on the red blood cells: “Sigma” and “SuperSap” (see experimental section for more details).

The cytotoxicity was first assessed in a macro-scale using the cell metabolic activity test (MTT), as described in the experimental section. The results comparing the cytotoxic effect of both saponin mixtures on A-549 and MRC-5 cell lines obtained from the MTT test are shown in Figure 3. Significant differences in general toxicity between the two extracts can be easily noticed. In the case of “SuperSap”, already at the lowest tested concentration (50 µg/mL), only 20% of the cells survived, and thus the half maximal inhibitory concentration, IC50, for both lines was below 50 µg/mL. The “Sigma” QBS was clearly less toxic, especially towards the cancer cells (A-549), for which a significant reduction of the cell viability could be noticed only at the highest concentration (200 µg/mL). Consequently, for the normal cell line, IC50 of “Sigma” could be estimated at around 50 µg/mL, and for the cancerous cell line, at around 200 µg/mL.

![Figure 3](image)

**Figure 3.** Viability of A-549 and MRC-5 cells (MTT test) after 24 h of incubation with (a) “Sigma” and (b) “SuperSap” *Quillaja saponaria* bark saponin (QBS) solutions on a macro scale. We assumed 100% cell viability for samples not treated with QBS solutions. Error bars were calculated from 18 measurements for three independent experiments.

In the next step, the cell viability was tested using a microfluidic setup. The setup allows for observation of the cell viability under a fluorescence microscope thanks to differential staining of alive (green) and dead cells (red). In line with the previously described results of the macroscopic MTT test, the representative microphotographs for 0, 50, 100, and 200 µg/mL collected in Figure 4 point to important differences between the responses of both cell lines to QBS. The effect of “SuperSap” is evident at lower concentrations than for “Sigma”. The viability results obtained from the analysis of the microphotographs of at least three independent microfluidic chips are collected in Figure 5. For both “SuperSap” and “Sigma” QBS, a dose-dependent cytotoxicity was observed. In both cases, QBS solutions were also more toxic towards the MRC-5 normal cells, with “SuperSap” being generally a stronger cytotoxic agent. For this mixture, the MRC-5 viability
was about 20% at merely 50 µg/mL, while above the concentration of 100 µg/mL both normal and cancer cells were already dead. For “Sigma”, the full toxicity towards both cell lines was observed only at the highest tested concentration (200 µg/mL). Below this dose, the A549 viability remained at a quite high level above 60%.

Figure 4. Photographs of A-549 and MRC-5 cell cultures after 24 h of incubation with “Sigma” and “SuperSap” QBS solutions (10× magnification). The top two rows (a) present the cells incubated with “Sigma” solution, and the bottom two (b) are these incubated with “SuperSap” QBS solution. The concentrations of saponin solutions in both cases were (from left) 0, 50, 100, and 200 µg/mL, respectively.

Figure 5. Viability of A-549 and MRC-5 cells after 24 h of incubation with (a) “Sigma” and (b) “SuperSap” QBS solution in the microfluidic device. The error bars were calculated as standard deviation from 12 and 8 independent experiments for A-549 and MRC-5 cells, respectively.
4. Discussion

Comparing the macro- and the microscale results, we observed a very similar toxic effect of “SuperSap” QBS, with the exception of the lowest saponin concentration (50 µg/mL), where the A549 viability was significantly higher in the microscale. In the case of “Sigma” QBS, the results obtained with the MTT assay and microfluidic setup also showed a similar pattern, although the viability was systematically higher for the MRC-5 line in the macroscale. In the microfluidic setup, the complete cytotoxic effect of “Sigma” QBS could only be achieved at the highest saponin concentration. The differences in cytotoxicity between the two QBS mixtures were reproducible in both experimental setups and for both cell lines were clearly related to their composition. Interestingly, the higher biological activity of “SuperSap” (yet lower than typically observed for steroidal saponins, e.g., digi- tonin [24]) was previously observed also in hemolytic tests [16]. In the same contribution, we showed that the two mixtures differed significantly in their total saponin content, with “SuperSap” containing 20% more saponins than “Sigma” (see Figure S1 in Supplementary Data and [16]). The generally higher cytotoxicity of “SuperSap” might thus at least partially be explained by its higher saponin content. On the other hand, the natural consequence of the lower saponin content of “Sigma” is the increased amount of a non-saponin fraction. In the HPLC chromatogram, it accounted for 17.4% of the total peak area (vs. only 2.7% in “SuperSap”, see Figure S1 in Supplementary Data and [16]). This fraction is probably rich in hydrophilic tannins and phenolic compounds, such as (+)-piscidic acid, p-coumaric acid, glucosyringic acid, and vanillic acid, to name just a few found by Maier et al. in commercially available QBS mixtures [63]. It might be speculated that the non-saponin fraction of “Sigma” protects both types of cells against the membranolytic activity of the saponin fraction, further weakening cytotoxicity of the mixture. It is commonly known that polyphenols may serve as protective agents against various toxins and pollutants, even though their mechanism of action is still not clear. Plant polyphenols may also take part in communication between cells and are speculated to play a crucial role in anticarcino- genic, vascular, and cardioprotective activities [64,65]. The third major difference between “Sigma” and “SuperSap” that might significantly affect their cytotoxicity is the saponin profile. The analysis of large sets of data on biological activity of triterpenoid saponin shows that there is no single factor determining their toxicity towards the cells [66–68]. The data on biological activities of individual saponins is still very scarce and fragmentary. Nevertheless, among the few identified saponins present in QBS (Figure S2), there are some with high (e.g., QS-17 and QS-18) and some with low (e.g., QS-7 and QS-21) hemolytic activity [13]. The fact that the HPLC peaks corresponding to QS-17 and QS-18 were indeed the highest for “SuperSap”, and those corresponding to QS-7 were the highest for “Sigma”, correlates well not only with the higher hemolytic activity of “SuperSap”, but also with its higher cytotoxicity described above.

To the best of our knowledge, the only data comparing cytotoxicity of saponins towards the tumor A-549 and normal MRC-5 lung cells concerns steroidal saponins from Dioscorea birmanica [69]. Our present results concerning triterpenoid saponins generally agree with the available literature data on a generally low cytotoxicity of QBS. The higher toxicity against the MRC-5 cells reaffirms the expected poor applicability of both QBS mixtures in fighting lung cancer but, more importantly, confirms the key role of cholesterol in determining their biological activity.

Cholesterol plays a key role in plasma membranes [70], and there is an on-going debate concerning its role in cancer development. Consequently, the contradicting opinions exist on whether the differences in cholesterol content could be eventually employed for any diagnostic or therapeutic purposes. Although the tumor cells are generally known to have more fluid plasma membranes than the normal ones [71], there is no general consensus as to whether they indeed contain more or less cholesterol than their normal counterparts [72]. For example, prostate cancer is clearly linked to an enhanced cholesterol level in the prostate epithelial cells and in blood [73]. On the other hand, the opposite examples can be easily found in the literature. One such example are leukemic cells in
mice and humans, where the unesterified cholesterol levels are lower than in the normal leukocytes [74]. Although in this study only the whole-cell lysates were assessed for the total cholesterol content, both spectrophotometric and fluorimetric analysis clearly point to about twice higher cholesterol content in the normal MRC-5 cells than in A-549. The cytotoxicity tests showed that high-cholesterol cells (MRC-5) are indeed more susceptible to the toxic effect of QBS than their low-cholesterol counterparts, both in the macroscopic and microscopic setups. In view of the generally accepted crucial role of cholesterol in saponins’ affinity to biological membranes, the present cytotoxicity results could be easily explained by the differences in cholesterol content of both investigated lung cell lines. Whether this relation holds also for other cells with low and high cholesterol content remains an open question. Furthermore, the distribution of cholesterol molecules within the cell probably plays even more important role than its total content.

5. Conclusions

The total cholesterol content in the normal lung cell lines (MRC-5) is about twice higher than that in the tumor (A-549) line. The cytotoxicity of two commercially available *Quillaja* saponin mixtures (QBS) was found to correlate well with the total cholesterol level found in the normal (high cholesterol, high cytotoxicity) and tumor (low cholesterol, low cytotoxicity) lung cell lines. Although both mixtures showed higher cytotoxicity toward the MRC-5 line, the QBS mixture that had previously been shown to be more abundant in the saponin fraction and more hemolytic (“SuperSap”) also showed higher cytotoxicity than “Sigma”. The latter is enriched in the non-saponin fraction of tannins and phenolic compounds, which, combined with a specific saponin profile, is probably responsible for the observed lower cytotoxicity of “Sigma”. The results were confirmed in both macro- and microscopic setups using the metabolic assay (MTT) and cell staining tests, respectively. Although in the present set of cell lines, the tumor cell line (A-549) was less abundant in cholesterol, which rendered it more resistant to QBS, many other cancer cells display elevated cholesterol levels. In such cases, QBS or other saponin mixtures may lead to enhanced cytotoxicity towards the unwanted tumor cells and could hopefully be employed as anticancer agents. This hypothesis requires experimental validation with more cell lines of known cholesterol content. The present observations may also help to explain why in some studies saponins prove effective against cancer cells, while in others this is not the case—their efficacy might in fact strongly depend on the actual cholesterol content in the given line, be it normal or tumor.

The microfluidic setup described in this contribution might serve in the future as a platform enabling for comparison of cytotoxicity of QBS and other potential anticancer drug candidates towards normal and cancerous cell lines. The main advantage of the setup is the minimization of the amount of sample required for single analysis, which might be especially useful for testing of individual components obtained by expensive purification procedures from the crude mixtures, e.g., of plant extracts.

**Supplementary Materials:** The following are available online at https://www.mdpi.com/article/10.3390/biophysica1020010/s1: Figure S1: RP-HPLC chromatograms of QBS mixtures (a) “Sigma”, (b) “SuperSap”. UV–VIS absorbance detection at 210 nm and the resulting relative content of major identified saponins in “Sigma” and “SuperSap” QBS extracts. Figure S2: Structures of identified saponins present in “Sigma” and “SuperSap”.

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