# Phytochemical screening and evaluation of the cytotoxicity

# of fruits of Solanum torvum Swartz (Solanaceae) on HFF

# cells (Human Foreskin Fibroblasts).

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- Objective: The aim of this work is to evaluate the cytotoxic activity of the 70% ethanolic
- 6 extract of the fruits of Solanum torvum Swartz (Solanaceae) on HFF (Human Foreskin
- Fibroblast) cells in *in vitro* culture and to determine its phytochemical composition.
- 8 **Study plan**: An ethnobotanical survey was conducted in August 2015 in the Haut-Sassandra
- 9 Region (Ivory Coast), on medicinal plants with multiple uses, and at the end of this survey
- the fruits of *Solanum torvum* have been selected then harvestedin the Sub-prefecture of
- Bédiala (Ivory Coast). Then after drying, the ethanolic extract of the fruits was prepared and
- sent to France in February 2016 at the Laboratory Adaptation and Pathogenesis of
- Microorganisms (LAPM) of Grenoble for cytotoxic tests. Phytochemical screening was
- carried out at the Faculty of Biological Sciences of Félix Houphouët Boigny University (Côte
- 15 d'Ivoire).
- Methods: From an ethnobotanical survey, the fruits of *Solanum torvum* were harvested. The
- 17 70% ethanolic extract prepared from the fruits of this plant was tested *in vitro* on divisional
- 18 HFF cells after phytochemical screening.
- 19 **Results**: The result revealed that this extract has cytotoxic activity on the tested HFF cells. At
- 20 800 μg/mL, the survival rate of HFF cells increased from 100% to 4% of living cells.
- 21 Phytochemical screening revealed the presence of compounds such as alkaloids, tannins,
- 22 polyphenols, saponins and flavonoids.
- Conclusion: This extract is cytotoxic on HFF cells. It is therefore necessary to continue
- studies on toxicity and to be cautious in the use of *solanum torvum* fruits in traditional
- 25 medicine.
- 26 **Key words**: HFF cell, Cytotoxic, Extracts, ethnobotanical survey, Solanum
- 27 *torvum*, Phytochemicals

#### 28 1. Introduction

- 29 Solanum torvum belongs to the family Solanaceae. Solanum torvum is native to Central and
- South America, from Mexico to Brazil and Peru. It has spread widely in the Caribbean [1]. In
- West and Central Africa, it is grown locally in gardens for cooking [2]. Fruits are widely used

in the treatment of shingles in Cameroon[3]. They are also used as a vegetable and considered an essential ingredient in the diet of the South Indian population[1]. A fruit decoction is used in Ghana for the treatment of cough, liver disease and spleen [4]. Solanum torvum are rich in antioxidant, Its extracts are used in the preparation of tonic and hemopoietic agents and also for the treatment of pain throughout the body [5,6]. Previous work has revealed that the fruits of Solanum torvum are used in the treatment of various conditions: microbial infections [7], hypertention [8], kidney disease [9] diabetes [10]. Pérez-Amador et al. [11] reported the presence of glycoalkaloids in the fruit of Solanum toryum and another study showed that very low doses of glycoalkaloids in some Solanaceae are toxic [12,13]. In view of the interest of Solanum torvum fruits in the traditional medicine, we have undertaken to evaluate scientifically the cytotoxicity and phytochemical screening of the fruits. An ethnobotanical survey was conducted in August, 2015 in the Haut-Sassandra Region (Ivory Coast). This survey showed that all parts of Solanum torvum are intensively used, especially leaves and fruits in the treatment of dermatosis and anemia. The aim of this work is to perform a phytochemical screening and a cytotoxicity study of Solanum torvum fruits on Human Foreskin Fibroblasts (HFF) cells.

#### 2.Material and methods

## 2.1 Plant material

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The fruits of *Solanum torvum* (Fig 1) were harvested in Biadiala in the Department of Daloa (Ivory Coast).





Fig 1: Leafy and fruiting twig of *Solanum torvum* (Solanaceae)

#### 2.3 Cellular material

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- The cellular support consists of human HFF (Human Foreskin Fibroblasts) cells. These are
- 63 human cells that testify to the toxic activity of an extract. When these cells are in culture for
- only 24 hours, they are in a state of mitosis (or dividing cells).

## **2.4 Preparation of plant extracts**

- After harvest, the fruits were freed of impurities, dried in the shade for a week and then
- 67 pulverized with an electric grinder. The fine powders obtained were stored in glass jars to
- 68 prevent mold.

## 69 2.5 Preparation of total aqueous extract (TAE)

- 70 The preparation of this extract was performed using themethod described by Zirihi et
- 71 al[14] which involves macerating 100 g of plant powder of species in 1L of sterile distilled
- vater using a Blinder type 7 SEVEN STAR. The homogenate wasfiltered over hydrophilic
- cotton and then on Whatman filterpaper  $(n^{\circ}_{3})$ . The aqueous filtrate thus obtained is evaporated
- in an oven of Med CenterVenticell type at 50°C to obtain powders that constitute thetotal
- 75 aqueous extract (TAE).

## 76 2.6Preparation of 70 % ethanolic extract (70 %FE)

- 77 The extract was obtained by dissolving 5 g of TAE in 100 mL of ethanol 70% (v;v) solution
- 78 and thenhomogenized. After decantation and filtration of thealcoholic fraction on hydrophilic
- 79 cotton and on Whatman filterpaper(n°<sub>3</sub>), the filtrate collected isevaporated in an oven at 50
- °C. The powder obtained constitutes 70 % ethanolic extract (70 %EE)[15].

#### 81 2.7Yield calculation

- 82 The yield is the quantity of extract obtained from the plant powder. It is expressed as a
- 83 percentage. In practice, it has been determined by the ratio of weight of the solids content after
- evaporation on theweight of the dry plant material powder used for the extraction, multiplied
- by 100. This results is indicated by the following formula:

## 86 $Yd = (m \times 100)/M$

- Yd: Extraction yield in percentage
- m: mass in grams of the dry extract
- 89 M: mass in grams of the drug powder.

### 2.8 Phytochimical screening

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- The identification of different chemical compounds in 70 % ethanolic was done by tubes
- 92 characterizationreactions. This method consists of detecting the different families of chemical
- 93 compounds that may exist in plant extracts on the basis of characteristic colorations or
- 94 precipitation reactions[16].

#### o Alkaloids characterization

- 96 The characterization of alkaloids was made using Bouchard (iodo-iodide) and Dragendorff
- 97 (tetraiodopotassium bismuthate)reagent. 6 mL of 70 % ethanolicextract solution was
- 98 evaporated to dryness. Theresidue was taken up in 6 mL of alcohol at 60 °C. Thefiltrate thus
- obtained was divided into two test tubes. In the first tube, two drops Dragendorffreagent
- 100 wereadded. The presence of alkaloids was characterized byobserving orange-coloured
- precipitates. In the secondtube, two drops of Bouchard reagentswas added. The appearance of
- a reddish-brown color indicates the presence of alkaloids. A control test was made
- withquinine.

## o Characterization of polyphenols

- The polyphenols colorimetry forms colored precipitates with a solution of ferric chloride
- (FeCl<sub>3</sub>). Thus, one drop of alcoholic solution of 2% ferric chloride and 2 mL of solution of 70 %
- 107 ethanolicextract wasadded. The formation of blue-black or green colouringmore or less dark
- 108 confirms to the presence of polyphenols. A control test was performed with asolution of
- 109 phenol.

#### Characterization of flavonoids

- 111 Flavonoids have been characterized by the reaction tocyanidin. Thus, 2 mL of 70 %
- 112 ethanolicextract were evaporated to the dry sand bath. The residue thus obtained was
- mixedwith mL dilute hydrochloric acid 2 times. The mixture was collected in a testtube, in
- which pink-orange or violet colouration will appear. The addition of 3 drops of soamyl
- alcohol intensifies this coloring and confirmspresence of flavonoids. An alcoholic solution

#### o Tannins characterization

- 118 The Stiasny reagent (Formalin 30%, concentrated HCl1/0.5) helped to distinguish the
- catechin tannins (byprecipitation) of gallic tannins (by saturation). Tannins cathéchiques: to

10 mg of 70 % ethanolicextract, were added 10 mL of Stiasny reagent. The mixturewas heated in a water bath at 80 °C for 30 minutes. After cooling in a stream of water, observation of precipitate in the form of clear-brown flakescharacterizes catechin tannins. An alcoholic solution of catechin was used as a control. Gallic tannins: For this test, the filtrate obtained from the reaction of catechol tannins characterization was saturated with sodium acetate. To this mixture was added a few drops of a dilute aqueous solution of FeCl<sub>3</sub> at 1% (approximately 1 mL). The appearance of an intense blue-black coloration indicates the presence of gallic tannins not precipitated by Stiasny reagent. An alcoholic solution of gallic acid was used as acontrol.

#### Terpenes characterization

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Sterols and terpenes characterizationwas made by the Liebermann-Burchard reaction. To 0.2 g of 70 % ethanolicextract, were added 5 mL of ethyl ether, then themixture was macerated for 30 minutes. The solution obtained after the maceration was filtered and then evaporated to dryness. The residue was then dissolved in 0.5 mL of acetic anhydride. Using a pipette, 2 mL of concentrated sulfuric acidwere laid down at the bottom of the test tube withoutstirring. The appearance of brownish red or purple ringreflects the two liquid contact zone. The upper liquid turns green or purple to green or purple indicating the presence of sterols and terpenes. A control test was performed with progesterone.

### Coumarins characterization

- For the detection of coumarins, 2 mg of 70 % ethanolicextract was added to 2 mL of warm water and thenhomogenized. The homogenate thus obtained was divided into two test tubes.
- Thereafter, 0.5 mL ofdiluted ammonia at 25% was added to the contents of one of the tubes.
- 142 After observation under UV 365 nm, the presence of fluorescence in the tube where
- ammoniac wasadded indicates the presence of coumarins.

#### Saponins characterization

- For the detection of saponins, 10 mLof 70 % ethanolicextract was introduced in the test
- tubes. Each tube wasstrongly stirred in a vertical position for 15 seconds, and thenleft to set
- 147 15 minutes. The height of persistentfoam is higher than 1 cm, testifying the presence
- ofsaponins.

## 2.9 Cytotoxicitytest

To measure the toxicity of the ethanolic extract, the Human Foreskin Fibroblasts (HFF) cells were inoculated in 96-well plates (CellStar) at the rate of 3000 to 5000 cells per well in 100 μl of D10 medium. These cells are kept in culture for 24 hours (dividing cells). Subsequently they were exposed for 24 hours at different concentrations (125-800 µg/mL) in plant extract solubilized in PBS buffer. This was done in triplicate, also for control control without plant extract. Viability was determined using 3- (4,5-dimethylthiazol-2-yl) -2,5-diphenyl tetrazolium bromide (MTT). The tetrazolium ring it contains is reduced in formazan by the mitochondrial succinate dehydrogenase of metabolically active cells, which precipitates and gives a purple color. The amount of precipitate formed is proportional to the number of living cells. In each well, MTT is added at a concentration of 500 µg / mL and incubated for 3 hours at 37 ° C. The formazan crystals are solubilized in 10 mM dimethylsulfoxide (DMSO). The measurement of the optical density at 544 nm was made using a Safir spectrophotometer (Tecan); this measurement of absorbance will make it possible to determine the relative quantity of living and metabolically active cells [17]. The results were expressed as a percentage of viability compared to control without plant extract. Viability rate = (Abs544 nm extract / Abs544 nm control)  $\times$  100

#### 3.RESULTS

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## 3.1Yield of different extracts of fruits of solanum torvum

We obtained from 200 g of powder, 20 g oftotal aqueous extract a yield of 10 % and from 5 g oftotal aqueous extract, we got 2 g of 70 % ethanolic extract or a yield of 40 %.

## 3.2Phytochimical sorting

The phytochemical sorting performed with the extractsof fruits of *Solanum torvum* allowed to detect the presence of various chemical groups (Table 1). They are thepolyphenols, tannins, flavonoids, saponins, and alkaloids in 70% ethanol extract.

Table 1: Chemical compounds in the fruits of Solanum torvum

		Chimical compounds							
Species	Extract	Sap	Flav	Terp/ster	Tanins		Coum	Alc	Poly
					Gall	Cathé			
Solanum torvum	EE 70 %	+	+++	-	++	++	-	+	+

- :negative reaction ; + : positive reaction

EE 70 %:70 % ethanolic extract

Sap: saponins; Flav: flavonoids; Terp / Ster: terpenes / sterols; Gall: gallic;

Cathé: cathechic; Coum: coumarines; Alc: alkaloids; Poly: polyphenol

# 3.3Cytotoxicity test

Figure 2 gives the percentage of viability of the HFF cells cultured in the presence of concentrations of 100 to 800  $\mu g$  / mL for the 70% ethanolic extract of the fruits of *Solanumtorvum* compared to the control without plant extract. The number of cells decreases considerably as the concentration of the 70% ethanol extract of the fruits of *Solanum torvum* increases. At 800  $\mu g$  / mL the number of dividing cells is 4%. The averages with the same superscript superscript letters are not different at 5% according to the turkey test.

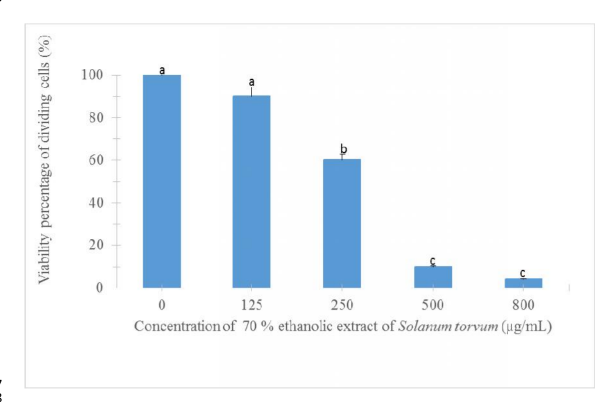


Figure 2: Cytotoxicity test of 70%ethanolic extract of fruits of *Solanum torvum* on HFF dividing cells

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#### 4.Discussion

Medicinal plants play a central role in traditional medicine. Ethnobotanical surveys conducted among traditional health practitioners have made it possible to harvest the fruits of Solanum torvum, which are used in the treatment of an emia, bacterial infections and several other diseases [18]. The recipes obtained from the fruits of this plant are monospecific, which is an advantage for the patients, because the associations of wrongly mixed plants, are sometimes dangerous for the health [19]. Phytochemical screening revealed the presence of chemical compounds such as alkaloids, tannins, polyphenols, saponins and flavonoids. The presence of these chemical compounds could justify the multiple activities of the fruits of this plant [20]. The cytotoxic essay performed on HFF cells showed a gradual decrease in purple staining in each well. Since the dye penetrates only in living cells, the coloring is weaker as the plant extract is cytotoxic by inhibition of HFF cells[21]. The sharp decrease in the relative amount of the dividing HFF cells could be explained by the fact that the HFF cells would be killed by the 70% ethanol extract of Solanum torvum. Indeed, extracts resulting in a cell death greater than 30% could be considered as cytotoxic[22]. This extract could therefore contain a chemical compound that inactivates succinate dehydrogenase, an enzyme important for mitochondrial respiration, the blockage of which would lead to cell death. This result demonstrates the cytotoxic effect of 70% ethanolic extract of Solanum torvum, a Solanaceae from the Ivorian pharmacopoeia on the cell line tested. Which means that the external use of the fruits of this plant would probably be dangerous for human health. This toxicity of fruit could also be explained by the presence of certain groups of chemical compounds such as glycoalcaloides which are toxic in some Solanaceae[13]. Our results on *in vitro* toxicity corroborate those [23] who worked on the same family of plants. Indeed according to the work of Busser and Baies[23]the fruits of Solanum nigrum L. (Solanaceae) another Solanaceae rich in glucoalcaloids and saponins are toxic in internal and external uses on an organism.

#### 5. Conclusion

- 224 Solanum torvum is an important medicinal plant of the family of Solanaceae. From the
- evaluation of the biological activity, it appears that 70% ethanolic extract of the fruits of
- 226 Solanum torvum is cytotoxic on the HFF cells. We therefore plan to continue the acute and
- subacute toxicity studies to compare the results with those obtained in vitro to confirm or
- refute the toxicity of the fruits in this study.

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