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Quilamine HQ1-44, an iron chelator vectorized toward tumor cells by the polyamine transport system, inhibits HCT116 tumor growth without adverse effect.

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Running title: Vectorized iron chelator HQ1-44 inhibits tumor cell growth
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Abstract

Tumor cell growth requires large iron quantities and the deprivation of this metal induced by synthetic metal chelators is therefore an attractive method for limiting the cancer cell proliferation. The antiproliferative effect of the Quilamine HQ1-44, a new iron chelator vectorized toward tumor cells by a polyamine chain, is related to its high selectivity for the Polyamine Transport System (PTS), allowing its preferential uptake by tumoral cells. The difference in PTS activation between healthy cells and tumor cells enables tumor cells to be targeted, whereas the strong dependence of these cells on iron ensures a secondary targeting. Here, we demonstrated in vitro that HQ1-44 inhibits DNA synthesis and cell proliferation of HCT116 cells by modulating the intracellular metabolism of both iron and polyamines. Moreover, in vivo, in xenografted athymic nude mice, we found that HQ1-44 was as effective as cis-platin in reducing HCT116 tumor growth, without its side effects. Furthermore, as suggested by in vitro data, the depletion in exogenous or endogenous polyamines, known to activate the PTS, dramatically enhanced the antitumor efficiency of HQ1-44. These data support the need for further studies to assess the value of HQ1-44 as an adjuvant treatment in cancer.
1- Introduction

1.1- Iron metabolism, a new target for anticancer therapy

Iron is essential in many biological functions and a potential target for anticancer therapy [1]. Initially developed for the treatment of iron overload, iron chelators have shown an antiproliferative action on tumor cells, thus demonstrating their potential value in the treatment of some cancers [2]. Thus, the high efficiency and selectivity of new thiosemicarbazone chelators against various tumors have been reported in vitro and in vivo.[3]. In fact, growing tumor cells require large quantities of iron, leading to an increase in the iron importing proteins and a decrease in the expression of iron exporting proteins [4-6]. In addition, iron exposure stimulates the proliferation of tumor cells in vitro and in vivo [7]. Therefore, decreasing the exchangeable intracellular iron level with synthetic iron chelators appears an attractive method to reduce the proliferation of cancer cells [8]. Our previous results dealing with deferasirox (ICL670), an iron chelator currently used in the treatment of secondary iron overload disease, suggested that its antiproliferative activity results from its inhibitory effects on both iron and polyamine metabolisms [9]. However, these chelators were designed to treat patients exhibiting iron excess. Therefore, their use in patients without iron excess may lead to major side effects, thus limiting their use as antitumoral agents in patients not affected by iron overload. Thus, for potential iron chelators to be used in cancer therapy, they must be vectorized toward tumoral cells.

1.2- Polyamine, a vector for anticancer agents

In tumor cells, polyamine metabolism, including biosynthesis and uptake, is particularly amplified [10, 11]. Targeting polyamine metabolism by using polyamine synthesis inhibitors or polyamine catabolism activators was demonstrated to prevent and reduce risk factors associated with the tumor development [12]. The efficiency of the polyamine transport system (PTS) is also greatly increased in tumoral cells. This property is of potential interest for targeting antitumor agents [13-28].

Based on these observations, we have developed a new generation of iron chelators, the Quilamines, in which an 8-hydroxyquinoline chelating subunit is grafted onto polyamine vectors, in order to vectorize the iron chelator inside the cancerous cell with an overactive PTS [29]. The difference in PTS activation
between healthy and tumor cells enables tumor cells to be targeted whereas the stronger dependence of these latter cells on iron ensures a secondary targeting.

We previously reported that the Quilamine HQ1-44 was highly taken up by the PTS and displayed an efficient antiproliferative activity [29]. As expected during the design of Quilamines, the polyamine chain, involved in the metal coordination, reinforced the affinity for iron (III) and the ferric complex was shown to adopt preferentially a 1:2 [Fe/HQ1-44] stoichiometry at pH=7.4.

We extended the screening of the antiproliferative activity of HQ1-44 to various tumor cell lines and compared it to two non-tumoral cell lines [30]. The relationship between PTS activity and antiproliferative efficiency of HQ1-44 in these various cancer cell lines clearly demonstrated that the Quilamine HQ1-44 was the most efficiently recognized by the active PTS of human colon adenocarcinoma HCT116 cell lines. Consequently, this Quilamine exhibited the higher antiproliferative efficiency on this cell line. In contrast, a low cytostatic effect was observed in normal cells. Colorectal cancer (CRC) is the second most common cause of death from cancer in France and the fourth most commonly diagnosed cancer in the world, so we choose to demonstrate the proof of the Quilamine concept on this kind of cancer.

In the present work, our aim was to investigate the impact of HQ1-44 on cell proliferation and the mechanisms involved as well as assessing its efficiency and tolerance in vivo as an antitumoral treatment in athymic nude mice xenografted with HCT116 cells, by comparison with cis-platin.

2- Materials and Methods

2.1- In vitro study

2.1.1- Chelator solutions

All reagents were obtained from Sigma-Aldrich (Saint Quentin Fallavier, France) at the highest available grade. Stock solutions of the Quilamine HQ1-44 (10 mM) were prepared in water. Cells were supplemented with iron by exposure to 20 µM exogenous iron-citrate complexes (1:10) for 72 h.

2.1.2- Cell studies
The involvement of the PTS in the selective uptake of Quilamine HQ1-44 was investigated in the HCT116 cell line, derived from human colon carcinoma (ATCC number 91091005). We previously demonstrated the higher sensitivity of this cell line to Quilamines, compared to other carcinoma cells [30].

These cells were grown in McCoy’s medium (Life Technologies, Invitrogen™, Saint Aubin, France), supplemented with 10% fetal calf serum (Eurobio, Les Ulis, France), 100 units/mL penicillin and 50 µg/mL streptomycin, at 37°C in 5% humidified CO2.

2.1.3- Cell treatment

For experiments, cells were harvested with trypsin and seeded 24 h before the treatments in microplates (Becton Dickinson, Oxnard, CA, USA) at a density of 10,000 cells/cm². In these conditions, cells reached confluency in 5-6 days. Chelator exposure was performed one day (D1) or one week (D8) after cell seeding in proliferating or confluent HCT116 cells, respectively. The effect of Quilamine on cell viability was also tested in the presence of DFMO (1 mM), a selective inhibitor of ODC, which induced a depletion of intracellular putrescine and spermidine and thus activated the PTS. In this case, preincubation, in the presence of the ODC inhibitor, was performed prior to the Quilamine treatment. In other experiments, 50 µM of spermidine was added to the culture medium to perform a competitive inhibition of Quilamine uptake through the PTS. Aminoguanidine (1 mM), an inhibitor of the serum amine oxidase, was added to the cell culture medium to prevent the oxidation of exogenous polyamine.

For the caspase 3/7 activity assay, Quilamine uptake quantification, polyamine concentration determination and iron status evaluation, cell lysates were obtained after 72 h of treatment. After supernatant collection for biochemical analysis, cells were washed twice with 1 mL of ice-cold phosphate-buffered saline solution (PBS). Cells were collected by scraping in 1 mL PBS, and then centrifuged for 5 min at 2,000 g. The cell pellet was sonicated for 30 min at 4°C in 500 µl of pure water. Protein content in cell extracts was measured according to a method adapted from Bradford (1976) (Bio-Rad Protein Assay, Bio-Rad, Ivry sur Seine, France) and absorbance was read at 595 nm.

2.1.4- Measurement of cytostatic and cytotoxic effects

Treatments was performed one day (D1) or one week (D8) after cell seeding in proliferating or confluent HCT116 cells, respectively. After 72 h of incubation at 37°C, cell supernatants were collected for
cytotoxicity evaluation (cytotoxicity detection kit – LDH, Roche, Penzberg, Germany). Results were reported as a percentage of extracellular LDH activity with respect to the control value.

Cell viability was determined by cell nuclei counting after Hoechst 33342 staining and the dose-effect curves were analyzed according to an established procedure [30]. The number of cell nuclei was reported as a percentage of the value obtained in control conditions. Due to their biphasic feature, the dose-response curves were fitted as the sum of two sigmoids (double 4-parameter fit) according to Rodbard [31]. The various parameters, including the percentage and the IC$_{50}$ values of each sigmoid, were deduced from these fits. Three independent replicates were performed for each experiment, which was repeated three times.

2.1.5- Measurement of DNA synthesis

DNA synthesis was assessed by tritiated thymidine incorporation. Twenty-four hours before cell harvesting, $[^3]$H methyl-thymidine (Amersham, Uppsala, Sweden) was added to the culture medium at a final concentration of 0.5 µCi/mL. Cells were washed twice with PBS and lysed in 0.1 M NaOH. DNA precipitation was achieved with 30% trichloroacetic acid (TCA), followed by two washes with 10% and 5% TCA successively, and dissolution in formic acid. The radioactivity of 200 µl of this formic solution was counted in 5 mL of Instagel using a Packard Tricarb 2100TR scintillation counter β (Perkin Elmer, Waltham, MA, USA) and expressed as cpm/g of protein.

2.1.6- Analysis of the cell cycle

The cell cycle was assessed by flow cytometry. Cells were detached with trypsin and nuclei were stained with propidium iodide (Cycletest Plus Kit, Becton Dickinson, Le Pont de Clais, France). Data were acquired and analyzed with a FACSCalibur flow cytometer (Becton Dickinson) using Cell Quest software (Becton Dickinson).

2.1.7- Measurement of caspase 3/7 activity

Apoptosis was assessed through caspase 3/7 activity quantification, using the “AMC caspase-3/7 assay” kit (AnaSpec, Fremont, CA, USA), in accordance with the manufacturer’s instructions. Caspase 3/7 activity measured by fluorescence at 440/460 nm was expressed as RFU/g of protein and reported as a percentage with respect to untreated control cells.
2.1.8- Measurement of gene expression

Total RNA extracted using “SV total RNA isolation system” kit (Promega, Wisconsin, USA) was tested by spectrophotometrical dosage and electrophoretic separation on a 1.5% agarose gel. RNA was reverse-transcribed to synthesize cDNA using Moloney Murine Leukemia Virus reverse-transcriptase (Promega, Wisconsin, USA). Gene expression was measured by real-time quantitative PCR performed in triplicate on an ABI Prism 7000 Sequence Detection System (Applied Biosystems, London, UK) using the qPCR Mastermix Plus for SYBR Green I kit and specific primer (Eurogentec, Seraing, Belgium) designed by Primer Express 1.0 software (Table 1). The relative quantification of mRNA levels was calculated after normalization using 18S RNA as an endogenous reference. The results were expressed as percentage compared to control culture.

2.1.9- Measurement of HQ1-44 uptake in HCT116 cells

HQ1-44 exposure at 1 µM was performed one day (D1) or one week (D8) after cell seeding in proliferating or confluent HCT116 cells, respectively. Quilamine uptake was also measured after activation of the PTS (1 mM DFMO, see below) or in the presence of 50 µM exogenous spermidine in the culture medium to perform a competitive inhibition of the HQ1-44 uptake by the PTS. Cell lysates were ultracentrifuged for 10 min at 15,000 g and HQ1-44 was quantified in the supernatant by LC/MSMS.

2.1.10- LC/MS quantification of HQ1-44 in HCT116 cells

Analyses were performed on a liquid chromatograph coupled with an ultra-high-resolution and accurate mass instrument (LC-HR-MS), Q ExactiveTM equipped with an Orbitrap mass analyzer (Thermo Scientific, San Jose, USA). An HESI source was used for the ionization of target compounds. Data acquisition, peak integration and calibration were performed using Xcalibur® 2.1 software (Thermo Scientific, San Jose, CA, USA).

LC separation was carried out using a gradient on a C18 Hypersil Gold column (100 mm x 2.1, 3 µm) from Thermo Scientific (San Jose, USA). The mobile phases used were 10 mM ammonium acetate buffer containing 0.1% (v/v) formic acid (solvent A) and methanol/water (95/5, v/v) (solvent B). The mobile phase was delivered at a flow rate of 300 µL/min using the following stepwise gradient elution program: initial conditions of 100% solvent A maintained for 3 minutes, run from 100% solvent B at 2 minutes,
maintained for 1 minute, run from 100% solvent A at 1 minute and conditions of 100% solvent maintained for 5 minutes for equilibration. All prepared samples were kept at 15°C in the autosampler until injection of 5 µL into the LC-HR-MS system (partial loop) in a thermostated column at 25°C.

The MS conditions were as follows: HESI in positive mode, capillary temperature: 300°C; spray voltage: 4500 V; sheath and auxiliary gas (nitrogen) flow-rate: 30 and 15 (arbitrary units), respectively. Data were acquired in full scan mode. Full mass spectra were scanned in the mass range 120 to 700 m/z, the AGC Target was 1e6 and the resolution was 140,000 (FWHM). The measured accurate m/z values of the protonated species was 317.2337. Quantitation was achieved by extracting the exact mass of the protonated species using a 10-ppm extraction window. Quilamine concentrations were then deduced from the calibration curves obtained from HCT116 cell lysate enrichments with various concentrations of HQ1-44, followed by their sonication and ultrafiltration.

Each HQ1-44 measurement was determined in triplicate and was corrected for protein content in the cell extracts.

2.2- In vivo study

2.2.1- Xenografted nude mice model studies

All animal experiments were approved by the Rennes Ethics Committee for Animal Experimentation and by the Ministry for Higher Education and Research. Mice were housed in individually ventilated cages in pathogen-free quarters in the animal handling facility of Rennes 1 University (ARCH-E-Biosit). The 5-week-old athymic female Swiss nude mice (Charles River, Saint Quentin Fallavier, France) were acclimated in a sanitary room for 1 week before the start of the experiments. Viable HCT116 colon cancer cells (3.0 ×10⁶ in medium/100 µl) were injected subcutaneously into the right flank of the nude mice.

2.2.2- Comparison of antitumoral efficiency of HQ1-44 and cis-platin

Ten days after tumor cell inoculation, mice showing a tumor volume between 80-120 mm³ were divided randomly into different treatment groups (6 mice per group) and were treated daily by intraperitoneal injection of Quilamine HQ1-44 at different doses (0, 6.6, 20 and 40 mg.kg⁻¹) or cis-platin (4 mg.kg⁻¹) for 17 days. The mice were observed daily. Mice weight and tumor volume were measured three times per
week with a digital caliper. Tumor volumes were calculated according to the formula for a spheroid: tumor volume = 0.5 x length x width². Growth inhibition was calculated from the start of treatment by comparing the change in tumor volume for vehicle and treated groups.

2.2.3- Effect of activation of PTS on the antitumoral efficiency of HQ1-44

PTS activation was induced by a nutritional deficiency in polyamine (polyamine deficient chow, PDC) as previously described [32] and/or by inhibition of the polyamine biosynthetic pathway with α–difluoromethylornithine (DFMO, 3% (w/w) in the drinking water). This activation was associated or not with the Quilamine treatment.

Ten days after tumor cell inoculation, mice showing a tumor volume between 80-120 mm³ were randomly divided into 6 treatment groups (9 mice per group). Mice received PDC supplemented or not with polyamine (21 mg.kg⁻¹ putrescine.2HCl, 153 mg.kg⁻¹ spermidine.3HCl, 48.7 mg.kg⁻¹ spermine.4HCl) and/or DFMO 3% (w/w) in their drinking water. Quilamine-treated mice groups were treated daily by intraperitoneal injection of Quilamine HQ1-44 at 40 mg.kg⁻¹ for 17 days. The mice were observed daily.

Mice weight and tumor growth were measured three times per week. Growth inhibition was calculated from the start of treatment by comparing the change in tumor volume for vehicle and treated groups.

Mice were sacrificed the day after the last injection. Blood was collected by intra-cardiac puncture after anesthesia. Liver, spleen, heart, kidneys and tumor were collected, weighed, and frozen at -80°C for biochemical and gene expression analyses and/or fixed in formaldehyde for histological analyses.

2.2.4- Blood analysis

Whole blood was collected in heparinized microtubes. After collection of the plasma for biochemical analyses (iron, transferrin, ferritin), 100 µL of red blood cells (RBC) was discarded for polyamine measurement according to a described method [33] [34]. The LC/MSMS analysis of the dansylated polyamines, adapted from a described method [35, 36], was performed with a Thermo Scientific TSQ Quantum Ultra (Thermo Fisher Scientific, Courtabeuf, France).

Levels of intracellular iron were measured by a colorimetric method using OSR6286 kits (Olympus Diagnostics GmbH, Lismeehan, Ireland) on an Olympus AU 2700 chemistry autoanalyzer (Olympus Diagnostics GmbH, Hamburg, Germany).
Ferritin and transferrin were quantified in plasma by ELISA using a mouse-specific assay kit in accordance with the manufacturer’s recommendations (Mouse Ferritin Elisa Kit, Abnova; Mouse Transferrin Elisa Kit, Genway).

2.2.5- Statistical analysis

Results from in vitro studies were replicated three times and expressed as means ± SD. Within the in vivo study, results from each animal group were expressed as means ± SD. Statistical analysis was performed using the non-parametric Mann-Whitney test. The significance level was set at 0.01.

3- RESULTS

3.1- HQ1-44-induced decrease in HCT116 cell growth involves the PTS

The dose-effect curves obtained after treatment of proliferating HCT116 cells (D1) with HQ1-44, could be deconvoluted into two components (Figure 1A). For Quilamine concentrations lower than 10 µM, the first component indicated the antiproliferative effect of the Quilamine (62% of component 1 with IC\textsubscript{50-1} = 1 µM). For higher concentrations, the second component reflected the cytotoxic effect (38% of component 2 with IC\textsubscript{50-2} = 165 µM), linked to membrane damage and a release of lactate dehydrogenase (LDH) into the supernatant (Figure 1B). In contrast, when confluent cells, eight days after cell seeding (D8), were treated with HQ1-44, the dose-effect curve was reduced to one component (IC\textsubscript{50} = 210 µM, Figure 1A), associated with LDH release into the supernatant (Figure 1B).

To explore the involvement of the PTS in HQ1-44 vectorization and consequently its antiproliferative effect, we used two approaches. Firstly, PTS activation was induced by a treatment with α-difluoromethylornithine, DFMO (1 mM), an inhibitor of ornithine decarboxylase (ODC), leading to polyamine depletion and an increase in PTS expression. This activation improved the antiproliferative effect of HQ1-44 at D1, as shown by the increase in the first component (70%) and the decrease in the IC\textsubscript{50-1} value (0.5 µM) (Figure 1A). In contrast, co-incubation of the HCT116 cells in the presence of 50 µM spermidine, a natural polyamine, at D1 led to a competitive inhibition of HQ1-44 uptake by the PTS and thus an inhibition of its antiproliferative effect (60% of cytostatic component 1 with IC\textsubscript{50-1} = 3.5 µM).
Secondly, the HQ1-44 uptake measured in cell lysates by LC/MSMS was six-fold higher in proliferating (D1) than in confluent (D8) HCT116 cells (Figure 1C), and the activation of the PTS by DFMO treatment in proliferating cells led to a two-fold increase in Quilamine uptake. In contrast, co-incubation of proliferating cells with exogenous spermidine (50 µM) competitively inhibited the Quilamine uptake by the PTS. In confluent cells, both DFMO and spermidine remained ineffective in activating or inhibiting, respectively, the Quilamine uptake.

### 3.2- HQ1-44 reduces DNA synthesis, stopping the cell cycle and inducing apoptosis

HCT116 cells were seeded at a cellular density allowing their analysis in the proliferative phase (10,000 cells/cm$^2$). In the untreated control cells, $^3$H-thymidine incorporation, evaluating DNA synthesis, increased with time 24 h and 48 h after cell seeding and reached a plateau at 72 h (Figure 2A). After 72 h, cells reached confluency and DNA synthesis was slowed down. In the HCT116 proliferating cells treated with a concentration of HQ1-44 close to its IC$_{50}$ value (2 µM), DNA synthesis was weakly inhibited compared to the control after 48 h incubation and, to a lesser extent, at 72 h (45% of inhibition).

Analysis by flow cytometry of the percentage of cells in the various phases of the cell cycle in untreated control cells at different growth times after cell seeding (24, 48 and 72 h) showed a progressive reduction of cells in phases G2/M and S with time, whereas the number of cells in phase G0/G1 increased (Figure 2B). Cell culture treatment with HQ1-44, at a concentration close to its IC$_{50}$ value (2 µM), led to an arrest of the cell cycle after 72 h of treatment, resulting in an increase in the percentage of cells in phases G2/M (13.6% versus 5.0% in the control), S (31.9% versus 11.3% in the control) and a drop in those in phase G0/G1 (54.6% versus 83.6% in the control). No significant apoptosis was detected by the flow cytometry analysis at this HQ1-44 concentration.

Under the same conditions, these data were confirmed by in situ analyses of the size and shape of cell nuclei after DNA staining, showing the drop in the percentage of viable cells with a small nucleus (G0/G1 phase) and the relative increase in the percentage of cells with a 50% bigger nucleus, characteristic of cells in G2/M (Figure 3C).
For concentrations lower than 5µM that are cytostatic but not cytotoxic, HQ1-44 was ineffective at inducing caspase 3/7 activity (Figure 2C), an early apoptosis marker, while for cytotoxic concentrations (e.g. up to 5 µM) HQ1-44 triggered a slight, but significant, increase in caspase 3/7 (+10%). The 20% increase in caspase 3/7 induced by 10 µM HQ1-44 remained lower than the effect of 50 µM hydrogen peroxide, used as a reference apoptosis inducer (+45%).

At sub-toxic HQ1-44 concentrations (<5 µM), an in situ analysis of nuclei after DNA staining showed no significant DNA fragmentation, which could reflect apoptosis (Figure 3A), thus supporting the caspase 3/7 activity data. In contrast, some fragmented nuclei were observed after HQ1-44 treatment with cytotoxic concentrations (10µM, Figure 3B).

### 3.3- HQ1-44 affects iron and polyamine metabolisms in HCT116 cells

Treatment with 20 µM iron-citrate induced a marked increase in the level of the L-ferritin protein subunit, and a decrease in the soluble transferrin receptor form in cell supernatants (Figure 4A). In contrast, treatment with the reference iron chelator, ICL670, at a concentration close to its IC$_{50}$ (8 µM, data not shown) caused an iron deficiency characterized by a reduction in the L-ferritin level and an increase in the level of the soluble transferrin receptor, also observed to a lesser extent at 2 µM (Figure 4A). At this concentration, HQ1-44 induced a larger reduction in the ferritin level and a greater increase in the level of soluble transferrin receptor 1 than ICL670, confirming a situation of iron deficiency.

Finally, an analysis of the TFRc mRNA, encoding the transferrin receptor 1 protein, showed a down-regulation of the expression of this gene in iron overload conditions (20 µM iron-citrate) while an up-regulation was observed in iron depletion conditions induced by both chelators, ICL670 and HQ1-44, at 2 µM (Figure 4B).

Polyamine metabolism was also affected by HQ1-44. The addition of 2 µM of HQ1-44 to the HCT116 cell culture medium induced a significant reduction in intracellular levels of putrescine and spermidine while the spermine level remained unchanged (Figure 4C). In the same way, a reduction in the level of acetylated spermine (ASpm) was found but with no effect on the acetylated spermidine level (data not shown). Under the same experimental conditions, ICL670 remained ineffective at modulating the polyamine level within HCT116 cells whereas it induced an inhibition of polyamine metabolism in rat and human hepatoma cells.
at higher concentrations [9]. These data suggest that HQ1-44 may induce a disturbance both in the polyamine biosynthetic pathway, by inhibiting the synthesis of putrescine and spermidine, and in their retro-conversion, by inhibiting the level of acetylated spermidine (Fig. 4C). Analysis of the mRNA levels of genes involved in polyamine biosynthesis, including ornithine decarboxylase (ODC), antizyme (OAZ1), S-adenosylmethionine decarboxylase (SamDC) and polyamine oxidase (PAO), showed a significant down-regulation of their expression (Figure 4D).

3.4- HQ1-44 inhibits tumor growth in vivo

To test the antiproliferative efficiency of HQ1-44 in vivo, we treated HCT116 xenografted athymic nude mice by i.p. injection with increasing doses of Quilamine (Figure 5, Table 2). After a treatment of 15 days, analysis of the tumor weights showed a non-significant reduction for doses of 6.6 and 20 mg/kg and a significant reduction for the dose of 40 mg/kg (-38%). Under the same conditions, cis-platin (4 mg/kg) induced a reduction of 29% of the tumor weight. As expected, the treatment with cis-platin also involved a marked reduction in the weight of the animal’s body, liver and spleen (-17%, -17% and -44%, respectively) highlighting the toxicity of this compound. Histological analysis of the spleen (Figure 6) showed that the 44% drop in spleen weight in animals treated with cis-platin was mainly due to a decrease in the red pulp of this organ (36% decrease with respect to the control), as suggested by its paler red color.

It is noteworthy that cis-platin administration also induced a behavioral change and reduced activity in mice, while no such changes were observed in Quilamine-treated mice.

By contrast, treatment with the efficient dose of HQ1-44 (40 mg/kg) did not affect the animal and organ weights, or behavior, thus suggesting a better tolerance of the Quilamine at this concentration.

Investigating the impact of HQ1-44 on different iron and polyamine parameters, we found that at doses of 6.6 and 20 mg/kg, the Quilamine did not significantly affect the plasmatic iron concentrations, whereas a significant decrease in iron level was observed for a dose of 40 mg/kg (-21%) (Table 2). By contrast, cis-platin had no effect on iron parameters. Finally, neither treatment (HQ1-44 and cis-platin) affected the intraerythrocytic concentration of polyamines.

3.5- Polyamine deprivation increases the inhibitory effect of HQ1-44 on tumor growth
As the *in vitro* data suggested an additive impact of DFMO with Quilamine, we investigated whether PTS activation might potentialize the Quilamine impact in our mouse model. Thus, polyamine deprivation was carried out by combining a blockade of endogenous sources, inhibition of intracellular polyamine biosynthesis with DFMO [37] and/or a reduction in exogenous sources by nutritional deficiency in polyamine with a polyamine deficient chow (PDC) [32, 38]. As shown in Table 3, DFMO treatment alone, associated to a nutritional deficiency in polyamine (PDC) or not (PA), inhibits tumor growth, leading to a 33% and 35% tumor growth inhibition for PA and PDC animal groups, respectively. Our results showed that polyamine deprivation was efficient in reducing tumor development. The monitoring of tumor size under the different treatments is summarized in Figure 7. After 15 days of treatment, tumour reduction induced by HQ1-44 (-30%) was amplified in the group receiving 3% DFMO in their drinking water (-56%). Polyamine deprivation in the group of animals receiving the PDC, in the absence of any other treatment, had no impact on tumor growth while the PDC diet increased the tumor growth inhibitory effect of the combined treatment with HQ1-44 and DFMO (-65% of tumor growth inhibition). The various treatments did not significantly modify the animal and organ weights, except for DFMO associated with HQ1-44 treatment, which induced a significant reduction in the spleen weights (Table 3). The decrease in spleen weight in animals receiving this combined treatment is low (-16% with respect to the control, Table 3) compared to the spleen toxicity of the treatment with 4 mg/kg *cis*-platin (-44%, Table 2). Contrary to the treatment by this alkylating agent, the histological analysis of the spleen after the combined treatment with PDC, DFMO and HQ1-44 did not show any significant decrease in the red pulp of this organ.

A significant decrease in circulating iron in serum was observed in the group of animals receiving the Quilamine in conjunction with the polyamine-free diet (Table 4). No change in the ferritin and transferrin levels was detected in these groups while an unexpected decrease in the serum ferritin level was observed in the group of untreated animals receiving the polyamine-free diet (PDC). The various treatments did not modify the polyamine levels in RBC (Table 4).

### 4- Discussion
Our objective was to investigate whether the vectorized iron chelator HQ1-44 might be of value in the treatment of cancer. Our data demonstrate in vivo that HQ1-44 is effective, by limiting tumor growth of HCT116 cells in xenografted nude mice. The antitumor efficiency of the Quilamine HQ1-44 is similar to that of the reference alkylating antitumor agent, cis-platin, and is not associated with toxicity, unlike the reference treatment. In addition, both in vivo and in vitro, the results obtained are helpful in understanding the mechanisms involved, suggesting an impact of HQ1-44 on the cell cycle through an alteration of both iron and polyamine metabolisms.

The decrease in the intracellular ferritin protein level and the increase in soluble transferrin receptor concentration in culture medium in the presence of HQ1-44 clearly show that HQ1-44 induces a reduction in the intracellular iron pool in tumor cells, which in turn regulates the expression of these two genes through the Iron Responsive Element/Iron Regulatory Protein (IRE/IRP) system. These data are in accordance with those we previously reported, demonstrating the iron-chelating activity of Quilamines, including HQ1-44. In HQ1-44, the iron-chelating activity of the chelating motif is reinforced by the polyamine backbone, with both the 8-HQ moiety and the polyamine part participating in the iron coordination [29]. In addition, a role of the amino groups of polyamines in the chelation of metallic cations and in the stability of iron chelates has previously been reported [39, 40]. Such a high capacity for chelating iron is likely to participate in the strong antiproliferative activity, as demonstrated in a previous study, showing that the higher antiproliferative activity of the bis-8-hydroxyquinoline benzylamine chelator S1 in hepatocytes, compared to the iron chelator o-Trensox, was closely connected to the high iron-chelating capacity of this tetradeutate ligand [41]. For low concentrations of chelating agents (2 µM), the higher iron-chelating capacity of Quilamine HQ1-44, compared to deferasirox, may explain the stronger impact on L-ferritin and transferrin receptor levels, through the IRE/IRP system.

It is noteworthy that in vivo data showing that the serum iron level is only slightly affected with the highest HQ1-44 dose, without a significant change in both systemic ferritin and soluble transferrin receptor levels, support the concept that HQ1-44 is an iron chelator targeting tumoral cells. As reported above, such a concept has been proposed to target other molecules. Our in vivo data, with a low level of side effects, reinforce those obtained in vitro showing the higher sensitivity of proliferating HCT116 cells compared to
confluent cells, as well as those previously reported, showing that cells not expressing the PTS are less sensitive to HQ1-44 [29]. In addition, the HQ1-44 targeting can be reinforced through polyamine metabolism manipulation using polyamine depletion tools by providing animals with a polyamine-deficient diet and DFMO, an inhibitor of polyamine biosynthesis, in their drinking water. The polyamine depletion induced by these experimental conditions are currently believed to activate the PTS in tumor cells, as a feedback regulation to maintain the higher polyamine levels required for tumor growth [42].

In addition to its impact on intracellular iron metabolism, HQ1-44 could also exert its antiproliferative activity by modulating polyamine metabolism. Indeed, polyamines are involved in the control of cell cycle progression and cell proliferation [11]. Thus, the ubiquitous polyamine molecules modulate the G1/S transition phase [11]. Our results show that the Quilamine HQ1-44, like deferasirox at higher concentrations, disturbs the metabolism of natural polyamines by inhibiting their biosynthetic pathway. HQ1-44 reduces the gene expression of the metabolism of polyamines, such as those of ornithine decarboxylase (ODC), antizyme (OAZ1) and S-adenosylmethionine decarboxylase (SamDC). This leads to a significant decrease in putrescine, spermidine and a slight, non-significant, reduction in spermine. Such findings are characteristic of a biosynthetic pathway inhibition, such as that of ODC by DFMO. HQ1-44 also modulates the retro-conversion oxidative pathway of polyamines, causing inhibition of the production of N1-acetyl spermine and increased levels of N1-acetyl spermidine. The contribution of the inhibitory effect of chelating iron, such as by Quilamine HQ1-44, to the metabolism of polyamines and their impact on cell proliferation is under study.

In conclusion, HQ1-44 Quilamine is effective in inhibiting HCT116 in vitro and in vivo in xenografted mice. This effect is related to the iron chelator impact of the iron chelator motif, which is reinforced by the polyamine backbone. An inhibitory impact of HQ1-44 on polyamine metabolism may also play a role by affecting cell cycle progression. The polyamine backbone of the molecule enables tumor cells to be targeted and limits the toxicity and side effects of the molecule. Taken together, these results suggest that Quilamines are a way of exploring and adding to the cancer therapeutic panoply in association with other treatments.
5- Acknowledgments

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The histological analysis were carried out in the histopathological platform H2P2 of the Structure Fédérative de Recherche BIOSIT in Rennes (BiogenOuest and Cancéropôle Grand Ouest network).

6- References


[40] Palmer BN, Powell HKJ. Polyamine complexes with seven-membered chelate rings: Complete formation of 3-azaheptane-1,7-diamine, 4-azaoctane-1,8-diamine (spermidine), and 4,9-diazadodecane-1,12-diamine (spermine) with copper(II) and hydrogen ions in aqueous solution. JCS Dalton. 1974:2089-92.


7- Legends of tables and figures

Table 1: Primer sequences.

Table 2: Comparison of the antitumor efficiency of Quilamine HQ1-44 and cis-platin in HCT116 xenografted immunocompromised nude mice. Effect on animal, organ and tumor weights and on blood levels of iron and polyamine parameters. Results from each animal group were expressed as means ± SD from three independent measurements. Statistical analysis was performed using the non-parametric Mann-Whitney test (* p≤0.05, ** p≤0.01).

Table 3: Effect of polyamine depletion on the antitumor efficiency of HQ1-44 in HCT116 xenografted immunocompromised nude mice. Effect on animal, organ and tumor weights. Results from each animal
group were expressed as means ± SD from three independent measurements. Statistical analysis was performed using the non-parametric Mann-Whitney test (* p≤0.05, ** p≤0.01).

**Table 4:** Effect of polyamine depletion on the antitumor efficiency of HQ1-44 in HCT116 xenografted immunocompromised nude mice. Effect on blood levels of iron and polyamine parameters. Results from each animal group were expressed as means ± SD from three independent measurements. Statistical analysis was performed using the non-parametric Mann-Whitney test (* p≤0.05, ** p≤0.01).

**Figure 1:** Relationship between PTS efficiency and the antiproliferative effect of HQ1-44 in HCT116 cell cultures.

A- The effect on cell viability was determined in proliferating cells (treatment at day 1, D1) and in confluent cells (-○-, treatment at day 8, D8) by measuring DNA content after Hoescht staining and fluorescence analysis. Dose-effect curves of Quilamine HQ1-44 (-○-), in the presence of DFMO (2 mM) (-△-) and spermidine (50 µM) (-▲-) in proliferating HCT116 cells.

B- Cytotoxic effects of HQ1-44 in proliferating (-○-) treatment at day 1, D1) and confluent HCT116 cells (-○-, treatment at day 8, D8). Membrane damage was deduced from lactate dehydrogenase activity in cell supernatants, collected after 72 h of incubation.

C- Effect of activation (2 mM DFMO) or inhibition (exogenous spermidine 50 µM) of the PTS uptake of Quilamine HQ1-44 (1 µM) in proliferating (grey bars) and confluent HCT116 cultured cells (black bars). Results were expressed as means ± SD from three independent measurements. * means p≤0.01 between treated and control cells.

**Figure 2:** Effect of HQ1-44 on DNA synthesis, cell cycle and apoptosis in proliferating HCT116 cultured cells.

(A) Effect of HQ1-44 (2 µM) on DNA synthesis measured by ³H-thymidine uptake (24 h incorporation).

(white bars, untreated control cells; grey bars, cells treated with 2 µM HQ1-44)
(B) Effect of HQ1-44 (2µM) on the cell cycle analyzed by FACS after propidium iodide staining. The percentage of cells in each phase of the cell cycle was deduced by using Cell Quest software. White bars; G0/G1 phase; grey bars; S phase, black bars; G2/M phase.

(C) Effect of HQ1-44 on apoptosis analyzed by measuring caspase 3/7 activity. 50 µM hydrogen peroxide treatment was used as a reference apoptosis inducer.

Figure 3: Effect of HQ1-44 on cell nuclei of HCT116 cells (shape and size). The impact of HQ1-44 was analyzed by Hoescht staining. (A) Morphology of nuclei of HCT116 cells treated with 2 µM HQ1-44 for 72h. (B) Morphology of nuclei of HCT116 cells treated with 10 µM HQ1-44 for 72 h Fragmented nuclei (FN).

(C) Analysis of the mean size of nuclei of HCT116 cells treated with increasing concentrations of HQ1-44 for 72 h.

Figure 4: Effect of a 72 h treatment of proliferating HCT116 cells with HQ1-44 (2 µM), ICL670 (2 µM) and iron-citrate (20 µM) on iron (A and B) and polyamine (C and D) metabolisms. The effect on both iron and polyamine metabolisms of the reference chelator ICL670 (2 µM deferasirox), which was previously shown to induce an iron deficiency, was compared to that of 2 µM HQ1-44 and to 20 µM iron-citrate (mimicking the situation of iron overload).

A- The soluble transferrin receptor1 (white histogram,□) in cell supernatants and the intracellular ferritin level (black histogram,■) were measured in proliferating HCT116 cells after 72 hours of treatment with the various iron effectors. These two parameters were corrected from total protein concentrations.

B- Gene expression of the transferrin receptor1 (white histogram, □) and ferritin (black histogram, ■) was analyzed by RT-qPCR in proliferating HCT116 cells after 72 h of treatment.

C- Intracellular polyamine levels (putrescine, □; spermidine, □; spermine, ■; and N1-acetyl spermine, ■) after treatment with the various iron effectors. These two parameters were corrected from total protein concentrations.

D- Gene expression of ornithine decarboxylase (□, ODC), antizyme (□, OAZ1), S-adenosylmethionine decarboxylase (■, SamDC) and polyamine oxidase (■, PAO), which were analyzed by RT-qPCR.

* significant differences according to the Mann-Whitney test (p<0.01).
Figure 5: Antitumor effect of HQ1-44 compared to cis-platin in HCT116 xenografted athymic nude mice. Tumor weight after 15 days intraperitoneal administration of Quilamine (0 to 40 mg/kg) and cis-platin (4 mg/kg). * p<0.01.

Figure 6: Histological analysis of spleen. Hematoxyline eosine saffron (HES) staining from HCT116-xenografted nude animals (A) untreated animals, (B) mice treated by the intraperitoneal route for a couple of weeks with 40 mg/kg HQ1-44 (B), and (C) mice treated by 4 mg/kg cis-platin (C). Red pulp (RP), White pulp (WP), Trabecule (T) and germinative center (GC) structures are indicated by arrows.

Figure 7: Effect of polyamine depletion induced by DFMO and/or by a polyamine-deficient diet (PDC) on the antitumor effect of the Quilamine HQ1-44 in HCT116 xenografted immunocompromised nude mice. Tumor growth curves in animals treated with polyamine containing diet (PA, - - - ◆---), polyamine deficient chow (PDC, −◇−), PA and 40mg/kg HQ1-44 (−■−), PDC and 40 mg/kg HQ1-44 (−□−), PA+3%DFMO and 40 mg/kg HQ1-44 (−▲−), PDC+3%DFMO and 40 mg/kg HQ1-44 (−△−)
Polyamine (PA) biosynthesis and uptake via the polyamine transport system (PTS) are overexpressed in tumor cells. HQ1-44 targeting via the PTS can be reinforced by polyamine depletion (polyamine-deficient diet and DFMO, an inhibitor of polyamine biosynthesis).
18S:
Rev: 5'-TCGAGGCCCTGTAATTGGAA-3'
Fw: 5'-CCCAAGATCCAATACGAGCTTT-3'

TfR1:
Rev: 5’-GCTTTCCCTTGCATATTCTT-3’
Fw: 5’-GTGGTACCAAAATAAGGATAATCTGT-3’

L-ferritine:
Rev: 5’-AGGGCCCTTGCAAATTTAA-3’
Fw: 5’-GCCTATTTGCTGGAGGAG-3’

PAO:
Rev: 5’-GGTAGGCTGGAGCCGTCATT-3’
Fw: 5’-AAGGTGCCCCCTGGTTACCA-3’

OAZ1:
Rev: 5’-AATTCCGCGAAAAAGAGAAG-3’
Fw: 5’-CTCCTAGGCCCCTCTGGGTAGTT-3’

SAMDC:
Rev: 5’-TCGTCTTTCCGAGTAAGACCTAT-3’
Fw: 5’-CTTTGCAGTAACACCACCATCTGGATG-3’

ODC:
Rev: 5’-CGGTACAGCCGCTTCTACA-3’
Fw: 5’-CGGTGCACGCTCAGAAG-3’

Table 1
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<th>Organ weight</th>
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<th>HQ1-44 6.6 mg/kg</th>
<th>HQ1-44 20 mg/kg</th>
<th>HQ1-44 40 mg/kg</th>
<th>CisPt 4 mg/kg</th>
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<td>24.7 ±1.5</td>
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<td>24.6±2.7</td>
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<td>Tumor (mg)</td>
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<td>511 ±213</td>
<td>567 ±197</td>
<td>381 ±45**</td>
<td>439 ±53*</td>
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<td>Spleen (mg)</td>
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<td>Kidney (mg)</td>
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<td>Liver (mg)</td>
<td>1347 ±97</td>
<td>1181 ±65</td>
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<td>1321 ±180</td>
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<td>Heart (mg)</td>
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<td>Iron (µM)</td>
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<td>Ferritin (µg/l)</td>
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<td>1775±453</td>
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<td>Transferrin (g/l)</td>
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<td>Polyamines (nM)</td>
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<tr>
<td>Putrescine</td>
<td>133±31</td>
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<td>102±23</td>
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<td>3545±579</td>
<td>6186±887</td>
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Table 2
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<td>PA</td>
<td>25.6±2</td>
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<td>PDC</td>
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<td>390±21</td>
<td>1550±98</td>
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<td>382±25</td>
<td>1344±67</td>
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Table 3
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<td>PDC+DFMO</td>
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<td>PA+HQ1-44</td>
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<td>PDC+HQ1-44</td>
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<td>PDC+HQ1-44+DFMO</td>
<td>28,94±6,62**</td>
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</table>

Table 4
Figure 1
Figure 2

A) \( ^3H \cdot dT \) incorporation (cpm/g) over incubation time (hours).

B) Cell percentage in the cell cycle phase over incubation time (hours).

C) % caspase 3/7 activity/control over HQ1-44 and \( H_2O_2 \) concentrations.
Figure 3
Figure 4
Figure 5
Figure 6
Figure 7