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ARTICLE

Mitochondria-Targeting Symmetric Diiminoguanidines: Potent and Selective Anticancer Agents Against Pancreatic Tumors

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Sigrid Lacaille*a and Andreea R. Schmitzer*a

Pancreatic cancer remains one of the deadliest malignancies of the 21st century, with a five-year survival rate below 12%. Its growing incidence is strongly linked to modern lifestyles, marked by obesity, diabetes, overnutrition, and physical inactivity. Current chemotherapies offer limited success and are often burdened by severe side effects, highlighting the urgent need for more effective and selective treatments. In response, we have developed a new class of easily synthesized, amphiphilic symmetric diiminoguanidines and evaluated their antiproliferative activity against pancreatic cancer cell lines. Several compounds demonstrated remarkable efficacy and selectivity, positioning them as strong candidates for further *in vivo* evaluation. Fluorescence microscopy revealed that these molecules rapidly localize into mitochondria. Preliminary mechanistic studies suggest their primary target is the mitochondrial respiratory chain. These findings support the potential of diiminoguanidines as affordable, mitochondria-targeting alternatives to existing pancreatic cancer therapies.

Introduction

Pancreatic cancer stands as one of the most aggressive and fatal malignancies in modern oncology. ¹⁻³ With a dismal five-year survival rate of less than 12%, it is projected to become the second leading cause of cancer-related deaths in the United States by 2040, surpassing colorectal cancer and second only to lung cancer. ³⁻⁴ This alarming trend reflects both the increasing incidence of the disease and the lack of effective therapeutic interventions. ⁵⁻⁶

The rising prevalence of pancreatic cancer in Western societies is closely tied to metabolic and lifestyle-associated risk factors. Type 2 diabetes, obesity, overnutrition, and physical inactivity have been consistently identified as key contributors to pancreatic tumorigenesis.⁷⁻¹¹ These conditions, highly prevalent in North America and Europe, create a metabolic and inflammatory environment that fosters cancer development and progression. One of the principal challenges in treating pancreatic cancer is the latestage diagnosis in most patients. Early clinical signs are either vague or absent, often mistaken for benign gastrointestinal disorders. By the time a definitive diagnosis is made, the disease is frequently at

Chemotherapy remains the cornerstone of treatment for unresectable or metastatic pancreatic cancer. However, current regimens offer only modest improvements in survival and are associated with significant toxicity. Antimetabolites such as 5fluorouracil (5-FU) and capecitabine disrupt RNA and DNA synthesis but are burdened with adverse effects including myelosuppression, neurotoxicity, and gastrointestinal damage. 12-17 Combination therapies like FOLFIRINOX (5-FU, folinic acid, irinotecan, and oxaliplatin) improve efficacy but also amplify systemic toxicity, often limiting their use to patients with good performance status. 18-20 Gemcitabine, a nucleoside analog, remains the standard of care in many settings, yet typically extends survival by only a few months.²¹⁻ ²⁵ These clinical limitations underscore the urgent need for novel therapeutic strategies that are both efficacious and well-tolerated. Mitochondria, the powerhouses of the cell, have emerged as promising targets in cancer therapy due to their essential roles in bioenergetics, apoptosis regulation, and reactive oxygen species (ROS) generation.²⁶⁻²⁸ Compounds that selectively disrupt mitochondrial function in cancer cells could represent a powerful new avenue for treatment, particularly in metabolically active and chemoresistant cancers such as pancreatic adenocarcinoma. 29-32

an advanced or metastatic stage, leaving few curative options. Surgical resection, while potentially curative, is only feasible in a minority of cases due to the pancreas's deep retroperitoneal location, rich vascular supply, and the high metastatic potential of pancreatic tumors.¹¹

^{a.} Département de chimie- Faculté des arts et des sciences, Université de Montréal. Campus MIL, 1375, Ave. Thérèse Lavoie-Roux, Montréal, Québec, H2V 0B3, Canada. *ar.schmitzer@umontreal.ca.

b. Address here.

c. Address here.

[†] Footnotes relating to the title and/or authors should appear here.
Supplementary Information available: [details of any supplementary information available should be included here]. See DOI: 10.1039/x0xx00000x

3f: R=5-pyrimidine

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Over the past decade, our group has concentrated on the development of guanidine-based molecules, particularly biguanides, as promising agents for targeting pancreatic cancer cells (**Fig. 1**). We previously demonstrated that amphiphilic biguanide salts, most notably a triflate salt of phenylethynylbenzyl (PEB)-substituted biguanide, exhibited potent antiproliferative activity, with IC₅₀ values in the low micromolar range against pancreatic cancer cell lines.³³⁻³⁴ Building on these findings, we further explored bifunctional constructs, including biguanide-based PROTACs and AUTACs, which induced mitochondrial accumulation and led to marked growth inhibition.³⁵⁻³⁶ These efforts validated the guanidine scaffold as a versatile and bioactive platform in anticancer drug discovery.

Fig.1 Structure of antiproliferative PEB-biguanide, biguanide-PROTAC and biguanide-AUTAC synthesized in our group.

Encouraged by these results, we hypothesized that another class of guanidine derivatives, iminoguanidines, could serve as valuable building blocks for the design of new amphiphilic molecules with anticancer potential. Like biguanides, iminoguanidines possess basic nitrogen atoms capable of protonation and hydrogen bonding, key mitochondrial targeting and intracellular accumulation.³⁷ Moreover, their synthetic accessibility enables rapid generation of structural analogs, facilitating structure-activity relationship (SAR) exploration. In this study, we report the synthesis of a library of amphiphilic symmetric diiminoguanidines, prepared as chloride salts, bearing a range of aromatic, polycyclic aromatic, and heterocyclic substituents. These compounds were evaluated for their antiproliferative activity on pancreatic cancer cell lines. Selectivity toward cancer versus normal cells was assessed, and mechanistic studies were conducted to elucidate their mode of action, with a particular focus on mitochondrial targeting.

Results and discussion

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Design and synthesis of amphiphilic symmetric diiminoguanidines as chloride salts

A series of 14 symmetric diiminoguanidines were synthesized as chloride salts (Fig. 2) in order to explore the relationship between molecular structure, specifically side-chain identity and antiproliferative activity against pancreatic cancer cells. Amphiphilicity was a critical design parameter, as it strongly

influences a molecule's ability to traverse cell membranes and reach amphiphilicity and molecular recognition features, we introduced aromatic rings (both substituted and unsubstituted), polycyclic aromatic systems, and heterocycles as side chains. Aromatic moieties are prevalent in medicinal chemistry due to their capacity for π - π stacking interactions, while polycyclic aromatics offer advantageous photophysical properties for intracellular tracking. Heterocycles, on the other hand, are biocompatible and capable of hydrogen bonding, enhancing their potential to interact with biological targets. The diiminoguanidines were obtained via double condensation of N,N'-diaminoguanidinium hydrochloride with two equivalents of the corresponding aldehydes in methanol, under mild conditions (room temperature to 80 °C, ≤24 h) (Fig. 2). It should be noted that certain compounds were obtained with low yields. In the case of the polycyclic aromatic compounds, the main reason was the low solubility of the final substituted diiminoguanidine hydrochloride in methanol. In the case of the heterocyclic compounds, even higher temperatures or longer reaction times did not allow the increase of the yields.

Generally, this synthetic route is operationally simple, efficient, and adheres to green chemistry principles, an important consideration in the development of sustainable therapeutic candidates.

Fig.2 Structure of the synthesized symmetric diiminoguanidines

Antiproliferative activity of the diiminoguanidine hydrochlorides

antiproliferative properties of the synthesized diiminoguanidines (were evaluated on two human pancreatic cancer cell lines: KP4, which is characterized by a high metastatic potential, and PANC1, associated with a comparatively more moderate phenotype. As shown in Table 1, all compounds exhibited IC50 (concentration that inhibit cancer cell growth by 50%) in the micromolar range, validating the iminoguanidine scaffold as a promising starting point for the development of anticancer agents. Remarkably, compound **3b**, bearing imidazole side chains, demonstrated the most potent activity with an IC₅o of 280 nM, while **2c**, bearing pyrene groups, was the least active (IC₅₀ = 163 μ M). SAR analysis revealed several trends:

• Heterocyclic side chains (**3a-3f**) generally conferred higher potency. Imidazole- and oxazole-bearing compounds showed IC₅₀ values well below 5 μM. The exception was **3f** (pyrimidine), which showed only modest activity (IC₅₀ = 119 μM), suggesting that not all heterocycles contribute equally to activity.

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- Polycyclic aromatic derivatives (2a-2c), such as those containing pyrene (2c) or phenanthracene (2b) moieties, were the least active. Their high lipophilicity may hinder cell membrane penetration or intracellular distribution.
- Phenylic derivatives (1a-1e) showed variable activity with no clear correlation to electronic effects. For example, the unsubstituted phenyl derivative 1a and its para-fluorinated analog 1b exhibited similar activity. Interestingly, a metasubstituted derivative bearing electron-donating methyl groups (1d) yielded an IC₅₀ of 29 μM, whereas its electron-withdrawing trifluoromethylphenyl counterpart (1e) had a significantly reduced activity (IC₅₀ = 134 μM). This observation points to a complex interplay of steric and electronic factors not solely explained by traditional SAR paradigms.

As imine bonds are known to have the ability to be hydrolyzed in acidic aqueous media, we investigated the stability of our compounds. To do so, we mimicked the biological experimental conditions, namely PBS buffer with 4% DMSO at 37°C for 3 days, and recorded the UV-Vis spectrum every 24 hours. All the recorded spectra were identical, confirming no deterioration of the diiminoguanidine structure for the duration of treatment (see Supporting information).

Role of Amphiphilicity (clogP) in Antiproliferative Activity Article Online

To further investigate the impact of amphiphilicity/อก/ชางเอเซล performance, we analyzed the correlation between the calculated logP (clogP) values and IC50 data (Fig. 3). The results revealed a compelling inverse relationship between amphiphilicity and antiproliferative efficacy. Compounds with clogP values between 0.5 and 4 consistently showed strong activity (IC₅₀ < 50 µM), indicating an optimal amphiphilic window that supports both membrane permeability and intracellular distribution. In contrast, compounds with clogP values >4 exhibited significantly reduced potency (IC₅₀ > 100 μM), likely due to excessive lipophilicity that hinders effective cellular uptake. Compound 3f, with a clogP of -0.42, displayed weak activity (IC₅₀ = 119 μM), suggesting that higher hydrophilicity may also be detrimental. However, the high activity of 3b despite its higher hydrophilicity suggests that this is not the only parameter that can explain the cytotoxicity of the studied diiminoguanidines. Overall, these results emphasize the necessity of fine-tuning amphiphilicity of the compounds to achieve an optimal balance for membrane penetration, intracellular targeting, and pharmacological activity. Nevertheless, only a deep study of the mechanism of action and the determination of the precise target of diiminoguanidines in the cells would allow to establish the relationship between the nature of the substituents and their cytotoxic activity.

Table 1 IC₅₀ values of diiminoguanidines on pancreatic cancer cell lines KP4 and PANC1, on healthy cell lines IMR90 and *hTERT*-HPNE and selectivity indexes

Compound	Substituent R	IC ₅₀ cancer cells (μM)		IC ₅₀ healthy cells (μM)		SI IMR90/KP4	SI hTERT-	clogP*
		KP4	PANC1	IMR90	hTERT-HPNE		HPNE/KP4	6
1a	phenyl	9.3 ± 1.4	29.6 ± 2.9	23.6 ± 0.9	62.9 ± 7.5	2.5	6.8	2.24
1b	p-fluorophenyl	9.4 ± 1.3	28.6 ± 4.3	16.3 ± 2.9	50.0 ± 5.4	1.7	5.3	2.83
1c	p-chlorophenyl	36.6 ± 5.5	59.1 ± 4.1	43.5 ± 5.2	48.5 ± 9.7	1.2	1.3	3.29
1d	3,5-dimethylphenyl	29.1 ± 3.5	56.2 ± 3.4	89.1 ± 5.3	140.7 ± 21.1	3.1	4.8	3.57
1e	3,5-bis(trifluoro)methylphenyl	134.0 ± 10.7	58.2 ± 8.4	197.0 ± 27.6	374.9 ± 30.0	1.5	2.8	6.4
2a	2-naphtyl	122.2 ± 21.9	104.7 ± 7.3	119.2 ± 27.4	276.0 ± 55.2	1.0	2.3	4.05
2b	9-phenantryl	138.6 ± 16.6	151.4 ± 13.6	190.5 ± 24.7	130.6 ± 10.4	1.4	0.9	5.83
2c	1-pyrenyl	163.2 ± 24.5	240.3 ± 28.3	280.9 ± 25.3	284.3 ± 54	1.7	1.7	6.76
За	2-thiophenyl	10.8 ± 1.7	50.1 ± 6.0	47.9 ± 7.7	120.8 ± 20.5	4.4	11.2	2.26
3b	2-imidazole	0.28 ± 0.02	9.7 ± 1.3	180.1 ± 10.8	257.9 ± 33.5	643.2	921.1	-0.63
3с	2-pyridinyl	5.4 ± 1.0	4.8 ± 0.3	95.0 ± 4.6	198.8 ± 23.8	17.6	36.8	0.78
3d	2-pyrole	0.52 ± 0.03	4.1 ± 0.3	21.6 ± 2.6	74.3 ± 7.4	41.5	142.9	0.62
3e	2-indole	4.9 ± 0.4	7.01 ± 1.2	5.9 ± 1.0	8.8 ± 0.8	1.2	1.8	2.47
3f	5-pyrimidine	118.9 ± 14.3	113.5 ± 22.7	95.6 ± 17.2	398.1 ± 55.7	0.8	3.3	-0.42

^{*}calculated with Swiss ADME consensus

Selectivity Toward Cancer Versus Normal Cells

To evaluate the selectivity of our diiminoguanidines for pancreatic cancer cells over non-cancerous cells, we assessed their antiproliferative activity on two healthy human cell lines: hTERT-HPNE (immortalized pancreatic epithelial cells) and IMR-90 (normal human lung fibroblasts). hTERT-HPNE serves as a physiological control representative of the targeted organ, while IMR-90

represents a non-tumorigenic, proliferative cell line that is often sensitive to chemotherapeutic toxicity, providing a stringent measure of off-target effects. Selectivity indices (SI), calculated as the ratio of IC₅₀ in healthy cells to that in cancer cells, revealed important trends. Compounds bearing phenylic (1a-1e) and polycyclic aromatic (2a-2c) side chains exhibited low selectivity indices (SI < 10), suggesting a narrow therapeutic window and potential for toxicity in non-cancerous tissues. In contrast, some heterocyclic derivatives

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(3a–3d) showed higher selectivity. Notably, compounds 3b–3d, which featured imidazole (3b), pyridine (3c), and pyrrole (3d) side chains, respectively, stood out as the most selective candidates. Compound 3b, in particular, achieved a selectivity index greater than 600 for KP4 cells relative to both hTERT-HPNE and IMR-90 lines, indicating strong preferential activity against malignant cells with minimal cytotoxicity toward normal tissue. Given their potent antiproliferative activity and favorable selectivity profiles, these heterocycle-functionalized diiminoguanidines, especially 3b, represent promising leads for further in vivo evaluation. Their strong discrimination between cancerous and normal cells positions them as attractive candidates for the development of targeted pancreatic cancer therapies with reduced systemic toxicity.

Selectivity indexes were calculated for PANC1 cancer cells as well, and can be found in Supporting Information. Overall, the selectivity indexes on PANC1 cancer cells were in the same range as the one for KP4, except for compound **3b**, suggesting no preference for a particular pancreatic cancer cell line.

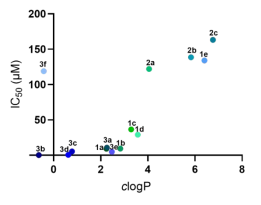


Fig 3. Correlation between clogP and IC₅₀ values of diiminoguanidines on KP4 pancreatic cancer cells

Mechanism of action

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In light of the potent antiproliferative activity demonstrated by our diiminoguanidines, we next investigated their cellular mechanism of action. Given the amphiphilic and cationic nature of these compounds, characteristics known to facilitate mitochondrial targeting, particularly among guanidinium-containing structures, we hypothesized that mitochondria might be their primary intracellular site of action. To test this, we exploited the inherent fluorescence properties of several of our diiminoguanidine derivatives and performed confocal fluorescence microscopy on live pancreatic cancer cells. Cells were co-stained with MitoTracker™ Deep Red (MTDR), a well-established mitochondrial marker, allowing us to assess potential colocalization. The resulting images revealed a strong spatial overlap between the fluorescent signal of our compounds and that of MTDR. This colocalization was visually confirmed by the emergence of yellow signals in merged images (Fig. 4 and Supporting information), and quantitatively supported by Pearson correlation coefficients exceeding 0.89 for all tested compounds—indicative of a robust mitochondrial localization.

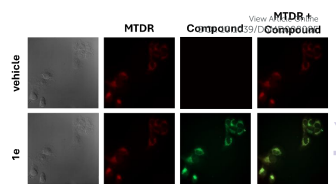


Fig.4 Confocal microscopy images of KP4 pancreatic cancer cells treated with 100 nM diiminoguanidine hydrochloride **1e** or vehicle (4% DMSO) and colocalization with MitoTracker Deep Red (zoom v63)

To further characterize this localization, we conducted timecourse experiments to evaluate the persistence of mitochondrial targeting. Cells treated with diiminoguanidines were imaged at 30 minutes and 24 hours post-incubation. Remarkably, mitochondrial accumulation was evident as early as 30 minutes and remained clearly detectable after 24 hours, indicating that these compounds are not rapidly cleared and instead exhibit sustained mitochondrial residency. Taken together, these findings confirm that our amphiphilic symmetric diiminoguanidines rapidly and durably accumulate in mitochondria. This prolonged retention is likely critical for their antiproliferative efficacy, enabling them to engage mitochondrial targets involved in energy production or apoptosis regulation over an extended period. The observed mitochondrial tropism not only highlights a key feature of the compounds' mechanism of action but also strengthens their candidacy as selective anticancer agents.

Action on the mitochondrial respiratory chain

Mitochondrial dysfunction is a common consequence of therapeutic agents that accumulate within these organelles. When the mitochondrial respiratory chain (MRC) is impaired, cancer cells often respond by shifting their energy production toward aerobic glycolysis, a metabolic reprogramming known as the Warburg effect. To determine whether our diiminoguanidines interfere with mitochondrial respiration, we employed a glycolytic suppression assay using 2-deoxyglucose (2-DG), a well-characterized glycolysis inhibitor. 2-DG is a glucose analog in which the hydroxyl group at the 2-position is absent. This structural modification prevents its metabolism during glycolysis, effectively blocking the glycolytic pathway. By co-administering 2-DG with our compounds, we can test whether the inhibition of glycolysis unmasks a dependence on mitochondrial respiration—a dependence that would become fatal in the presence of mitochondrial impairment. To this end, we treated KP4 pancreatic cancer cells with a non-toxic concentration of 2-DG (1 mM) in combination with each diiminoguanidine at 0.5×IC50 (Fig. 5). A simple 0.5×IC₅₀ treatment served as the baseline for partial growth inhibition. If the addition of 2-DG had no further effect, this Journal Name ARTICLE

would suggest the mitochondrial function remains intact, and cells are not compensating via glycolysis. In contrast, a further reduction in cellular proliferation upon co-treatment would indicate that our compounds impair mitochondrial function and that cells, unable to compensate via glycolysis, undergo energy crisis-induced death.

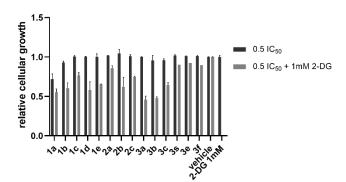


Fig.5 Relative cellular growth of KP4 cancel cells with 0.5 x IC_{50} of diiminoguanidines and a combination of 0.5 x IC_{50} of diiminoguanidines and 1mm 2-DG.

The results were unambiguous: all diiminoguanidines tested exhibited enhanced antiproliferative activity when combined with 2-DG, strongly supporting their inhibitory effect on the mitochondrial respiratory chain. Notably, compound **3b**, already identified as the most potent and selective molecule in the series, produced a ~50% further reduction in cellular viability when used in combination with 2-DG, compared to treatment with **3b** alone. This synergistic effect underscores a robust mitochondrial mechanism of action, reinforcing our earlier microscopy-based conclusions.

Together, these findings provide compelling evidence that diiminoguanidines, particularly compound **3b**, disrupt mitochondrial energy metabolism, and that this disruption is critical to their antiproliferative efficacy. These results also suggest the potential of using glycolytic inhibition as a co-therapeutic strategy to potentiate the effects of mitochondria-targeting agents in pancreatic cancer.

Conclusions

In this study, we successfully synthesized a series of 14 amphiphilic symmetric diiminoguanidinium hydrochloride salts through straightforward, green condensation reactions between N,N'-diaminoguanidine hydrochloride and a diverse set of aldehydes. By systematically varying the side chains—aromatic, polycyclic aromatic, and heteroaromatic—we modulated both the amphiphilicity and biological activity of the resulting compounds. Biological evaluation on pancreatic cancer cell lines revealed potent antiproliferative activity, with most compounds exhibiting IC50 values below 100 μ M. Among them, compound 3b, bearing imidazole side chains, emerged as the lead candidate, with submicromolar IC50 values and an excellent selectivity index exceeding 600 when compared to healthy pancreatic and lung fibroblast cell lines. This high efficacy—selectivity profile positions 3b as a strong candidate for *in vivo* validation.

We identified molecular amphiphilicity (logP) wastica key determinant of activity: compounds with dogP1betweef050an084 showed significantly improved potency, suggesting optimal membrane permeability is essential for mitochondrial access. Confocal microscopy confirmed that diiminoguanidines rapidly accumulate in mitochondria, where they persist for at least 24 h—indicating prolonged mitochondrial engagement. Furthermore, combination assays with 2-deoxyglucose provided mechanistic evidence that the mitochondrial respiratory chain is a critical functional target, particularly for compound 3b.

Taken together, this work introduces a new class of synthetically accessible, mitochondria-targeting anticancer agents with a favorable pharmacological profile for pancreatic cancer. The modular synthesis and tunable side chains offer a promising platform for future optimization toward enhanced selectivity, potency, and translational potential.

Materials and Methods

General information

All chemicals were purchased from Sigma Aldrich, Oakwood Chemicals, and Combi-Blocks at their highest purity and were used without further purification. NMR spectra were recorded on AVANCE II 400, Bruker AVANCE NEO 400, Bruker AVANCE 500, and Bruker AVANCE 700 spectrometers. High-resolution mass spectra (HRMS) and LCMS purity analysis were performed on Agilent TOF instruments by the regional mass spectrometry center (University of Montreal).

Synthesis and characterization of compounds General method for the synthesis of compounds 1a-1e and 2a-2c

To a stirring solution of *N*,*N'*-diaminoguanidine hydrochloride (1 eq.) in methanol (10 mL), aldehyde (2 eq.) was added and the solution was stirred at 80°C for 6h. The reaction mixture was then cooled down and concentrated in vacuo. *N*,*N'*-diiminoguanidines hydrochloride salts were obtained from trituration in acetone to obtain the pure product (yield: 18%-93%).

Synthesis of N,N'-di(phenylmethanimino)guanidine hydrochloride (1a)

Following the general procedure, *N,N'*-diaminoguanidine hydrochloride (1 eq.) and benzaldehyde (2 eq.) yielded compound **1a** as a white powder with a 53% yield. ^1H NMR (400 MHz, DMSO- d_6) δ 12.32 (s, 2H), 8.54 (s, 2H), 8.46 (s, 2H), 7.99 – 7.90 (m, 4H), 7.55 – 7.46 (m, 6H). ^{13}C NMR (101 MHz, DMSO- d_6) δ 153.3, 149.3, 133.7, 131.2, 129.2, 128.3. HRMS: m/z [M+H] $^+$ calculated for $C_{15}\text{H}_{16}\text{N}_5$, 266.1400; found 266.1396.

Synthesis of *N,N'*-di(4-fluorophenylmethanimino)guanidine hydrochloride (1b)

Following the general procedure, *N,N'*-diaminoguanidine hydrochloride (1 eq.) and 4-fluorobenzaldehyde (2 eq.) yielded compound **1b** as a white powder with a 61% yield. 1 H NMR (400 MHz, DMSO-d6) δ 12.36 (s, 2H), 8.57 (s, 2H), 8.45 (s, 2H), 8.04 (dd, J = 8.7, 5.7 Hz, 4H), 7.35 (t, J = 8.8 Hz, 4H). 13 C NMR (101 MHz, DMSO-d6) δ 165.2, 162.7, 153.3, 130.7, 130.6, 130.4, 130.4, 116.4, 116.1. HRMS: m/z [M+H] $^+$ calculated for C₁₅H₁₄N₅F₂, 302.1212; found 302.1212.

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Synthesis N,N'-di(4-chlorophenylmethanimino)guanidine hydrochloride (1c)

Following the general procedure, N,N'-diaminoguanidine hydrochloride (1 eq.) and 4-chlorobenzaldehyde (2 eq.) yielded compound 1c as a white powder with a 76% yield. ¹H NMR (400 MHz, DMSO-d6) δ 12.43 (s, 2H), 8.62 (s, 2H), 8.45 (s, 2H), 8.00 (d, J = 8.6Hz, 4H), 7.57 (d, J = 8.5 Hz, 4H). ¹³C NMR (101 MHz, DMSO-d6) δ 153.3, 148.1, 135.7, 132.7, 130.0, 129.3. HRMS: m/z [M+H]+ calculated for C₁₅H₁₄N₅Cl₂, 334.0621; found 334.0628.

Synthesis of N,N'-di((3,5-dimethyl)phenylmethanimino)guanidine hydrochloride (1d)

procedure, N,N'-diaminoguanidine Following the general hydrochloride (1 eq.) and 3,5-dimethylbenzaldehyde (2 eq.) yielded compound 1d as a white powder with a 93% yield. ¹H NMR (400 MHz, DMSO-d6) δ 12.30 (s, 2H), 8.49 (s, 2H), 8.39 (s, 2H), 7.55 (s, 4H), 7.13 (s, 2H), 2.34 (s, 12H). 13 C NMR (101 MHz, DMSO-d6) δ 153.1, 149.5, 138.3, 133.6, 132.7, 126.0, 21.2. HRMS: m/z [M+H]+ calculated for $C_{19}H_{24}N_5$, 322.2026; found 322.2031.

Synthesis of N, N'-di((3,5-

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bis(trifluoromethyl))phenylmethanimino)guanidine hydrochloride

Following the general procedure, N,N'-diaminoguanidine hydrochloride (1 eq.) and 3,5-bis(trifluoromethyl)benzaldehyde (2 eq.) yielded compound 1e as a white powder with a 93% yield. ¹H NMR (700 MHz, DMSO-d6) δ 12.64 (s, 2H), 8.90 (s, 2H), 8.70 (s, 4H), 8.58 (s, 2H), 8.21 (s, 2H). 13 C NMR (176 MHz, DMSO-d6) δ 154.0, 146.6, 136.7, 131.6, 131.4, 131.2, 131.0, 128.8, 126.0, 124.5, 123.9, 122.9. 121.4.

HRMS: m/z [M+H]⁺ calculated for $C_{19}H_{12}F_{12}N_5$, 538.0896; found 538.0903.

Synthesis N,N'-di(2-naphtylmethanimino)guanidine hydrochloride (2a)

N,N'-diaminoguanidine Following the general procedure, hydrochloride (1 eq.) and 2-naphtaldehyde (2 eq.) yielded compound 2a as a white powder with a 78% yield. ¹H NMR (400 MHz, DMSO-d6) δ 12.49 (s, 2H), 8.65 (s, 4H), 8.33 (dd, J = 13.6, 4.9 Hz, 4H), 8.10 – 7.92 (m, 6H), 7.61 (p, J = 5.4 Hz, 4H). ¹³C NMR (101 MHz, DMSO-d6) δ 153.2, 149.5, 134.5, 133.1, 131.5, 130.3, 128.9, 128.8, 128.3, 127.9, 127.3, 123.7. HRMS: m/z [M+H]⁺ calculated for C₂₃H₂₀N₅, 366.1713; found 366.1722.

Synthesis of N,N'-di(phenantrylmethanimino)guanidine hydrochloride (2b)

the general procedure, N,N'-diaminoguanidine Following hydrochloride (1 eq.) and phenantrene-9-carboxaldehyde (2 eq.) yielded compound 2b as an off-white powder with a 18% yield. ¹H NMR (400 MHz, DMSO-d6) δ 12.51 (s, 2H), 9.45 (s, 2H), 9.05 – 8.94 (m, 2H), 8.92 (d, J = 8.1 Hz, 2H), 8.75 (s, 2H), 8.72 (s, 2H), 8.59 (dd, J= 6.4 Hz, 3.2 Hz, 2H), 8.16 (d, J = 6.9 Hz, 2H), 7.80 (dddd, J = 15.8 Hz,14.7 Hz, 6.2 Hz, 1.6 Hz, 8H). 13 C NMR (101 MHz, DMSO-*d6*) δ 153.0, 131.2, 131.0, 130.5, 129.9, 129.4, 128.6, 128.1, 127.9, 127.7, 124.5, 124.2, 123.5. HRMS: m/z [M+H]⁺ calculated for $C_{31}H_{24}N_5$, 466.2026; found 466.2026.

Synthesis of N,N'-di(pyrenylmethanimino)guanidine hydrochloride (2c)

Following the general procedure, N,N'-diaminoguanidine hydrochloride (1 eq.) and pyrene-1-carbaldehyde (2 eq.) yielded

compound 2c as a green powder with a 52% yield. ¹H NMR (400 MHz, DMSO-d6) δ 12.60 (s, 2H), 9.79 (s, 2H), 9.07 (d; ½0=1832/Hz,√2H)0,8080 (s, 2H), 8.63 (d, J = 9.5 Hz, 2H), 8.50 - 8.34 (m, 8H), 8.30 (q, J = 9.0 Hz,4H), 8.15 (t, J = 7.6 Hz, 2H). ¹³C NMR (101 MHz, DMSO-d6) δ 132.9, 131.2, 130.5, 129.7, 129.4, 129.2, 127.8, 127.1, 126.7, 126.4, 126.4, 125.6, 124.4, 124.2, 122.1. HRMS: m/z [M+H]+ calculated for C₃₅H₂₄N₅, 514.2026; found 514.2021.

General method for the synthesis of compounds 3a-3f

To a stirring solution of N,N'-diaminoguanidine hydrochloride (1 eq.) in methanol (10 mL), aldehyde (2 eq.) was added and the solution was stirred at room temperature for 24h. The reaction mixture was then cooled down and concentrated in vacuo. N.N'diiminoguanidines hydrochloride salts were obtained from trituration in dichloromethane to obtain the pure product (yield: 15%-71%).

Synthesis N,N'-di(thiophenylmethanimino)guanidine hydrochloride (3a)

Following the general procedure, *N,N'*-diaminoguanidine hydrochloride (1 eq.) and 2-thiophenecarboxaldehyde (2 eq.) yielded compound 3a as an orange powder with a 34% yield. ¹H NMR (500 MHz, DMSO-d6) δ 12.29 (s, 2H), 8.65 (s, 2H), 8.24 (s, 2H), 7.79 (d, J =5.0 Hz, 2H), 7.62 (dd, J = 3.6, 0.9 Hz, 2H), 7.19 (dd, J = 5.0, 3.6 Hz, 2H), 3.40 (s, 2H). 13 C NMR (126 MHz, DMSO-d6) δ 152.6, 144.5, 137.8, 132.7, 130.8, 128.4. HRMS: m/z [M+H]⁺ calculated for $C_{11}H_{12}N_5S_2$, 278.0529; found 278.0530.

Synthesis of N,N'-di(imidazolemethan-2-imino)guanidine hydrochloride (3b)

Following the general procedure, N,N'-diaminoguanidine hydrochloride (1 eq.) and imidazole-2-carboxaldehyde (2 eq.) yielded compound **3b** as a beige powder with a 15% yield. ¹H NMR (400 MHz, DMSO-d6) δ 12.53 (s, 4H), 7.97 (s, 2H), 7.38 (s, 2H), 7.26 (s, 4H). ¹³C NMR (176 MHz, DMSO-*d6*) δ 144.3, 141.8, 133.8, 123.8. HRMS: m/z [M+H]⁺ calculated for C₉H₁₂N₉, 246.1210; found 246.1220.

Synthesis of N,N'-di(pyridinylmethan-2-imino)guanidine hydrochloride (3c)

procedure, N,N'-diaminoguanidine Following the general hydrochloride (1 eq.) and pyridine-2-carboxaldehyde (2 eq.) yielded compound 3c as a yellow powder with a 36% yield. ¹H NMR (400 MHz, DMSO-d6) δ 12.73 (s, 2H), 8.75 (s, 2H), 8.67-8.66 (d, J = 4Hz, 2H), 8.52 (s, 2H), 8.48-8.40 (d, J = 8Hz, 2H), 8.00-7.96 (t, J = 7.2 Hz, 2H), 7.52-7.49 (t, J = 4.8 Hz, 2H). ¹³C NMR (126 MHz, DMSO-d6) δ 152.5, 149.7, 137.8, 125.5, 121.9. HRMS: m/z [M+H]+ calculated for C₁₃H₁₄N₇, 268.1305; found 268.1315.

Synthesis of N,N'-di(pyrole-2-methanimino)guanidine hydrochloride (3d)

the general procedure, *N,N'*-diaminoguanidine hydrochloride (1 eq.) and pyrole-2-carboxaldehyde (2 eq.) yielded compound 3d as a khaki powder with a 71% yield. ¹H NMR (400 MHz, DMSO-d6) δ 12.69 (s, 2H), 11.80 (s, 2H), 8.24 (s, 4H), 7.05 (s, 2H), 6.52 (s, 2H), 6.15-6.14 (m, 2H). 13 C NMR (126 MHz, DMSO-d6) δ 152.8, 139.7, 127.5, 123.0, 114.2, 109.8. HRMS: m/z [M+H]+ calculated for C₁₁H₁₄N₇, 244.1305; found 244.1315.

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Synthesis of N,N'-di(indole-5-methanimino)guanidine hydrochloride (3e)

Following the general procedure, N,N'-diaminoguanidine hydrochloride (1 eq.) and indole-5-carboxaldehyde (2 eq.) yielded compound 3e as a pink powder with a 68% yield. ¹H NMR (400 MHz, DMSO-d6) δ 12.04 (s, 2H), 11.49 (s, 2H), 8.48 (s, 2H), 8.32 (s, 2H), 8.03 (s, 2H), 7.82-7.80 (d, J = 8.5 Hz, 2H), 7.50 (s, 2H), 7.48 (s, 2H), 7.44 (s, 2H), 7.82-7.80 (d, J = 8.5 Hz, 2H), 7.50 (s, 2H), 7.48 (s, 2H), 7.44 (s, 2H), 7.82-7.80 (d, J = 8.5 Hz, 2H), 7.50 (s, 2H), 7.48 (s, 2H), 7.44 (s, 2H), 7.82-7.80 (d, J = 8.5 Hz, 2H), 7.50 (s, 2H), 7.48 (s, 2H), 7.44 (s, 2H), 7.82-7.80 (d, J = 8.5 Hz, 2H), 7.50 (s, 2H), 7.48 (s, 2H), 7.44 (s, 2H), 7.82-7.80 (d, J = 8.5 Hz, 2H), 7.82 (s, 2H), 7.84 (s, 2H), 72H), 6.54 (s, 2H). 13 C NMR (126 MHz, DMSO-d6) δ 131.0, 127.1, 122.3, 120.7, 112.6, 102.5. HRMS: m/z [M+H]+ calculated for C₁₉H₁₈N₇, 344.1618; found 344.1629.

Synthesis N,N'-di(pyrimidine-5-methanimino)guanidine hydrochloride (3f)

Following general procedure, N,N'-diaminoguanidine hydrochloride (1 eq.) and 5-formylpyrimidine (2 eq.) yielded compound 3f as a yellow powder with a 53% yield. ¹H NMR (400 MHz, DMSO-d6) δ 12.70 (s, 2H), 9.37 (s, 4H), 9.26 (s, 2H), 8.77 (s, 2H), 8.50 (s, 2H). 13 C NMR (126 MHz, DMSO-d6) δ 159.4, 156.3, 143.6, 128.3. HRMS: m/z [M+H]⁺ calculated for $C_{11}H_{12}N_9$, 270.121; found 270.1220.

Biological assays

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General information

Absorbance measurements for the determination of IC₅₀ values were performed using a SpectraMax 190 microplate reader from Molecular Devices. The obtained data were analyzed with PRISM software by fitting a curve with nonlinear regression (log(inhibitor) vs. response -- Variable slope (four parameters)). hTERT-HPNE cells (hTERT-immortalized epithelial cells) were obtained from Dr. M. Ouellette (University of Nebraska, Medical Center). The pancreatic cells PANC-1 (epithelial morphology cell line isolated from the pancreatic duct of a 56-year-old white male with an epithelioid carcinoma) and KP-4 (human pancreatic ductal carcinoma cell line derived from human ascites) were obtained from ATCC or the Massachusetts General Hospital Center for Molecular Therapeutics

Cellular growth assay

KP4, PANC1, IMR90, and hTERT-HPNE cells were cultured in DMEM (319-015-CL, Wisent) supplemented with 10% fetal bovine serum (FBS, Wisent) and Penicillin/Streptomycin/L-Glutamine (Wisent) at 37°C with 5% CO₂. The cells were seeded at a density of 1x10³ cells per well for KP4 and PANC1, and 4x103 for IMR90 and hTERT-HPNE in 96-well plates for 24 hours. Following this, treatments with different concentrations of vehicle (4% DMSO) and compounds were carried out for 72 hours. The culture medium was then removed. The cells were washed twice with cold PBS to remove any dead cells. A 1% glutaraldehyde solution was added to the wells for 10 minutes to fix the remaining viable cells. The cells were then washed with PBS, stained with 0.2% violet crystal for 30 minutes at room temperature, washed with water, and dried. The adherent violet crystal was solubilized in 10% acetic acid, and absorbance was measured at a wavelength of 590 nm to calculate cell viability.

UV-Vis Stability Analysis

Solutions of diiminoguanidine hydrochlorides were prepared in PBS buffer with 4% DMSO to mimic the dilution conditions used for the in vitro assays. A first UV-Vis spectrum was recorded and named "day 0" to be used as a reference for the maximum wavelength of absorption. The solutions were heated at 37°C for 3 days, and a spectrum was recorded everyday and names "day 1" "day 2" and "dav 3". DOI: 10.1039/D5MD00808E

Fluorescence microscopy

KP4 cancer cells were seeded at a density of 8x10⁴ cells per well on microscope slides in 6-well plates overnight. They were then treated with either the vehicle or the studied compounds for the desired treatment times (30 min, 2 h, 4 h, 24 h). The culture medium was replaced, and a solution of MitoTracker® Deep Red FM (100 nM in DMEM) was added. After 30 minutes at 37°C, the medium was removed. The cells were washed twice with PBS. A 4% paraformaldehyde solution was added to the wells for 10 minutes to fix the cells onto the microscope slides. The cells were then washed three times with a 0.1 M glycine solution in PBS. 8 µL of mounting medium for fluorescence without DAPI was used to mount each slide onto a microscope slide. The slides were sealed, dried, and observed under a microscope. Microscopy images were acquired using the Zeiss Axio-Imager Z2 Microscope. The fluorescence filter sets used were: GFP HE (Zeiss Set 38): BP 470/40x FT 495 BP 525/50m; Cy5 (Chroma 49009): BP ET640/30x BS TLPXR660 BP ET690/50m, along with the DIC analyser.

Author contributions

A.R.S. and S.L. for conceptualization of the project. S.L. for investigation and synthesis of the compounds and biological studies. S.L. for the original draft preparation. S.L. and A.R.S. for writing, review and editing. All authors have given approval to the final version of the manuscript.

Conflicts of interest

There are no conflicts to declare.

Data availability

The data supporting this article have been included as part of the Supplementary Information.

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Notes and references

- 1. Cabasag, C. J.; Ferlay, J.; Laversanne, M.; Vignat, J.; Weber, A.; Soerjomataram, I.; Bray, F., Gut 2022, 71 (8), 1686-1687.
- 2. Halbrook, C. J.; Lyssiotis, C. A.; Pasca di Magliano, M.; Maitra, A., Cell 2023, 186 (8), 1729-1754.
- 3. Rahib, L.; Wehner, M. R.; Matrisian, L. M.; Nead, K. T., JAMA Network Open **2021**, 4 (4), e214708-e214708.
- 4. Rahib, L.; Smith, B. D.; Aizenberg, R.; Rosenzweig, A. B.; Fleshman, J. M.; Matrisian, L. M., Cancer Research 2014, 74 (11), 2913-2921.
- 5. Klein, A. P., *Nat Rev Gastroenterol Hepatol* **2021,** *18* (7), 493-502.
- Montalvo-Javé, E. E.; Nuño-Lámbarri, N.; López-Sánchez, G. N.; Ayala-Moreno, E. A.; Gutierrez-Reyes, G.; Beane, J.; Pawlik, T. M., J Gastrointest Surg 2023, 27 (5), 1001-1010.
- 7. Pothuraju, R.; Rachagani, S.; Junker, W. M.; Chaudhary, S.; Saraswathi, V.; Kaur, S.; Batra, S. K., Journal of Experimental & Clinical Cancer Research 2018, 37 (1), 319.
- 8. Zheng, J.; Guinter, M. A.; Merchant, A. T.; Wirth, M. D.; Zhang, J.; Stolzenberg-Solomon, R. Z.; Steck, S. E., Nutr Rev 2017, 75 (11), 883-908.
- 9. Grant, T. J.; Hua, K.; Singh, A., Progress in molecular biology and translational science 2016, 144, 241-275.
- 10. Kern, S. E.; Hruban, R. H.; Hidalgo, M.; Yeo, C. J., Cancer Biology & Therapy 2002, 1 (6), 607-613.
- 11. Murimwa, G. Z.; Karalis, J. D.; Meier, J.; Nehrubabu, M.; Thornton, M.; Porembka, M.; Wang, S.; Zeh, H. J.; Yopp, A. C.; Polanco, P. M., Journal of Surgical Oncology n/a (n/a).
- 12. Choi, C. W.; Choi, I. K.; Seo, J. H.; Kim, B. S.; Kim, J. S.; Kim, C. D.; Um, S. H.; Kim, J. S.; Kim, Y. H., Am J Clin Oncol 2000, 23 (4), 425-8.
- 13. Longley, D. B.; Harkin, D. P.; Johnston, P. G., Nature Reviews Cancer 2003, 3 (5), 330-338.
- 14. Wang, W. B.; Yang, Y.; Zhao, Y. P.; Zhang, T. P.; Liao, Q.; Shu, H., World J Gastroenterol 2014, 20 (42), 15682-90.
- 15. Siddiqui, N. S.; Godara, A.; Byrne, M. M.; Saif, M. W., Expert Opin Pharmacother 2019, 20 (4), 399-409.
- 16. Smith, D. B.; Neoptolemos, J. P., Core Evid 2007, 2 (2), 111-9.
- 17. Cartwright, T. H.; Cohn, A.; Varkey, J. A.; Chen, Y.-M.; Szatrowski, T. P.; Cox, J. V.; Schulz, J. J., Journal of Clinical Oncology 2002, 20 (1),
- 18. Chllamma, M. K.; Cook, N.; Dhani, N. C.; Giby, K.; Dodd, A.; Wang, L.; Hedley, D. W.; Moore, M. J.; Knox, J. J., Br J Cancer 2016, 115 (6),
- 19. Nitipir, C.; Vrabie, R.; Parosanu, A. I.; Tulin, R.; Cretu, B.; Cursaru, A.; Slavu, I.; Miron, A.; Calu, V.; Orlov Slavu, M. C., Cureus 2021, 13 (11), e19361.
- 20. Kindler, H. L., American Society of Clinical Oncology Educational Book 2012, (32), 232-237.
- 21. Chi, D.-C.; Brogan, F.; Turenne, I.; Zelonis, S.; Schwartz, L.; Saif, M. W., Anticancer Research 2012, 32 (9), 4147.
- 22. Burris, H. A.; Moore, M. J.; Andersen, J.; Green, M. R.; Rothenberg, M. L.; Modiano, M. R.; Cripps, M. C.; Portenoy, R. K.; Storniolo, A. M.; Tarassoff, P.; Nelson, R.; Dorr, F. A.; Stephens, C. D.; Hoff, D. D. V., Journal of Clinical Oncology 2023, 41 (36), 5482-5492.
- 23. Min, Y. J.; Joo, K. R.; Park, N. H.; Yun, T. K.; Nah, Y. W.; Nam, C. W.; Park, J. H., Korean J Intern Med 2002, 17 (4), 259-62.
- 24. Plunkett, W.; Huang, P.; Xu, Y. Z.; Heinemann, V.; Grunewald, R.; Gandhi, V., Semin Oncol 1995, 22 (4 Suppl 11), 3-10.
- 25. Hryciuk, B.; Szymanowski, B.; Romanowska, A.; Salt, E.; Wasąg, B.; Grala, B.; Jassem, J.; Duchnowska, R., Oncol Lett 2018, 15 (2), 1912-1916.

- 26. Armstrong, J. S., *Br J Pharmacol* **2007**, *151* (8), 1154-65, Wilew Article Online
- 27. Wang, C.; Youle, R. J., Annu Rev Genet 2009,143,095/1018/ID00808E
- 28. Fontanesi, F., Mitochondria: Structure and Role in Respiration. In eLS, pp 1-13.
- 29. Wang, H.; Fang, B.; Peng, B.; Wang, L.; Xue, Y.; Bai, H.; Lu, S.; Voelcker, N. H.; Li, L.; Fu, L.; Huang, W., Frontiers in Chemistry 2021,
- 30. Bock, F. J.; Tait, S. W. G., Nature Reviews Molecular Cell Biology 2020, 21 (2), 85-100.
- 31. Nguyen, T. T.; Wei, S.; Nguyen, T. H.; Jo, Y.; Zhang, Y.; Park, W.; Gariani, K.; Oh, C.-M.; Kim, H. H.; Ha, K.-T.; Park, K. S.; Park, R.; Lee, I.-K.; Shong, M.; Houtkooper, R. H.; Ryu, D., Experimental & Molecular Medicine 2023, 55 (8), 1595-1619.
- 32. Green, D. R.; Reed, J. C., Science 1998, 281 (5381), 1309-12.
- 33. Hébert, A.; Parisotto, M.; Ferbeyre, G.; Schmitzer, A. R., Supramolecular Chemistry **2019,** 31 (3), 127-139.
- 34. Hébert, A.; Parisotto, M.; Rowell, M.-C.; Doré, A.; Fernandez Ruiz, A.; Lefrançois, G.; Kalegari, P.; Ferbeyre, G.; Schmitzer, A. R., Scientific Reports 2021, 11 (1).
- 35. Vatté, J.; Bourdeau, V.; Ferbeyre, G.; Schmitzer, A. R., Molecules 2024, 29 (16).
- 36. Vatté, J.; Bourdeau, V.; Ferbeyre, G.; Schmitzer, A. R., Molecules 2024, 29 (22).
- 37. Juliana de Oliveira Carneiro, B.; Tanos Celmar Costa, F.; Steven, R. L.; José Daniel Figueroa, V., Mini-Reviews in Medicinal Chemistry 2020, 20 (5), 342-368.

The data supporting this article have been included as part of the Supplementary Information.