








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New diphenylphosphane derivatives of ketoconazole are promising antifungal agents

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Four new derivatives of ketoconazole (Ke) were synthesized: diphenylphosphane (KeP), and phosphane chalcogenides: oxide (KeOP), sulphide (KeSP) and selenide (KeSeP). These compounds proved to be promising antifungal compounds towards *Saccharomyces cerevisiae* and *Candida albicans*, especially in synergy with fluconazole. Simulations of docking to the cytochrome P450 14 α -demethylase (azoles' primary molecular target) proved that the new Ke derivatives are capable of inhibiting this enzyme by binding to the active site. Cytotoxicity towards hACSs (human adipose-derived stromal cells) of the individual compounds was studied and the IC₅₀ values were higher than the MIC₅₀ for *C. albicans* and *S. cerevisiae*. KeP and KeOP increased the level of the p21 gene transcript but did not change the level of p53 gene transcript, a major regulator of apoptosis, and decreased the mitochondrial membrane potential. Taken together, the results advocate that the new ketoconazole derivatives have a similar mechanism of action and block the lanosterol 14 α -demethylase and thus inhibit the production of ergosterol in *C. albicans* membranes.

In recent times, the emergence of pathogenic fungi resistant to commonly used antifungal drugs has been occurring at unprecedented rates¹. Antifungal resistance is especially a problem with *Candida* infections, and *Candida albicans* is the predominant cause of invasive fungal infections in humans², responsible for ~15% of nosocomial sepsis³. Invasive candidiasis are mortal in 50–80% rates^{4,5}, reaching ~70% even among patients undergoing antifungal therapy^{6,7}. Thus, the search for new antifungals and alternative therapies has become increasingly important. To tackle this problem from a medicinal chemistry perspective, and considering that a modification of existing drugs is more straightforward than developing a new class of therapeutic agents, we chose ketoconazole (**Ke**), a synthetic imidazole antifungal drug approved by FDA in 1981⁸, as a starting molecule for our present study. **Ke** was, for the following decade, the only antifungal available for oral treatment of systemic fungal infections caused by pathogenic yeasts. Its mechanism of action is well established. Similarly to other azole based drugs, its primary molecular target is cytochrome P450 14- α -demethylase (P45014DM)^{9–15}, acting as a competitive inhibitor. **Ke** binds the Fe atom of cytochrome P450 through the N atom of its imidazole ring. Still in the 1980's the interest on **Ke** expanded to other areas, namely as a potential anticancer agent, including in combinational therapy^{16,17}, treatment of prostatic cancer^{18–21} and Cushing's syndrome^{22,23}. Moreover, it was shown that **Ke** could inhibit growing of several malignant and cancer cell lines²⁴, inducing apoptosis through a p53 dependent pathway²⁵ or inducing G₀/G₁ arrest by triggering mitophagy through down-regulation of COX-2 (cyclooxygenase-2)²⁶.

Nowadays, due to all these diverse effects, **Ke** is regaining strong interest and new ketoconazole derivatives^{27–29} or complexes with several metal ions^{30–32} are being extensively studied. In this work, we decided to follow the pathway of modifying the existing drug and to investigate the properties of aminomethylphosphane derivatives of ketoconazole. Although a long time has passed since the first reports³³, aminomethylphosphanes (or α -aminophosphanes) have been not considered interesting as potential drugs or components of biologically active metal complexes until the last decade. Recently, however, due to their flexibility and diversity, combined

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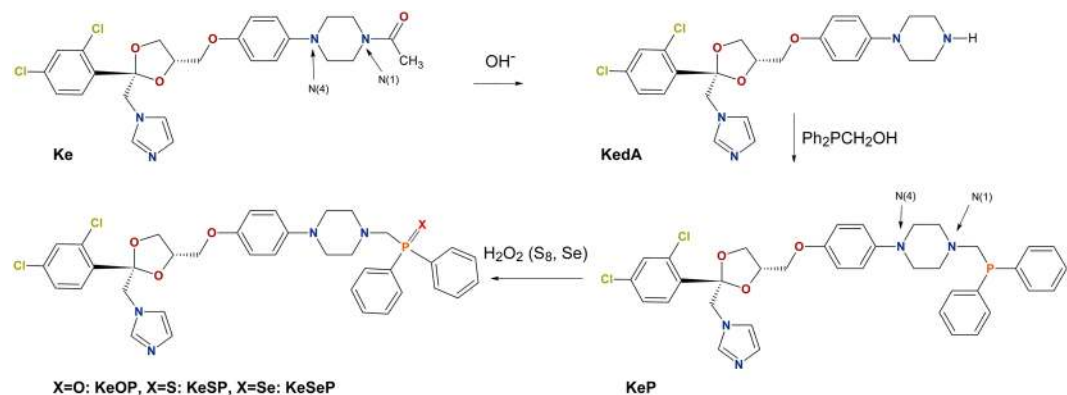


Figure 1. Molecular schemes of ketoconazole (**Ke**) and its derivatives.

with the ease of synthesis, they started to gain a great attention. These compounds can bear an almost infinite variety of substituents with different hydrophilicities and steric demands. The potential presence of auxiliary coordinating atoms allows them to act as mono-, bi- or multidentate ligands. Our team has been working with this class of ligands and their Cu(I), Pt(II) and Ru(II) complexes and, in many cases, the properties of the derivatized molecules and their complexes are more interesting than the parent ones. For example, we worked with trisaminomethylphosphanes derived from morpholine and thiomorpholine (see for example refs.^{34–37}) and a variety on N4-substituted piperazines^{34,35,38–41}, as well as with monoaminomethyldiphenylphosphanes^{40–42}, including a derivative of a model dipeptide⁴³ and the derivatives of selected fluoroquinolones: ciprofloxacin, norfloxacin^{44–46}, lomefloxacin⁴⁷ and sparfloxacin^{48–50}. Some of the compounds listed above showed interesting antimicrobial and/or anti-tumour activity.

Herein we present the synthesis and characteristics of the diphenylphosphanomethyl derivative of ketoconazole and its chalcogenide derivatives (oxide, sulphide and selenide) in order to understand better the influence of a diphenylphosphanomethyl(chalcogenide) moiety on the activity of the ketoconazole molecule.

Results and Discussion

Synthesis and characteristics of the compounds. **KeP** (Fig. 1) was synthesized from deacylated ketoconazole (**KedA**) in a modified Mannich condensation reaction of hydroxymethylphosphanes with amines³³. Briefly, we added **KedA** to the solution of $\text{PPh}_2\text{CH}_2\text{OH}$ obtained *in situ* from $\text{PPh}_2(\text{CH}_2\text{OH})_2\text{Cl}$ by adding the excess of NEt_3 . Chalcogenides were synthesized in a reaction of **KeP** with a stoichiometric amount of H_2O_2 , resublimed sulphur or metallic selenium in the ultrasound bath^{34,44,45,48}. To confirm the assumed structures and check the purity of the products we employed mass spectrometry and elemental analysis as well as NMR spectroscopy.

Mass spectra (Fig. S2 in ESI) confirmed the structures of the synthesized compounds. In the positive mass spectrum of the **Ke** molecule the most intense signals are $[\text{M} + \text{H}]^+$ (46%) with the most intense peak at 531.2 m/z, $[\text{M} + \text{Na}]^+$ (100%) at 553.1 and $[2\text{M} + \text{Na}]^+$ (63%) at 1083.5 showing strong tendency to form dimers under the experiment conditions. Phosphane **KeP** undergoes a strong fragmentation. The phosphane signal $[\text{M} + \text{Na}]^+$ at 709.2 was very weak (15%) and the most intense fragmentation signals found and identified were at 489.2 $[\text{KedA} + \text{H}]^+$ (6%) at 489.2 and $[\text{KedA} + \text{CH}_2]^+$ (100%) at 501.2. For the chalcogenide derivatives we observed a strongly increased stability of the molecules. In the case of **KeOP** the strongest signals were at 703.2 and 725.2 corresponding to $[\text{M} + \text{H}]^+$ (6%) and $[\text{M} + \text{Na}]^+$ (100%) ions respectively. A weak tendency to form dimers was reflected in the presence of the $[2\text{M} + \text{Na}]^+$ signal (5%) at 1429.4. Mass spectra of **KeSP** and **KeSeP** are very similar to one another. For both compounds we observed a small amount of the fragmentation products with dominating $[\text{KedA} + \text{CH}_2]^+$ at 501.2 with 13% or 19% intensity for **KeSP** and **KeSeP**, respectively. For **KeSP** the molecular signals were of $[\text{M} + \text{H}]^+$ (29%) at 719.2 and $[\text{M} + \text{Na}]^+$ (100%) at 741.2. The dimers were observed as the intense signals of $[2\text{M} + \text{H}]^+$ (51%) at 1439.4 and $[2\text{M} + \text{Na}]^+$ (32%) at 1461.4. For **KeSeP** the molecular signals were of $[\text{M} + \text{H}]^+$ (48%) at 767.1 and $[\text{M} + \text{Na}]^+$ (100%) at 789.1. The intensity of the signals of the dimers $[2\text{M} + \text{H}]^+$ (98%) at 1533.3 and $[2\text{M} + \text{Na}]^+$ (30%) at 1555.3 indicates that the tendency for dimer formation of the selenide derivative is the strongest among the compounds studied.

We measured the ^1H , $^{13}\text{C}\{^1\text{H}\}$ and $^{31}\text{P}\{^1\text{H}\}$ (when applicable) NMR spectra of ketoconazole and its derivatives in CDCl_3 . Using two-dimensional COSY, HMQC and HMBC spectra we were able to assign unambiguously all the signals to the corresponding atoms in **Ke** molecule (see Figs S4–S10 and Table S1 in ESI) as well as in its derivatives (Table S2). As expected, ^1H NMR spectrum of **Ke** is very complex and shows a set of characteristic patterns due to a presence of three different aromatic rings, racemic mixture of the chiral 2,2,4-trisubstituted 1,3-dioxolane ring and acetylated piperazine ring. It should be mentioned that the piperazine moiety in the **Ke** molecule shows four independent CH_2 signals (carbon singlets and corresponding proton multiplets) indicating a strong binding of the acetyl group leading to the restrained rotation of the N-C bond.

Formation of **KeP** from $\text{PPh}_2(\text{CH}_2\text{OH})$ and **KedA** and further formation on the chalcogenides and the completion of the reaction is best observed by recording $^{31}\text{P}\{^1\text{H}\}$ spectra (see Tables 1, S2). In CDCl_3 solution, $\text{PPh}_2(\text{CH}_2\text{OH})$ phosphorus atom exhibits a singlet at -10.03 ppm. Formation of the **KeP** molecule shifts this signal upfield to -27.39 ppm. Subsequent formation of chalcogenides transfers the electron density from the P atom towards a substituent, deshielding the phosphorus atom. Chemical shift for **KeOP** was 27.18, for **KeSP** was

		Ke	KeP		KeOP		KeSP		KeSeP	
$^{31}\text{P}\{^1\text{H}\}$			δ [ppm]		δ [ppm]		δ [ppm]		δ [ppm]	$^1J_{\text{SeP}}$ [Hz]
			-27.39		27.18		35.15		25.93	721.5
$^{13}\text{C}\{^1\text{H}\}$		δ [ppm]	δ [ppm]	$^3J_{\text{CP}}$ [Hz]	δ [ppm]	$^3J_{\text{CP}}$ [Hz]	δ [ppm]	$^3J_{\text{CP}}$ [Hz]	δ [ppm]	$^3J_{\text{CP}}$ [Hz]
-CH ₂ PPh ₂	C(1) ^{x=1}		61.4	3.6	58.4	88.1	63.0	73.6	63.5	66.3
	C(i) ^{x=1}		138.38	12.71	132.42	97.2	131.94	77.2	130.56	69.6
	C(o) ^{x=2}		132.83	18.16	131.28	9.1	131.91	10.0	132.54	10.0
	C(m) ^{x=3}		128.36	6.36	128.50	11.8	128.42	11.8	128.44	11.8
	C(p) ^{x=4}		128.52	S	131.86	2.7	131.64	2.7	131.74	2.7
pip	C(2,6) ^{x=3}	41.41/46.31	54.55	9.08	55.54	8.2	55.24	7.3	55.15	6.4
	C(3,5)	50.63/50.89	50.46	S	50.43	s	50.40	s	50.39	s
^1H		δ [ppm]	δ [ppm]	$^3J_{\text{CP}}$ [Hz]	δ [ppm]	$^3J_{\text{CP}}$ [Hz]	δ [ppm]	$^3J_{\text{CP}}$ [Hz]	δ [ppm]	$^3J_{\text{CP}}$ [Hz]
-CH ₂ PPh ₂	H(1) ^{x=2}		3.28	2.86	3.31	6.94	3.53	5.34	3.66	4.58
pip	H(2;6) m	3.76/3.61	2.83		2.81		2.69		2.67	
	H(3,5) m	3.02/3.05	3.12		3.03		2.97		2.96	

Table 1. Selected NMR data for ketoconazole and its derivatives (CDCl₃). *Observed as a singlet.

<i>S. cerevisiae</i> strains	Ke	KeP	KeOP	KeSP	KeSeP
WT (BY4741)	0.865	9.023	0.00416	>200	1.244
<i>erg6</i> Δ	>200	0.654	0.00482	>200	0.578
WT (W303)	6.25	12.5	3.13	>200	50
AD1-8	0.02	0.2	0.05	0.2	0.2

Table 2. Activity expressed as MIC₅₀ values [μM] of Ke and its derivatives against *S. cerevisiae* cells.

35.15 and for **KeSeP** the chemical shift was 25.93 ppm with the 1J satellite coupling to ^{77}Se equal to 721.5 Hz. All these values are typical for the diphenylphosphanomethyl ligands with N4-substituted piperazine ring attached to the P atom via methylene group and do not depend on N4 substituent^{40,45}. In ^1H and $^{13}\text{C}\{^1\text{H}\}$ NMR spectra, phosphane formation from KedA results in the new signals of methylene group being a link between **KedA** and **-PPh₂** moieties. Both proton and carbon signals are doublets with relatively small coupling constants to the phosphorus atom. This process is reflected also in the carbon signals of neighboring CH₂ groups (position 2,6) in the piperazine ring, where a P-coupled doublet with $^3J(\text{P-C})$ around 10 Hz is observed. As for the tris(aminomethyl) phosphanes³⁴ and the aminomethyldiphenylphosphanes^{40,45}, formation of the chalcogenide derivatives induces dramatic changes in the signals of the groups directly bound to the P atom. This is best seen in the carbon signal of -CH₂-P group, where formation of the P = X bond not only shifts the signal, but also significantly alters the values of $^1J(\text{C-P})$ coupling constant: from 3.6 Hz for **KeP** to 88.1, 73.6 and 66.3 Hz for **KeOP**, **KeSP** and **KeSeP**, respectively. These changes are almost repeated in the signals of the phenyl ring carbon atoms directly bound to the P atom (C(i)).

Activity against yeast *Saccharomyces cerevisiae*. For the preliminary studies of the biological activity of the studied compound we chose *S. cerevisiae*, which is generally considered a non-pathogenic organism, but currently recognized as a potentially important agent causing opportunistic infections against immunocompromised patients⁵¹. In order to assess whether all the **Ke** derivatives share the mechanism of action with the parent compound, or if they might present different or additional modes of action, the inhibitory concentration MIC₅₀ (Table 2) was determined on the basis of the dose-response curves (Fig. S11 in ESI) in a 48 h assay against *S. cerevisiae* wt strain BY4741 and the deletion mutant *erg6*Δ, which lacks the gene coding for the S-adenosylmethionine: 24-methyltransferase⁵², accumulating cholesta-5,7,24-trienol and zymosterol instead of ergosterol⁵³. There are several studies reporting that the ERG6 gene deletion may increase the resistance of *S. cerevisiae* cells to azole compounds^{54–56}. Thus, we first studied **Ke**, and confirmed this behavior, as a MIC₅₀ of 0.865 μM was determined against wt cells, whereas this compound showed no activity against the mutant cells. The value determined for the wt strain is in very good agreement with previous studies carried out under similar condition (48 h, 30 °C) (e.g.⁵⁷).

Regarding the **Ke** derivatives, they exhibit clearly distinct behavior from the parent compound, and notably, they were all active against the resistant strain to similar or even greater extent as they were against the wt strain (except **KeSP**, which was also inactive against wt cells). These results strongly suggest that the **Ke** derivatives may present different or additional modes of action underlining their antifungal activity, or override mechanism of drug resistance operative against **Ke**, since not only they are active against yeast cells that are resistant to azole compounds, and shown to be **Ke** resistant in this study, but also the behavior in relation both to the parent compound and between the two strains can change. For **KeP**, the wt cells are less sensitive (MIC₅₀ 9.023 μM) than for **Ke**, while *erg6*Δ cells are very sensitive, presenting a MIC₅₀ value (0.654 μM) which is even smaller than the one found for the parent compound against the wt. **KeSeP** and **KeOP** are active against both strains with similar

	Flc	Ke	KeP	KeOP	KeSP	KeSeP
CAF2-1	6.25	0.05	3.13	0.78	3.13	3.13
DSY1050	0.78	0.02	0.78	0.2	0.39	0.78
B3	6.25	0.05	0.39	1.56	1.56	3.13
B4	50	0.39	3.13	3.13	3.13	6.25
Gu4	6.25	25	3.13	12.5	50	12.5
Gu5	>200	50	50	25	>200	>200

Table 3. MIC₅₀ values [μM] for fluconazole (Flc), Ke and its diphenylphosphanomethyl derivatives towards *C. albicans* CAF2-1, DSY1050, B3, B4, Gu4 and Gu5.

MIC₅₀ values (Table 2). Finally, as mentioned above, **KeSP** is inactive against both *S. cerevisiae* strains. **KeOP**, on the other hand, presents very high activity (MIC₅₀ values on the nM range) against both wt and *erg6Δ* cells.

The lack of activity of **KeSP** is somewhat surprising, as it contrasts markedly with the behavior exhibited by the two other calchogenide derivatives, **KeOP** and **KeSeP**. We have, thus, hypothesized that this compound could be more efficiently exported by plasma membrane pumps. To test this hypothesis, we measured the activity of the compounds against a strain that lacks all the major ABC transporters in *S. cerevisiae* responsible for drug export, which is in general very susceptible to most xenobiotic compounds⁵⁸. As control, we have also used the respective wt strain, W303. As with BY4741, W303 cells are susceptible to **Ke** and its derivatives, except for **KeSP**. This suggests that the inactivity of this compound against *S. cerevisiae* is a general trend, as it was observed in two different genetic backgrounds. On another hand, the MIC₅₀ values are always considerably higher for W303 cells than for BY4741. This result is not unexpected, since it has been previously reported that, when grown in YPD medium, as in this study, BY4741 cells are more sensitive to Hygromycin B than W303, and they also differ in alkali-metal cation tolerance and plasma membrane potential, and in other important physiological parameters, as consequence of their different genome^{59,60}. Notwithstanding, it is noteworthy that for both wt strains; the order of MIC₅₀ values for the different compounds tested is almost the same, the only difference being between **KeSeP** and **KeP**, which are the second and third most active against BY4741 and in reverse order for W303 cells.

Regarding the strain lacking the major plasma membrane drug exporters, AD1-8, it is clear that all the compounds show much higher antifungal activity. The most striking result is the one for **KeSP** which displays a MIC₅₀ value that is similar to **KeP** and **KeSeP**, and is more than 3 orders of magnitude lower than against the W303 cells, the wt with the same genetic background. The most active derivative is, again, **KeOP**. These results strongly suggest that the inactivity of **KeSP** is due to a very efficient efflux of this compound by *S. cerevisiae*.

Candida albicans viability. *Minimal inhibitory concentration (MIC).* Fungistatic (MIC₅₀) effect of diphenylphosphanomethyl derivatives of **Ke** was evaluated in comparison to **Ke** and a triazole antifungal – fluconazole (**Flc**) (Table 3). **Flc** is the most commonly prescribed antifungal azole drug⁶¹ but approximately 16-fold less active *in vitro* than **Ke**⁶². **Ke** based phosphanes were less active than **Ke**; however, two (**KeP**, **KeSP**, **KeSeP**) or eight (**KeOP**) times more active than **Flc** towards *C. albicans* CAF2-1 strain, referential in our study.

High resistance of *C. albicans* towards azole based agents is related to the development of a multi-drug resistance (MDR) phenotype found among clinical *C. albicans* isolates, which connects with overproduction of efflux transporters located in the plasma membrane (Cdr1 and Cdr2, belonging to ATP-binding cassettes (ABCs) and Mdr1, belonging to major facilitator superfamily (MFS))⁶³. In order to determine the role of MDR efflux pumps in resistance towards **Ke** derivatives, we used three pairs of *C. albicans* strains - CAF2-1 and DSY1050; B3 and B4; Gu4 and Gu5. The mutant with no functional *CDR1*, *CDR2* and *MDR1* genes (DSY1050) was four (**KeP**, **KeOP**, **KeSeP**) or eight (**KeSP**) times more susceptible than its parental strain (CAF2-1). *C. albicans* B3 and B4 strains were isolated from a patient before and after fluconazole administration, respectively. Azole-resistance of B4 results from overexpression of *MDR1*, therefore two (**KeOP**, **KeSP**, **KeSeP**) and eight (**Ke**, **KeP**) times higher resistance of B4 than B3 indicates a possible role of Mdr1p in **Ke** derivatives efflux. High levels of Mdr1p are related to the development of resistance towards voriconazole (**Vor**) and **Flc**^{64–67} but to lesser extent for other azoles, e.g. **Ke**^{68–71}.

C. albicans Gu4 and Gu5 strains were also isolated from a patient before and after fluconazole administration, respectively. Contrary to B4, azole-resistance of Gu5 origins in overexpression of *CDR1* and *CDR2*. Cdr1p plays a major role in antifungal resistance in *C. albicans* cells by exporting most non-related xenobiotics outside fungal cells⁷². **KeSP** and **KeSeP** displayed no inhibitory effect on Gu5 strain and this strain was two and sixteen times more resistant than Gu4 strain towards **KeOP** and **KeP**, respectively. This result indicates a strong involvement of Cdr1p and Cdr2p in **Ke** derivatives resistance and that **Ke** derivatives are possibly substrates for Cdr1/Cdr2 pumps.

KeOP/KeP synergism with fluconazole. In case of candidiasis treatment, a synergistic combination of clinically used azoles with other drugs is highly desirable^{73,74}. One of the approaches is to inhibit activity of MDR efflux pumps by administration of competitive pump substrates among azole drugs⁷⁵. Thus, we combined **Flc** with **KeP** or **KeOP**, since both **Ke** derivatives had most promising MIC₅₀ values. **KeP** (0.39 μM) and **KeOP** (1.56 μM) reduced MIC₅₀ of **Flc** in the range from 6.25 μM to 1.56 μM and when combined with higher (3.25 and 6.25 μM) **Flc** concentrations lead to fungicidal effect (Fig. 2 and Table S3 in ESI). Since azoles are fungistatic^{2,76}, biocidal combination of **KeP/KeOP** with **Flc** is promising, especially, because simultaneous administration of different azoles may either lead to synergistic, subadditive or antagonistic effects⁷⁷.

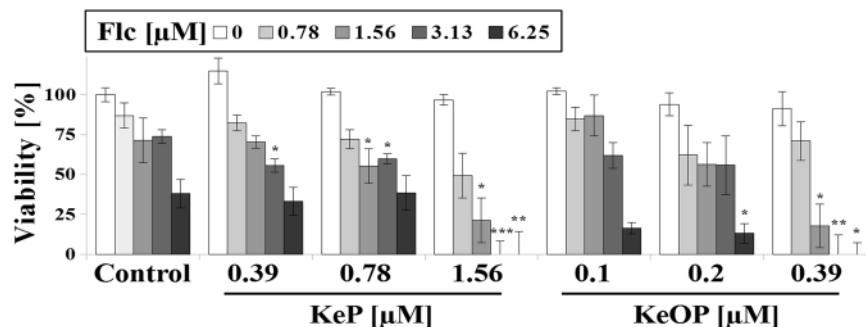


Figure 2. KeP and KeOP synergism with fluconazole (Flc). Viability [%] of *C. albicans* CAF2-1. Statistical analysis was performed comparing viability at corresponding Flc concentrations with/without KeOP/KeP (*0.01 < P < 0.05; **0.001 < P < 0.01; ***P < 0.0001). \pm SD, n = 3.

Human adipose-derived stromal cells (hASCs) studies. *Half maximal inhibitory concentration (IC₅₀) assessment.* In order to assess the activity of the ketoconazole and its derivatives against human cells, IC₅₀ values for Ke, KeP, KeOP as well as KeSeP for hASCs (human adipose-derived stromal cells) were obtained from the dose-response curves that were prepared using TOX-8 Assay after a 24-hours stimulation (Fig. 3).

The cytotoxicity assay indicated KeP (IC₅₀ = 37.65 μ M) as the most active compound. The cytotoxicity of KeOP (IC₅₀ = 120.8 μ M) and KeSeP (IC₅₀ = 109.8 μ M) were significantly lower, and the lowest cytotoxic effect was found for Flc and KeSP (Fig. 3), which were not cytotoxic in the tested concentration range. These data stand in good agreement with other results, that show a relatively small cytotoxicity of Flc⁷⁸. In this study, the IC₅₀ value for Ke is 120.8 μ M. However, contradictory data regarding cytotoxicity of Ke has been reported⁷⁸, which might be due to the use of different cell sources. It was previously shown by Benkoš *et al.* that the IC₅₀ value of Ke for whole population of human bone marrow cells was equal to 6,27 mg/L (11.8 μ M). Moreover, the IC₅₀ of Ke was almost 3-fold higher for murine bone marrow cells in comparison to the human cells⁷⁸. In turn, Haegler *et al.* showed that in HepG2 cells (liver hepatocellular carcinoma cell line), Ke cytotoxic effects started to be noticed at 50 μ M. In contrast, in HepaRG cells, cytotoxicity started at 100 μ M⁷⁹.

Apoptosis level of hASCs after 24-hours stimulation with Ke, KeP and KeOP (20 μ M). Apoptosis is characterized by cell shrinkage, alternations in the cell membrane and mitochondria, nuclear condensation, as well as DNA fragmentation and protein degradation by caspases⁸⁰. It was shown that BCL-2 transcript protects cells from apoptosis by associating with mitochondria and inhibiting their release of cytochrome c, whereas BAX promotes apoptosis. Moreover, p53 gene expression is associated with induction of apoptosis^{80,81}. Interestingly, p21 transcript plays a dual role in cell death and survival, i.e., it inhibits apoptosis through induction of cell cycle arrest and DNA repair, but p21-arrested cells may undergo apoptosis following activation of other pro-apoptotic genes⁸¹. In this study, we observed significant (p < 0.001) elevated expression of p21 transcript in cells challenged with Ke, KeP or KeOP when compared to the control group. Moreover, the relative expression levels of both BAX and BCL-2 were higher in cells treated with Ke, KeP or KeOP, but the differences were not significant. Interestingly, none of the compounds increased the expression of a master regulator of apoptosis, which is a p53 transcript (Fig. 4). Furthermore, no significant changes in the expression of p53 gene were observed when compared to control cells. This observation might suggest that the tested dosage of Ke, KeP and KeOP, which was 20 μ M, did not induce apoptosis in hASC cultures, which might be a positive feature light for their future potential clinical application in antifungal therapy.

Mitochondria network and F-actin in hASCs. Cell morphology was evaluated using MitoRed and Phalloidin Atto 590 staining after 24 h stimulation with Ke, KeP and KeOP (20 μ M). MitoRed is a vital dye which interacts with functional mitochondria due to their high membrane mitochondrial potential. Decreased fluorescence of MitoRed stain has been associated with reduced mitochondrial membrane potential⁸². Patalano *et al.* showed that Ke treatment induces depolarization of mitochondria which leads to the activation of apoptosis in mouse corticotroph tumour cell line (AtT20-D16)⁸³. Moreover, Haegler *et al.* showed that Ke decreases the mitochondrial potential in HepG2 cells, which was associated with decreased mitochondrial DNA synthesis and accumulation of mitochondrial superoxide that induce apoptosis⁸⁴. In this work, the mitochondrial staining indicates the lowest mitochondrial membrane potential in KeP treated cells, as well as the lowest number of mitochondria. (Fig. 5). This might suggest a reduced antioxidative capacity. In turn, phalloidin staining showed a multilayer growth pattern in cells treated with Ke and KeOP. These cells were characterized by close contact with a relatively dense growth pattern, while hASCs treated with KeP were characterized by a loose growth pattern. It can be concluded that the compounds tested, i.e. Ke, KeP and KeOP, affect in different ways the morphological features of progenitor cells. Moreover, KeP seems to affect negatively cellular morphology and mitochondrial activity.

Docking to CYP51 of *C. albicans*. As it has been said in the introduction, primary molecular target of Ke is cytochrome P450 14- α - demethylase. Mechanism of action is pretty simple. Ke, like other azoles, acts as an competitive inhibitor binding cytochrome Fe atom by N atom from imidazole ring and therefore blocking the enzyme active site. It was proven by several X-ray structures of the enzyme with different azoles. For Ke there one

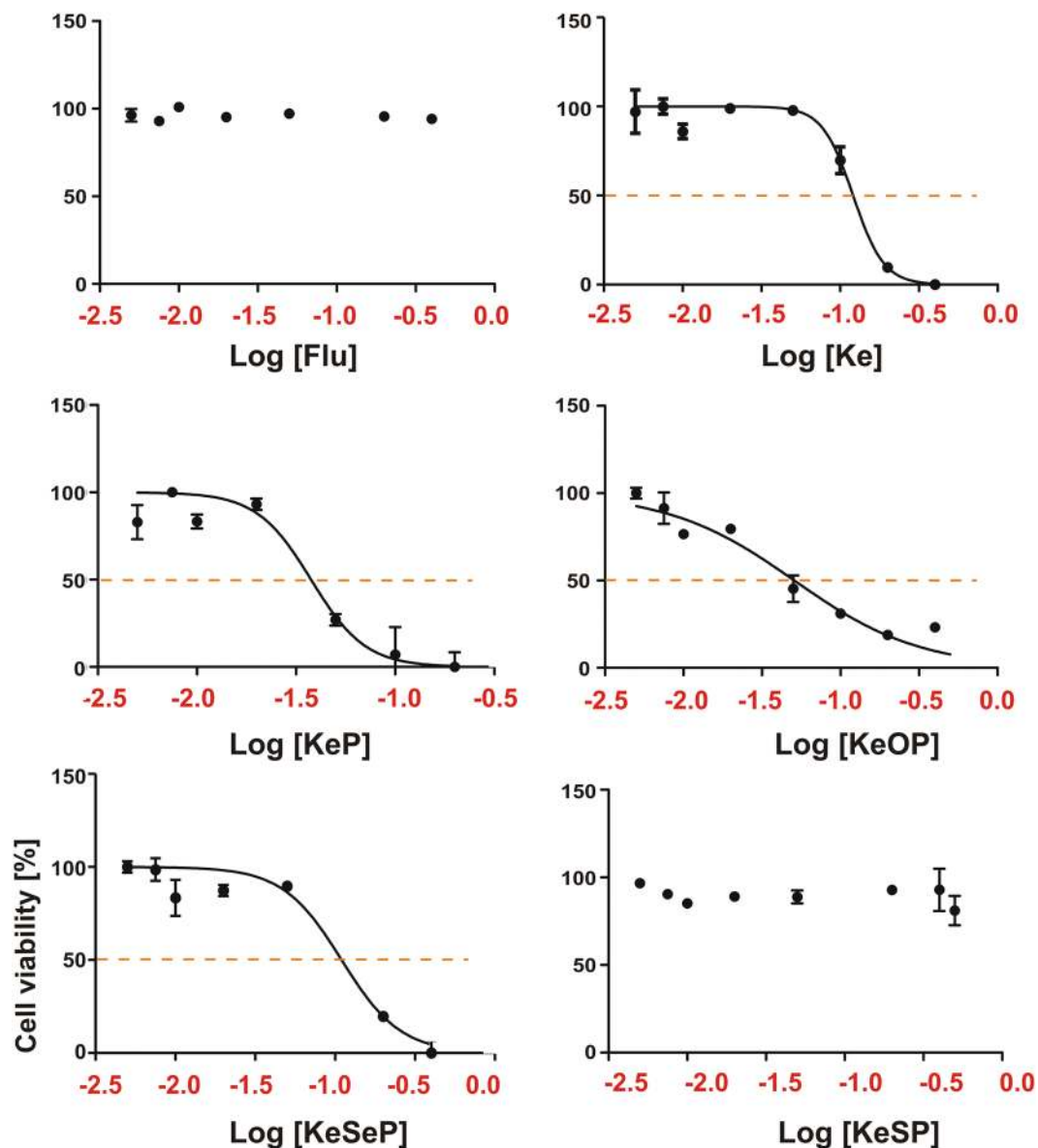


Figure 3. hACSs viability in the presence of different azole compounds. Percentage of cell survival is plotted against the logarithm of treatment concentrations. Calculated IC₅₀ values are: 120.8 for Ke, 37.65 for KeP, 50.76 for KeOP and 109.8 μM for KeSeP. It was not possible to determine the IC₅₀ values for Flu and KeSP.

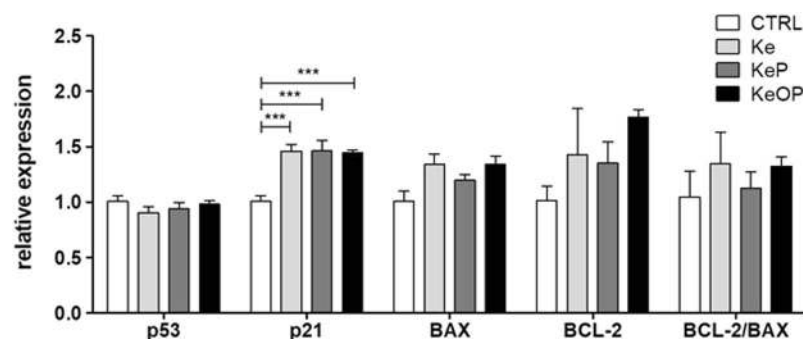


Figure 4. Apoptosis level of hACSs treated with 20 μM Ke, KeP and KeOP for 24 hours. Results expressed as mean ± SD. Statistical significance indicated as three asterisks ***p < 0.001.

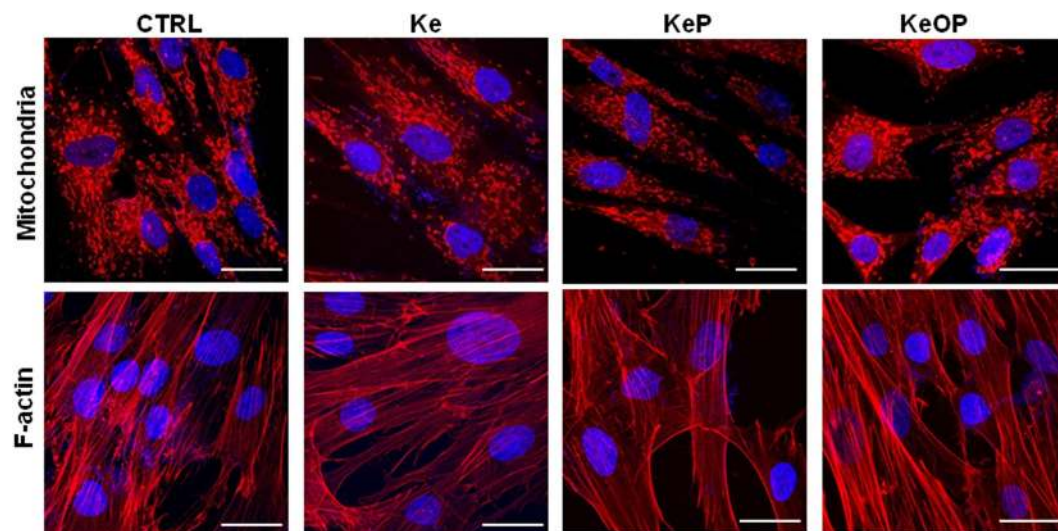


Figure 5. Visualization of mitochondrial network and F-actin of hASCs treated with **Ke**, **KeP** and **KeOP** (20 μ M; 24 h) using MitoRed and Phalloidin Atto 590 staining, respectively. Scale bar: 50 μ m.

can find a structure of human CYP51 inhibited by the **Ke** molecule^{13,14}, and only recently a structure of fungal CYP51 from *C. albicans* in complex with posaconazole was determined¹⁵. To check if the presented here derivatives can also target this enzyme we performed Molecular Docking using AutoDock Vina software⁸⁵.

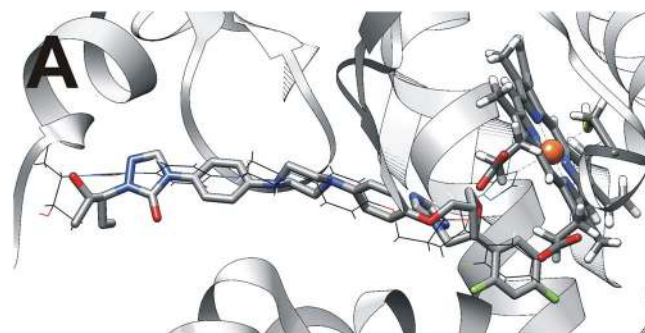
The docking results together with the best-fit structures are summarized in Fig. 6. The reference structure is the CYP51-posaconazole complex and the location of posaconazole serves to determine the correct binding mode. Posaconazole itself was also subjected to the docking to validate the docking procedure. Interestingly, while the correct overall binding mode is found for posaconazole, the triazole ring is rotated so that the Fe-N distance is much larger than in the experimental crystal structure (3.96 Å vs. 2.08 Å). However, the calculated binding affinity for posaconazole is the second strongest among the molecules studied, next only to **KeP**. For all five **Ke** derivatives, the proper location was found by the docking algorithm, which is a good indication of their binding affinity and the proper shape of the binding site cavity. The search space was large enough (cubic box, $a = 80$ Å) not to force the ligand into the cavity. The best-fit structures are always located in the binding site and oriented properly (that is, the heme iron atom is coordinated by the N atom in the imidazole ring of a given derivative), with the two important exceptions of **KeSP** and **KeSeP**, for which the sulfur or selenium atoms are oriented towards the heme iron. The resulting structure is “head to tail” with respect to posaconazole and the remaining **Ke** derivatives. This might arise from the stronger preference of the Fe-S(Se) interaction over the Fe-N contact, or from an increased volume of the P-S(Se) fragment. The matter of possible Fe-S(Se) coordination will be further investigated experimentally; nevertheless, both **KeSP** and **KeSeP** are also located in the correct cavity. Taking into account the approximate nature of the docking score function, it is not possible to determine whether the investigated **Ke** derivatives bind more or less strongly to the protein than **Ke** itself, but no dramatic changes were noted, suggesting that binding affinities among the analyzed **Ke** derivatives should be of similar order.

For **KeP** and **KeOP**, the ligands oriented as **Ke**, an analysis of the vicinity of the phosphorus atom was carried out. For **KeP**, the P atom interacts via its lone pair with the Pro230 ring (the closest distance P-C(Pro230) is 3.811 Å). Moreover, the P atom is also in potential contact with the phenyl ring of Phe233 (the closest P-C(Phe233) distance is 4.766 Å). In the case of **KeOP**, the direct environment around the ligand is somewhat different: Pro230, Phe233, Phe235 and Phe380 form a hydrophobic cage surrounding the $-PPh_2$ phenyl rings of the ligand, while the P=O moiety is in contact with the Tyr64 phenol group (the closest (**KeOP**)O...C(Tyr64) distance is 3.313 Å, and the O...O(phenol,Tyr64) distance is 4.667 Å). These values must be taken as a rough estimation, due to the aforementioned nature of the docking scoring function, but they represent a possible diversity of bonding between the protein and the phosphane moiety of the **Ke** derivatives. In particular, the possibility of additional stabilization by interactions with a hydrophobic cage could add to the binding strength of **KeP**.

Conclusions

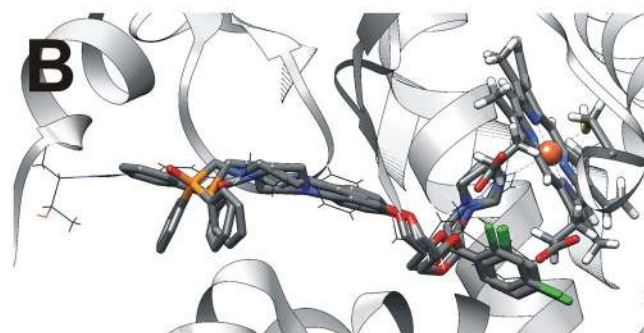
Derivatization of ketoconazole with diphenylphosphanomethyl moiety gave an opportunity to develop a set of similar molecules differing only by the direct environment of the phosphorus atom. Four new derivatives of ketoconazole (**Ke**) were synthesized: diphenylphosphane (**KeP**) and its chalcogenides: oxide (**KeOP**), sulphide (**KeSP**) and selenide (**KeSeP**). Their structures were analyzed using NMR spectroscopy and mass spectrometry.

The derivatives of ketoconazole synthesized and tested in this work, especially **KeP** and **KeOP**, proved to be promising antifungal compounds. A strong antifungal activity, in some occasions surpassing largely the parent compound, was shown for selected strains of *S. cerevisiae* and also in the detailed study for *C. albicans*, especially in synergy with fluconazole. All compounds tested are substrates for pumps expelling drugs (mainly Cdr1 and Cdr2) from *C. albicans* cells, and simultaneous administration of fluconazole allowed a significant reduction in concentrations of ketoconazole derivatives tested and low survival of the fungus. Simulations of docking of the derivatives to the cytochrome P450 14- α -demethylase (the azoles' primary molecular target) did not show large



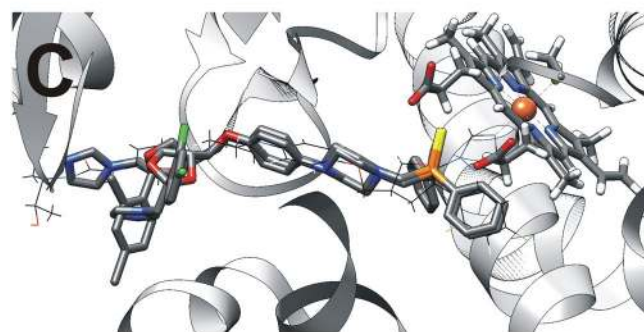
Ke:
-11.6 kcal/mol
Fe-N: 2.814 Å

posaconazole:
-13.2 kcal/mol
Fe-N: 3.960 Å



KeP:
-13.5 kcal/mol
Fe-N: 2.802 Å

KeOP:
-12.7 kcal/mol
Fe-N: 2.960 Å



KeSP:
-12.4 kcal/mol
Fe-S: 4.285 Å

KeSeP:
-12.2 kcal/mol
Fe-Se: 4.485 Å

Figure 6. Docking results: heme of the *C. albicans* CYP51 and the best-fit docked compounds; posaconazole position from the original crystal structure¹⁵ is also indicated (as a black wire). (A) docked posaconazole, (B) Ke, KeP, KeOP, C: KeSP, KeSeP.

differences between their binding strength, also in comparison with ketoconazole, proving that they are capable of inhibiting this enzyme. While **KeP** and **KeOP**, like **Ke**, form a bond between nitrogen atom from the azole ring and the heme iron atom, for **KeSP** and **KeSeP**, a reversed placement in the hydrophobic cage was the preferred one. This opens a possibility of a different/secondary action mechanism for these compounds and will be studied in detail.

In what concerns cytotoxicity towards hACs, the individual compounds had displayed some activity, except for **KeSP** which was not toxic. Two ketoconazole derivatives were further studied (**KeP** and **KeOP**) and in both cases we observed an increased level of the p21 gene transcript, but no changes in the level of p53 transcript. A decreased mitochondrial membrane potential was also apparent. This may indicate that all the new ketoconazole derivatives have a similar mechanism of action and block the lanosterol 14 α -demethylase and thus inhibit the

production of ergosterol in *C. albicans* membranes. The IC₅₀ value in the cytotoxicity assay with human cells was significantly higher than the MIC₅₀ for *C. albicans*, especially when the compounds were used with fluconazole. Therefore, we conclude that phosphane derivatives of ketoconazole are a promising group of compounds with potential use in the treatment of mycoses.

Taking into account all our results, we conclude that **Ke** based phosphane is a very interesting ligand and enables a significant expansion of possible metal complexes bearing the **Ke** moiety. Currently, we are working on copper(I) and ruthenium(II) complexes with **KeP**. Moreover, we are going to study phosphanes bearing more than one **Ke** moieties.

Experimental Part

Syntheses. Reagents and solvents used for synthesis were purchased from Sigma-Aldrich, Poland and used without further purifications. All syntheses were performed under inert (N₂) atmosphere using standard Schlenk techniques. Deacylated ketoconazole (**KedA**; 1-4-[2-(2,4-Dichlorophenyl)-2-imidazol-1-ylmethyl-[1,3]-dioxolan-4-ylmethoxy]phenyl-piperazine; M = 489.4 g/mol) and PPh₂(CH₂OH)₂Cl (M = 282.7 g/mol) were prepared following literature procedures^{86,87}.

Ph₂PCH₂-Ke (KeP). Triethylamine (3 mL) was added drop wise to a solution containing 0.3207 g of PPh₂(CH₂OH)₂Cl (1.134 mmol) in 10 mL of MeOH in an ice bath. After stirring for 30 min, an MeOH solution (15 mL) of **KedA** (0.5550 g; 1.134 mmol) was added. The mixture was stirred for 1 h at room temperature and its volume was reduced by 1/3 of initial volume under a stream of nitrogen. Then 100 mL of water was slowly added. The obtained white solid was filtered, washed five times with water and dried under reduced pressure for 24 h. Yield 61%. Anal. Calc. for C₃₇H₃₇Cl₂N₄O₃P (M = 687.6 g/mol): C, 64.63; H, 5.42; N, 8.15. Found: C, 64.34; H, 5.45; N, 8.11%. MS ESI (+): 489.2 [KedA + H]⁺ (6%), 501.2 [KedA + CH₂]⁺ (100%), 709.2 [M + Na]⁺ (15%).

Ph₂P(O)CH₂-Ke (KeOP). **KeOP** was obtained in a dichloromethane-acetonitrile mixture (1:1) (10 mL) through the reaction of **KeP** (0.1081 g; 0.157 mmol) placed in an ice bath with an equimolar amount of H₂O₂ (30% solution in water: 16 μL; 0.157 mmol). After stirring for 30 min, the solvent was evaporated under vacuum. The remaining white solid was washed twice with water and dried for 24 h under vacuum to remove any occluded solvent molecules. Yield 94%. Anal. Calc. for C₃₇H₃₇Cl₂N₄O₃P (M = 703.6 g/mol): C, 62.79; H, 5.33; N, 7.92. Found: C, 62.83; H, 5.33; N, 7.92%. MS ESI (+): 703.2 [M + H]⁺ (6%), 725.2 [M + Na]⁺ (100%), [2M + Na]⁺ 1429.4 (5%).

Ph₂P(S)CH₂-Ke (KeSP). To **KeP** (0.0983 g; 0.143 mmol) in chloroform (15 mL) an equimolar amount of resublimed sulfur (0.00457 g; 0.143 mmol) was added. The mixture was placed in an ultrasonic bath for 1 h at room temperature. The yellowish solid obtained after removing the solvent was redissolved in a minimal volume of methanol and precipitated with large amounts of water. The solid obtained was filtered and dried for 24 h under vacuum. Yield 68%. Anal. Calc. for C₃₇H₃₇Cl₂N₄O₃PS (M = 719.7 g/mol): C, 61.75; H, 5.18; N, 7.78. Found: C, 61.40; H, 5.22; N, 7.74%. MS ESI (+): 501.2 [KedA + CH₂]⁺ (13%), 719.2 [M + H]⁺ (29%), 741.2 [M + Na]⁺ (100%) 1439.4 [2M + H]⁺ (51%), 1461.4 [2M + Na]⁺ (32%).

Ph₂P(Se)CH₂-Ke (KeSeP). An equimolar mixture of **KeP** (0.1026 g; 0.149 mmol) and metallic selenium (0.01178 g; 0.149 mmol) in chloroform (15 mL) was placed in an ultrasonic bath until the traces of solid Se disappeared (~1 h). The yellowish solid obtained after removing the solvent was redissolved in a minimal volume of methanol and precipitated with a large amount of water. The solid obtained was filtered and dried for 24 h under vacuum. Yield 73%. Anal. Calc. for C₃₇H₃₇Cl₂N₄O₃PSe (M = 766.6 g/mol): C, 57.97; H, 4.87; N, 7.31. Found: C, 57.66; H, 4.90; N, 7.31%. MS ESI (+): 501.2 [KedA + CH₂]⁺ (19%), 767.1 [M + H]⁺ (48%), 789.1 [M + Na]⁺ (100%), 1533.3 [2M + H]⁺ (98%), 1555.3 [2M + Na]⁺ (30%).

Methods. Elemental analyses were performed on a Vario EL3 CHN analyzer for C, H, and N, and they were within ±0.3% of the theoretical values. NMR spectra were recorded on a Bruker AMX 500 spectrometer (at 298 K) with traces of solvent as an internal reference for ¹H (CDCl₃: 7.27 ppm) and ¹³C spectra (CDCl₃: 77.0 ppm) and 85% H₃PO₄ in H₂O as an external standard for ³¹P. All the NMR data are presented in ESI (Tables S1, S2) Mass spectra were recorded on a Bruker Daltonics micrOTOF-Q mass spectrometer equipped with an electrospray ionization (ESI) source and operated in positive ion mode.

Saccharomyces cerevisiae. The following strains of *S. cerevisiae* were used in this study: BY4741 (wild type towards *erg6Δ*, ACC Y00000; MATα; *his3Δ1*; *leu2Δ0*; *met15Δ0*; *ura3Δ0*), *erg6Δ* (ACC Y00568; BY4741 isogenic with YML008c::*kanMX4*), W303 (wild type towards AD1-8; MATα; *leu2-3,112 trp1-1 can1-100 ura3-1 ade2-1 his3-11,15 AD1-8* (W303 isogenic with *pdr1-3, ura3, his1, yor1Δ::hisG, snq2Δ::hisG, pdr5Δ::hisG, pdr10Δ::hisG, pdr11Δ::hisG, ycf1Δ::hisG, pdr3Δ::hisG, pdr15Δ::hisG*). BY4741 and *erg6Δ* were acquired from EUROSCARE, Frankfurt, Germany; W303 and AD1-8 were kind gifts from Prof. A. Goffeau, Chaire Internationale Blaise Pascal, Lab. Genet. Mol., École Normale Supérieure, Paris, and Prof. M. Ghislain, Université Catholique de Louvain, Faculté des sciences Agronomiques, Louvain-la-Neuve, Belgium. Visible turbid overnight cultures of *S. cerevisiae* strains were prepared and diluted to A₆₀₀ ≈ 0.1 in fresh Yeast Extract – Peptone - Dextrose (YPD) medium. The cultures were transferred to 96 microtiter well plates containing aliquots of serially diluted antifungal drugs (**Ke** and its derivatives: **KeP**, **KeOP**, **KeSP**, **KeSeP**), prepared from DMS stock solutions and drug-free controls. The yeast cultures were statically incubated at 30 °C for 48 hours with compounds⁸⁸. The dose-response curves were analysed with GraphPad Prism 5 software (San Diego, USA) or a microplate reader ASYS UVM (Biogenet). All values presented are the mean of at least three independent experiments.

Candida albicans. *C. albicans strains and growth conditions.* The *C. albicans* CAF2-1 (Genotype/Description: *ura3Δ::imm434/URA3*⁸⁹) and DSY1050 (Genotype/Description: *cdr1Δ::hisG/cdr1Δ::hisG/cdr2Δ::hisG/cdr2Δ::hisG/mdr1Δ::hisG-URA3-hisG/mdr1Δ::hisG*⁹⁰) strains used in this study were generous gift from D. Sanglard (Lausanne, Switzerland)^{91,92} whereas *C. albicans* B3 and Gu4 (Genotype/Description: clinical isolates fluconazole sensitive)⁹³, B4 (Genotype/Description: Clinical isolate fluconazole resistant due to the overexpression of *MDR1*⁹³) and Gu5 (Genotype/Description: Clinical isolate fluconazole resistant due to the overexpression of *CDR1* and *CDR2*⁹³) were a generous gift from prof. S. Milewski (Gdańsk, Poland)⁹⁴ and prof. J. Morschhäuser (Wurzburg, Germany)⁹³. All strains were grown at 37 °C on Sabouraud medium (Biomerieux).

Minimal inhibitory concentration (MIC) determination. The antifungal activity was determined on the 96-well polystyrene microtiter plates (Sarstedt) according to Clinical and Laboratory Standards Institute M27-A3⁹⁵. The different strains were incubated with antifungal compounds (fluconazole (**Flc**) (Sigma-Aldrich), **Ke** and **Ke** derivatives) in the range of concentrations 0.01–200 μM. The compounds were dissolved in DMSO (10 mM) and then serially diluted in RPMI-1640 (ThermoFisher) medium to obtain the final concentration range. Inocula of all strains were prepared as suspensions with turbidity of 0.5 McFarland standard in sterile 0.85% saline and diluted in RPMI-1640. After inoculation, each well contained appr. 5×10^2 CFU/mL. The plates were cultivated for 48 h at 37 °C. Optical density was measured at A_{490nm} using a microplate reader ASYS UVM 340 (Biogenet). The concentrations which resulted in $\geq 50\%$ growth inhibition were determined as MIC₅₀.

KeOP/KeP synergism with fluconazole. The synergistic effects of **KeOP/KeP** with **Flc** were determined using the M27-A3 method with modifications. **Flc** and **KeOP/KeP** stock solutions in DMSO (10 mM) were (alone and simultaneously) diluted in RPMI-1640 medium to obtain final concentrations of $1/8, 1/4, 1/2, 1 \times$ MIC₅₀ towards *C. albicans* CAF2-1 strain in microdilution sterile plates (Sarstedt). Statistical significance analysis was performed using Student's *t*-test (binomial, unpaired) comparing viability in the presence of corresponding **Flc** concentrations with/without **KeOP/KeP**.

Human Adipose Tissue-derived Stem Cells (hASCs). Unless indicated otherwise, all reagents used in this study were purchased from Sigma-Aldrich (Poland).

Cells isolation and culture. Human Adipose Tissue-derived Stem Cells (hASCs) were isolated from subcutaneous adipose tissue fragments harvested from non-diabetic female donors (age range 33–38, $n = 2$) during standard surgical procedure of total hip arthroplasty. Immediately after collection of tissue, the samples were placed in sterile tissue-transport medium (Hank's Balanced Salt Solution (HBSS) supplemented with 1% Penicillin/Streptomycin/Amphotericin B solution (P/S/A). Mesenchymal stromal cell isolation was performed using enzymatic digestion of extracellular matrix with collagenase type I. Before digestion, tissue specimens were washed twice with HBSS and incubated in collagenase type I solution (1 mg/mL) at 37 °C for 40 min. The digest was subsequently centrifuged ($1200 \times g$; 10 minutes; room temperature) and the cell pellet was resuspended in Dulbecco's modified Eagle's medium (DMEM) with Nutrient F-12 Ham, 10% of Fetal Bovine Serum (FBS), and 1% of P/S/A, transferred to a culture flask, and kept in culture under optimal conditions (37 °C in a humidified atmosphere of 5% CO₂). The medium was refreshed every 2–3 days. At 80–90% confluence, the cells were passaged using trypsin solution (TrypLE Express, Life Technologies, California, USA). At passage 3, hASCs were collected and cellular phenotype was confirmed by the high expression of markers CD44, CD29, CD73, CD105 and low expression of CD45. Moreover, the trilineage differentiation capacity of ASCs was confirmed as previously described⁹⁶.

Half maximal inhibitory concentration (IC₅₀) determination. The IC₅₀ values were determined using *in vitro* Toxicology TOX-8 Assay Kit (resazurin based) according to the manufacturer's protocol. For this assay, cells were seeded in a 96-well plate at a density of 5×10^3 cells/well. The cells were pre-incubated in drug-free complete medium for 24 h before adding various concentrations of the compounds to be investigated. Stock solutions of the compounds were prepared in DMSO following serial dilutions in DMEM with Nutrient F-12 Ham complete medium. The cells exposure period was 24 h. After that time media were replaced with 10% v/v resazurin dye solution in fresh complete culture medium. Incubation was carried out for 2 h at 37 °C in the CO₂ incubator. Reduction of the dye was measured spectrophotometrically at 600 nm and 690 nm reference wavelength (Epoch, Biotek). Absorbance measurements allowed the determination of viable treated cells relative to untreated controls using the inflection point of a dose-response graph. IC₅₀ values were estimated as duplicates of triplicate readings in two independent experiments.

Evaluation of cellular apoptosis. Cellular apoptosis level was assessed using quantitative reverse transcriptase real-time polymerase chain reaction (qRT-PCR). First, the cells were seeded in a 24-well plate at a concentration of 3×10^4 cells/well in DMEM with Nutrient F-12 Ham complete medium and left in the CO₂ incubator overnight to allow cells to attach. The compounds (**Ke**, **KeP** and **KeOP**) were diluted in the fresh complete medium to a final concentration of 20 μM. After 24 h of challenge with the compounds, the cells were homogenized with TRI Reagent and total RNA was isolated by phenol-chloroform extraction and ethanol precipitation⁹⁷. RNA purity and quantity were measured at a wavelength of 260 and 280 nm (Epoch, Biotek). Genomic DNA was removed with DNase I, RNase-free (1 U/L) (Thermo Scientific, Poland) and cDNA was synthesized using RevertAid First Strand cDNA Synthesis Kit (Thermo Scientific, Poland). 150 ng of total RNA served as a template for a single reaction using T100 Thermo Cycler (Bio-Rad, USA). The expression levels of p53 tumor suppressor, cyclin dependent kinase inhibitor 1A (p21), apoptosis regulators BAX and BCL-2 were determined. qRT-PCR was performed on a CFX Connect™ Real-Time PCR Detection System (Bio-Rad) using SensiFast SYBR & Fluorescein Kit (Bioline, Cincinnati, OH, USA) and primer pairs as listed in Table S4 in ESI. 500 nM of specific primers and 1 μL of cDNA

in a 10 μ L final volume were applied in each reaction. The quantitative expression of the genes was calculated by the $2^{-\Delta\Delta CT}$ method using GAPDH as housekeeping genes.

Statistical analysis was performed with GraphPad Prism 5 software (San Diego, USA). All data values are presented as mean \pm SD, measured in triplicates or more. Statistical significance between groups was determined using two-way ANOVA grouped analysis.

Confocal microscopy imaging. The mitochondrial network was visualized using MitoRed staining. After 24-hours treatment with either **Ke**, **KeP** or **KeOP**, the media were removed and replaced with MitoRed dye in 1:1000 dilution in fresh complete media. Cells were incubated for 30 minutes at 37 °C, then washed with Phosphate Buffered Saline (PBS), fixed with 4% paraformaldehyde (PFA) solution, washed three times and nuclei were counterstained with diamidino-2-phenylindole (DAPI).

In order to perform F-actin imaging, cells were fixed with 4% PFA solution, washed with PBS and then permeabilized using 0.5% TX-100 (incubation 15 minutes, room temperature). The cells were rinsed and incubated with Phalloidin-Atto 590 solution in PBS (dilution 1:1000) at 37 °C for 30 minutes in the dark. After washing, the cells were counterstained with DAPI.

Cells were observed and images were taken using confocal microscopy performed with a Leica TCS-SP8 laser scanning confocal microscope. Data were further processed with the use of ImageJ software.

Docking studies. Docking calculations were performed using the X-ray structure of *C. albicans* CYP51 in complex with posaconazole¹⁵ (PDB code 5FSA) as the macromolecular host, and the set of ketoconazole derivatives as the ligands. Posaconazole was also included in the set as a validation test. The DFT-optimized structures (the Gaussian 16, Rev. B.01 program⁹⁸ with ω B97XD functional⁹⁹ and the 6-311 + G(d,p) basis set) served as initial ligand structures. The macromolecule consisted of the protein chain A, together with the heme cofactor, but without posaconazole. The docking of the ligands was carried out using a cubic box with an 80 Å edge, centered on the iron atom of the heme cofactor, as the search space; up to nine structures for each search were retained for further analysis. For two of the ligands (**KeSP**, **KeSeP**), docking runs were also repeated with a smaller box (a = 40 Å). A docking run for each ligand was replicated three times with different random seeds, and generally the results were consistent between the replicas. Preprocessing of the structures and visualization of the results was carried out with AutoDockTools program of the MGLTools package¹⁰⁰ and UCSF Chimera 1.11.2_b41376 graphic program¹⁰¹, while the docking itself was performed with the AutoDock Vina software ver. 1.1.2⁸⁵.

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Author contributions

R.S., R.A. and A.K. designed the scientific rationale; R.A. designed *S. cerevisiae* related experiments; F.S. performed *S. cerevisiae* related experiments, analyzed data and generated figures; A.K. designed *C. albicans* related experiments; J.S. performed *C. albicans* related experiments, analyzed data and generated figures; A.J. and J.P. performed molecular docking experiments, analyzed data and generated figures; R.S. administrated the project, performed syntheses and spectroscopic characterization of the compounds, analysed the data and generated figures; K.M. designed hASCs related experiments; M.A. performed hASCs related experiments, analyzed data and generated figures; R.S., R.A., K.M., A.K., J.S., J.P. and A.J. wrote the manuscript. The manuscript was reviewed by all Authors prior to submission.

Competing interests

The authors declare no competing interests.

Additional information

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