ABSTRACT: Chagas disease, which was once thought to be confined to endemic regions of Latin America, has now gone global, becoming a new worldwide challenge with no cure available. The disease is caused by the protozoan parasite *Trypanosoma cruzi*, which depends on the production of endogenous sterols, and therefore can be blocked by sterol 14α-demethylase (CYP51) inhibitors. Here we explore the spectral binding parameters, inhibitory effects on *T. cruzi* CYP51 activity, and antiparasitic potencies of a new set of β-phenyl imidazoles. Comparative structural characterization of the *T. cruzi* CYP51 complexes with the three most potent inhibitors reveals two opposite binding modes of the compounds (\((R)-6, EC_{50} = 1.2 \text{ nM}\), vs \((S)-2/(S)-3, EC_{50} = 1.0/5.5 \text{ nM}\)) and suggests the entrance into the CYP51 substrate access channel and the heme propionate-supporting ceiling of the binding cavity as two distinct areas of the protein that enhance molecular recognition and therefore could be used for the development of more effective antiparasitic drugs.

**INTRODUCTION**

American trypanosomiasis (Chagas disease) is a vector-borne anthropozoonosis, the life-long infection caused by the protozoan pathogen *Trypanosoma cruzi*. For centuries the disease has been a major cause of mortality and morbidity in South and Central America, where it still remains endemic in 21 countries, resulting in about 14,000 deaths per year, mainly due to heart failure, that is the most typical pathology of the chronic form of this infection. During the last years, Chagas disease has begun receiving attention as an emerging global medical problem, predominantly because of human and insect vector (kissing bug) migration, but also as a result of the lack of awareness and diagnostics in nonendemic areas, often leading to transfusion of infected blood; transplantation of infected organs, food, and drink contaminations; as well as HIV coinfections and congenital transmission. Thus, recent estimates indicate that there could be up to 1 million cases of Chagas disease in the USA, a significant portion of which, particularly in the southern states, are of autochthonous vector-borne origin.
Despite the severity of the problem and ~25 million people at risk of infection, nifurtimox and benznidazole are the only drugs available for treatment of Chagas disease. Although generally rather helpful in the acute stage, they vary significantly in their efficiency against different T. cruzi strains, they have considerable adverse side effects, and their success in curing chronic Chagas disease is still debated.10,11 Despite these limitations, pharmaceutical companies remain reluctant to invest resources in the development of new antischagasic chemotherapies, because of the lack of assurances that they can make a return on their investment, since historically Chagas disease has been known to mostly affect the poorest. As a result, the majority of efforts to fill the gap for new antischagasic drugs have come from academia. Repurposing of antifungal azoles, the drugs that act via sterol biosynthesis by inhibiting the cytochrome P450 enzyme sterol 14α-demethylase (CYP51)12 so far have been particularly successful (reviewed in refs 1 and 13). The FDA-approved drug posaconazole and an investigational prodrug of ravuconazole (Eisai) are presently in phase 2 clinical trials for Chagas.3 Some other azole derivatives, e.g. the anticancer drug candidate tipifarnib14,15 or (S)-2-((1-(biphenyl-4-ylmethyl)-1H-imidazol-5-yl)methylamino)-biphenyl-2-ylcarboxamido)-4-(methylthio)butanoic acid (FTI-2220),16 were also shown to display potent antiparasitic effects and are under development. Later, several new experimental heterocyclic compounds, both azoles17,18 and pyridines19,20 were identified as potent and selective inhibitors of T. cruzi CYP51 (the protein has less than 25% amino acid sequence identity to its fungal orthologs21) and structurally characterized in complex with the target enzyme.18,20,22,23 Most recently, one of these inhibitors, VNI, has been shown to cure both the acute and chronic forms of Chagas disease in mice.24

Being the most potent T. cruzi CYP51 inhibitors that we have discovered,27 VNI and VNF share high structural similarity. The phenethylimidazole portion of these molecules is connected via the polar linker (carboxamide fragment) to the lipophilic arm that consists of either a 2-ring (VNF) or a 3-ring (VNI) linear polycycle (Figure 1A). In the CYP51 costructures, VNI and VNF are coordinated to the P450 heme iron through their imidazole ring nitrogen (N3). The other two portions of the inhibitor molecules, however, adopt an opposite orientation,25 while the 3-ring arm of VNI lies in the CYP51 substrate access channel, the 2-ring arm of VNF is positioned within the deepest segment of the CYP51 binding cavity, the hydrophobic area that accommodates the aliphatic tail of the sterol substrate (Figure 1B).

In this work we prepared a set of 12 β-phenyl imidazoles and analyzed their structure–activity relationship in terms of T. cruzi CYP51 binding parameters, inhibition of reconstituted enzymatic reaction in vitro, and antiparasitic effects against GFP-expressing T. cruzi amastigotes. Three most efficient compounds were co-crystallized with T. cruzi CYP51, the X-ray costructures uncovering two basic approaches that can be utilized to further enhance potencies of CYP51 inhibitors.

## RESULTS AND DISCUSSION

**Medicinal Chemistry.** Compound 1 (MW 368, clogP 5.3, tPSA 41.9 (ChemDraw)) has been previously characterized as a potential antifungal agent and revealed quite promising results.25 Its structural resemblance to VNF has prompted us to expand our work on this inhibitory chemotype by modifying the chemical structure of 1 as shown in Table 1 followed by testing the original molecule and its derivatives against T. cruzi and its potential target enzyme T. cruzi sterol 14α-demethylase. In all cases the polar linker between the phenethylimidazole moiety and the opposite arm of the new structures was replaced with the carbamate group, because it was previously found to have higher hydrolytic stability in liver microsomes than the ester group of 1.26 Our major focus on modification of the side chain arm of the compounds was based on the observation that variations in the composition of this portion of a β-phenyl imidazole molecule (a) could be crucial for its potency to inhibit CYP51 activity17 and (b) may alter its orientation within the enzyme active site.25 In compounds 2 (MW 384, clogP 4.9, tPSA 53.9) and 3 (MW 367, clogP 4.3, tPSA 53.9), the side chain arm is one aromatic ring shorter than it is in 1, and in the para-position of the β-phenyl ring they have either Cl atom (2) or smaller and more polar F atom (3). Compound 4 (MW 357, clogP 1.8, tPSA 69.5), similarly to 3, also has fluorine in the para-position of the β-phenyl ring; however, its side chain arm, instead of the bulky aromatic ring, carries a flexible three-carbon atom aliphatic chain ending with the polar imidazole ring. The arm of compound 5 (MW 494, clogP 5.6, tPSA...
105.5) bears two aromatic rings linked via the sulfur atom, with the distal ring ending with a nitro group in the para-position. Compound 6 (MW 605, clogP 7.2, tPSA 60.4) is the largest of the molecules. It has two chlorine substituents in the ortho- and para-positions of the β-phenyl ring, while its long arm is composed of three rings linked in a linear sequence, with the distal aromatic ring being complemented with two chlorine atoms, in the 3- and 4- positions so that the total length of the arm (∼18 Å) is close to the length of the CYP51 substrate access channel.22

In this study the compounds were prepared as (R)- and (S)-stereoisomers using enantioselective synthesis.26 Briefly, the 2-(1H-imidazol-1-yl)-1-phenylethanones were obtained by condensation of 1H-imidazole and commercial bromoacetophenones25 substituted in the ortho/para position with different halogens (Scheme 1a). In order to obtain the (S)-2-(1H-imidazol-1-yl)-1-phenylethanols, the keto group of 2-(1H-imidazol-1-yl)-1-phenylethanones was reduced to hydroxyl using as the catalyst RuCl(p-cymene)[(R,R)-Ts-DPEN] (Scheme 1b). Accordingly, RuCl(p-cymene)[(S,S)-Ts-DPEN] has been used as the catalyst to prepare the (R)-2-(1H-

Table 1. Structural Formulas, T. cruzi CYP51 Spectral Binding Parameters, Inhibition of Enzymatic Activity, and Antiparasitic Effects of Compounds 1−6a

| Compound | Structure | CYP51 spectral response | CYP51 substrate conversion at 1/E=2 (%)b | T. cruzi amastigotesc (%) |
|----------|-----------|-------------------------|------------------------------------------|---------------------------|
|          | Amax/cm/ nMol P450 (optical units) | Ks (µM) | Binding efficiency (ΔAmax/Ks) | EC50 (nM) | Inhibition at 10 nM (%) |
| 1        | (R)-     | 0.042±0.001 0.087±0.02 0.5 | 79.5±1.5 | 15±0.3 | 29.5±2.1 |
|         | (S)-     | 0.043±0.001 0.046±0.01 0.9 | 0.7±0.2 | 1.2±0.1 | 76.3±7.1 |
| 2        | (R)-     | 0.039±0.001 0.056±0.01 0.7 | 90.0±5.0 | 22±1.7 | 30.5±3.5 |
|         | (S)-(LFT)c | 0.044±0.001 0.032±0.01 1.4 | 1.9±0.2 | 1.0±0.1 | 74.5±13.4 |
| 3        | (R)-     | 0.058±0.002 0.083±0.02 0.6 | 86.5±6.5 | 20±2.5 | 32.1±2.8 |
|         | (S)-(LFS)c | 0.047±0.001 0.032±0.01 1.4 | 8.5±3.5 | 5.5±0.7 | 69.3±4.2 |
| 4        | (R)-     | 0.030±0.001 0.087±0.02 0.3 | 89.2±4.8 | Nd      | Nd      |
|         | (S)-     | 0.030±0.0002 0.600±0.02 0.1 | 93.9±3.8 | Nd      | Nd      |
| 5        | (R)-     | 0.063±0.09 0.715±0.29 0.1 | 52.5±2.5 | 2.5±0.1 | 62.5±3.5 |
|         | (S)-     | 0.046±0.004 0.162±0.08 0.3 | 12.5±1.5 | 1.5±0.1 | 70.0±14.1 |
| 6        | (R)-(LFD)c | 0.042±0.001 0.064±0.03 0.7 | 0.8±0.1 | 1.2±0.1 | 83.0±9.9 |
|         | (S)-     | 0.047±0.006 0.123±0.11 0.4 | 3.8±0.4 | 1.3±0.1 | 73.5±7.8 |

The values represent mean ± standard deviation from three independent experiments.1 h reaction (examples of HPLC profiles for compounds 2 and 6 are shown in Figure 3). The corresponding values for posaconazole are EC50 = 5.0 nM ; inhibition at 10 nM = 64%. Compounds cocrystallized with T. cruzi CYP51.
imidazol-1-yl)-1-phenylethanols. The −OH group was subsequently deprotonated with sodium hydride (c) or activated with triphosgene (d) in anhydrous CH$_3$CN in order to obtain sodium alkoxide and chloroformate. Then commercial 4-isopropylphenyl isocyanate, 2-(1H-imidazol-1-yl)propan-1-amine and 4-((4-nitrophenyl)thio)aniline have been added for preparing the side chain arms of compounds 2/3, 4, and 5, respectively. (Scheme 1 ef).

The long arm of compound 6 has been synthesized as shown in Scheme 2. The synthesis was performed by condensation of 1-fluoro-4-nitrobenzene with 1-(3,4-dichlorophenyl)piperazine (a). Then the nitro group of 1-(3,4-dichlorophenyl)-4-(4-nitrophenyl)piperazine was converted to an amino group by reaction with H$_2$ using Pd/C as the catalyst (b) to produce 4-[4-(3,4-dichlorophenyl)piperazin-1-yl]aniline; and the final amine was stabilized as 4-[4-(3,4-dichlorophenyl)piperazin-1-yl] anilinium chloride by HCl$_{(g)}$ flow in the same reaction environment (c).

_T. cruzi_ CYP51 **Spectral Responses vs Inhibition of Reconstituted Sterol 14α-Demethylase Activity in Vitro.**

In the resting ferric form the heme iron of CYP51 enzymes is present in the low spin state, with a water molecule serving as its sixth axial (distal) ligand. This results in the P450 absorbance spectrum with the Soret band maximum at around 417 nm. Binding of azoles or other heterocyclic compounds replaces the water molecule in the heme iron coordination sphere with the basic nitrogen atom, causing a so-called red shift in the Soret band maximum. Accordingly, in the difference spectra a trough and a peak appear on the left and the right sides of an isobestic point. These spectral changes, also known as a type 2 spectral response, are widely used to identify new P450 binding ligands. However, even very low spectral dissociation constants do not necessarily reflect the compound’s potency to inhibit CYP51 activity, because during the reaction many ligands can still be replaced in the enzyme active center by the substrate.\textsuperscript{17,20,22}

Twelve imidazole derivatives tested in this study act in good agreement with this observation. As expected, all of them cause typical type 2 spectral responses in the CYP51 heme iron, with the apparent spectral dissociation constants ($K_s$) being mostly in the nanomolar range, and therefore should be defined as tight binding ligands (Table 1, Figure 2). However, particularly in the case of compounds 1–3, the inhibitory effects of the (R)- and (S)-enantiomers on the CYP51 activity appeared to be quite “independent” of the binding parameters. Thus, (S)-2 displayed only 2-fold higher apparent binding efficiency than (R)-2, yet its inhibitory effect on the substrate conversion is ~50-fold stronger (Table 1, Figure 3A). The same tendency was observed for 1 and 3. In all these three cases, the (S)-enantiomers are much more potent as _T. cruzi_ CYP51 inhibitors than the (R)-enantiomers. The (R)- and (S)-enantiomers of compounds 4–6, on the opposite, do not differ that drastically in their potencies to inhibit _T. cruzi_ CYP51 reaction (Table 1, Figure 3B), although significant variations can be seen in the apparent $K_s$’s, particularly for 4 ((S) $\gg$ (R)). Only the enantiomers of compound 6 presented
rather good correlation between the CYP51 inhibition and binding parameters. Nevertheless, their spectral dissociation constants are either higher ((S)-, \( K_s = 0.123 \ \mu M \)) or within ((R)-, \( K_s = 0.064 \ \mu M \)) the range of the values calculated for the (R)- enantiomers of compounds 1–3 (weak \( T. cruzi \) CYP51 inhibitors).

Another unusual observation, which we encountered upon titration of \( T. cruzi \) CYP51 with compound 6 (both (R)- and (S)-enantiomers), was the very slow increase in the amplitude of the type 2 spectral response over time. This increase was most pronounced when the enzyme concentration considerably exceeded the concentration of the ligand. To our knowledge, such an increase has not been previously reported for any other CYPs. Since none of the other compounds tested in this work produced such an effect, we conducted the same experiments with other known \( T. cruzi \) CYP51 inhibitors, and we found that a somewhat similar pattern is also produced by ketoconazole, another CYP51 inhibitor with a rigid long arm but not by VNI (Figure 4). The current lack of any information on the topic might be related to the fact that traditionally the measurements of P450 spectral responses have been mostly conducted at higher ligand/P450 molar ratios.

**Crystallographic Analysis.** In order to expand our understanding of the molecular basis that underlies the potencies of CYP51 inhibitors as well as to shed light on the features that may have caused the observed peculiarities in their binding behavior, we crystallized \( T. cruzi \) CYP51 in the presence of (S)-2, (S)-3, and (R)-6, which are among the strongest inhibitors of the enzyme identified in this study (Table 1), and we determined the X-ray structures of their complexes. Table 2 summarizes the diffraction and refinement statistics. As expected, (R)-6 (ligand PDB ID LFD, molecular volume 1,040 Å³, surface area 880 Å²) adopts a VNI-like orientation, with its longer arm protruding ∼7 Å further toward the entrance into the substrate access channel (Figure 5A), while the vectors of (S)-2 (ligand PDB ID LFT, molecular volume 870 Å³, surface area 725 Å²) and (S)-3 (ligand PDB ID LFS, molecular volume 849 Å³, surface area 709 Å²) follow that of VNF, being directed toward helix C (Figure 5B). The amino acid residues that contact each inhibitor are listed in Table 3, and their location in the \( T. cruzi \) CYP51 structure is seen in Figure 5.

Overall, the structures suggest that the potency of LFD (20 inhibitor contacting residues) must be enhanced by the formation of the surface binding subsite (Figure 6A–C, Supporting Information Figure S1), the rearrangement that was first observed in the structure of posaconazole-bound \( T. cruzi \) CYP51 [3K1O].23 When two molecules of CYP51-LFD

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**Figure 2.** Spectral responses of \( T. cruzi \) CYP51 to the binding of (A) (R)- and (S)-enantiomers of compound 2; \([P450] = 1.1 \ \mu M \) and (B) (R)- and (S)-enantiomers of compound 6; \([P450] = 0.9 \ \mu M \). Optical path length 5 cm. Upper: absolute absorbance spectra, the Soret band maximum shifts to the right (from 418 to 425 nm). Lower: difference spectra upon titration with the ligands (titration step 0.2 \( \mu M \)). Insets: titration curves showing absorbance changes per 1 cm optical path/1 nmol P450 upon increasing ligand concentrations (processed with the Morrison equation).
(this structure has two molecules in the asymmetric unit (Table 2)) are superimposed with CYP51-LFT and CYP51-LFS, it is clearly seen that in the complex with LFD the secondary structural elements that form the entrance into the CYP51 substrate access channel are getting better organized, particularly helix F″, whose density in the complexes of T. cruzi CYP51 with smaller molecules is either missing (VNF [3KSW], fluconazole [3KHM], NEU [4H6O]) or appears to be more looplike (LFT and LFS (Figure 6C)). Moreover, some residues, e.g. F214, P210 (F″), I45 (A′), I72 (β1-1), move 1−1.5 Å closer toward the LFD 1,2-dichlorobenzene ring (Figure 6B), which, as a result, forms van der Waals contacts with 8 amino acids around the channel entrance (a possible example of a "localized" induced fit in CYP51). These multiple contacts between the enzyme and inhibitor are clearly decreasing the entrance flexibility (Supporting Information Figure S2A) and may prevent the CYP51 channel from opening, making it harder for the substrate molecule to replace the inhibitor, e.g. upon P450 reduction. On the other hand, LFD does not induce any significant rearrangements around the CYP51 heme binding area, which might explain why the (R)- and (S)-enantiomers of this compound would have similar inhibitory potencies and display similar apparent binding efficiencies. The observed time-dependent increase in the spectral response of the P450 heme iron to the binding of the LFD nitrogen might be connected with some repositioning of the proximal to the heme portion of the inhibitor molecule, as even in the crystal lattice the two medium rings of this compound display obvious differences in their conformations (Figure 6B,E).

Although only 16 of the T. cruzi CYP51 amino acid residues lie at the distance <4.5 Å from LFT and LFS (Figure 6B, Table 3), which are much smaller molecules than LFD, their complexes with the enzyme are strengthened by the H-bonds with Y116. In both these structures the side chain hydroxyl of Y116 looses its contact with the heme ring D propionate, shifts 1.5 Å toward the inhibitor, and interacts with the O and N atoms of its carbamic fragment (Figure 6E,F). This change in the Y116 position increases the flexibility of the heme (Supporting Information Figure S2B) and allows the p-cymene moiety of the inhibitors to protrude deeper into the CYP51 binding cavity. Enhancement of the LFT/LFS complexes with CYP51 by the H-bond formation elucidates the importance of (S)-stereochemistry for the inhibitory potency of compounds 2 (LFT) and 3 (LFS), because the opposite orientation of the carbamic fragment in the (R)-enantiomers is not favorable for hydrogen bonding. It also provides a possible explanation for the observed “discrepancies” between the spectral binding...
parameters and inhibition (Table 1), since the H-bond formation might not influence the spectral response of the P450 heme iron to the imidazole nitrogen coordination, which apparently occurs fast and easily for both (R)- and (S)-enantiomers. Finally, combination of biochemical and structural data implies that compounds 4 and 5 are most likely to bind to T. cruzi CYP51 in the LFD orientation: (1) the arm of 5 (≈15 Å) is probably too long and the imidazole ring of 4 is too polar to fit well into the hydrophobic deepest segment of the cavity, (2) relatively small differences between the potencies of their (R)- and (S)-enantiomers suggest the lack of H-bonding.

**Antiparasitic Effects in T. cruzi Cells.** T. cruzi is a unicellular eukaryotic parasite with the complex life cycle involving insect vectors and mammalian hosts. In insects, multiplying midgut epimastigotes transform into metacyclic trypomastigotes that infect humans. In humans, T. cruzi exists either as infective bloodstream trypomastigote or as multiplying intracellular amastigote (mostly in the heart muscles, but also in lungs, liver, spleen, and other organs and tissues). In this work the antiparasitic effects of the CYP51 inhibitors were analyzed in T. cruzi amastigotes within cardiomyocytes. Because compound 4 did not inhibit CYP51 activity *in vitro*, it was not selected for cellular experiments. For the majority of the other compounds, a very good correlation between their potencies to inhibit T. cruzi CYP51 enzyme in the reconstituted reaction *in vitro* and antiparasitic effects against the pathogen cells has been observed (Table 1, Figure 7), although quite pronounced activity of 5, both the (R)- and (S)-enantiomers, might suggest that this structure can potentially have alternative target(s), better cellular permeability, or perhaps even an additional benzimidazole-like mode of action (oxidative stress) due to the presence of the reactive nitro group. Overall, strong antiparasitic activity combined with simple and cost-effective syntheses make these compounds, particularly T. cruzi CYP51 inhibitors (S)-2 (LFT) and (R)-6 (LFD) as well as compound (S)-5 (e.g., as an option for combination therapy) potentially promising new antichagasic drug candidates, which are worth testing in animal models.

**CONCLUSIONS**

By now T. cruzi sterol 14α-demethylase has been validated as essential for survival and multiplication of T. cruzi. The antifungal drugs posaconazole and ravuconazole may well pass clinical trials and reinforce the scarce arsenal of antichagasic chemotherapy. However, posaconazole is too expensive (>1,000 euro per patient), and ravuconazole still has unresolved problems with its bioavailability, which are most likely the major reason for its suppressive but not curative effect. We hope that VNI, or one of its derivatives, will also eventually proceed to translational research. Nevertheless, additional options for Chagas disease treatment are still highly needed, and therefore better understanding of the target CYP51 structure/function and inhibition should be greatly advantageous, particularly if the parasite acquires drug resistance upon treatment or if humans are infected with strains of T. cruzi that display natural drug resistance.

This work, in addition to identifying new potential drug candidates, also provides the structural rationale for the compounds’ inhibitory potencies and offers two alternative approaches that can be followed for further CYP51 structure-guided drug design and development. One approach is to use the high rigidity of the CYP51 substrate binding cavity. This feature is best displayed by the LFT/LFS costructures. Both these small molecules ensure their tight fit into the deepest segment of the CYP51 cavity via formation of the hydrogen bonding that not only strengthens the enzyme/inhibitor

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**Figure 4.** Time course of spectral responses of T. cruzi CYP51 (1.0 μM) to the addition of 0.2 μM of the heme-coordinating ligands.
interaction but also disrupts the heme support from the protein moiety, thus affecting the environment (and most likely redox potential) of the catalytic iron. The second approach is to block the entrance into the CYP51 substrate access channel by building a surface binding subsite (as, e.g., in the \textit{T. cruzi} complex with \textit{LFD} or posaconazole\textsuperscript{31}), since even weak van der Waals contacts between the enzyme and inhibitor in this area appear to prevent the substrate access channel from opening, the feature that must be functionally essential for sterol 14α-demethylase catalysis and therefore for the possibility of the substrate to replace the inhibitor in the enzyme active site.

**Table 2. Data Collection and Refinement Statistics**

| T. cruzi CYP51–inhibitor complex | (R)-6 (ligand PDB ID LFD)\textsuperscript{a} | (S)-2 (ligand PDB ID LFT)\textsuperscript{b} | (S)-3 (ligand PDB ID LFS)\textsuperscript{c} |
|----------------------------------|------------------------------------------|------------------------------------------|------------------------------------------|
| **Data Collection**              |                                          |                                          |                                          |
| Wavelength, Å                    | 0.9787                                   | 0.9787                                   | 0.9787                                   |
| Space group                      | P22(1)2(1)                               | P3(1)2(1)                                | P3(1)2(1)                                |
| Cell dimensions                  |                                          |                                          |                                          |
| \(a, b, c, \) Å                  | 59.900; 137.180; 152.430                 | 62.953; 62.953; 222.435                  | 62.582; 62.582; 221.068                  |
| \(\alpha, \beta, \gamma, \) deg | 90.00, 90.00, 90.00                       | 90.00, 90.00, 120.00                    | 90.00, 90.00, 120.00                    |
| No. of molec in asymm. unit      | 2                                        | 1                                        | 1                                        |
| Solvent content, %               | 58.2                                     | 48.8                                     | 47.9                                     |
| Resolution (last shell), Å       | 100–2.61 (2.67–2.61)                     | 30–2.6 (2.69–2.6)                        | 30–2.7 (2.8–2.7)                         |
| \(R_{merge}\) (last shell)      | 0.049 (0.558)                            | 0.052 (0.675)                            | 0.05 (0.583)                             |
| \(I/\sigma\) (last shell)        | 39.3 (2.9)                               | 31.0 (3.0)                               | 31.1 (3.4)                               |
| Completeness (last shell), %     | 99 (99)                                  | 99 (99)                                  | 99.5 (100)                               |
| Redundancy (last shell)          | 7.2 (5.6)                                | 7.2 (7.3)                                | 7.0 (7.2)                                |
| **Refinement**                   |                                          |                                          |                                          |
| Resolution, Å                    | 30.0–2.62                                | 30.0–2.74                                | 28.8–2.7                                |
| R-factor                         | 0.235                                    | 0.272                                    | 0.263                                    |
| R-free                           | 0.286                                    | 0.296                                    | 0.284                                    |
| Reflections used                 | 36336                                    | 13383                                    | 13738                                    |
| Test set size, %                 | 5.0                                      | 5.2                                      | 5.0                                      |
| rms deviations from ideal geometry |                                        |                                          |                                          |
| Bond lengths, Å                  | 0.005                                    | 0.001                                    | 0.003                                    |
| Bond angles, deg                 | 1.14                                     | 0.93                                     | 1.28                                     |
| Ramachandran plot                |                                          |                                          |                                          |
| Residues in favorable regions (%)| 95.3                                     | 95.6                                     | 94.2                                     |
| Residues in allowed regions (%)  | 99.8                                     | 100                                      | 100                                      |
| Outliers (%)                     | 0.2                                      | 0                                        | 0                                        |
| **Model**                        |                                          |                                          |                                          |
| No. of atoms (mean B-factor, Å)  | 7377 (69.7)                              | 3562 (90.5)                              | 3541 (90.9)                              |
| Number of residues per molecule  | A/B                                      | A                                        | A                                        |
| Protein                          | 449 (71.8)/450 (71.5)                    | 434 (88.1)                               | 434 (91.1)                               |
| Heme                             | 1 (30.2)/30.1                            | 1 (67.5)                                 | 1 (36.2)                                 |
| Ligand LFD                       | 1 (73.9)/1 (81.9)                        | 1 (75.6)                                 | 1 (72.7)                                 |
| Ligand LFT                       |                                            | 24 (55.6)                               | 4 (68.1)                                 |
| Ligand LFS                       |                                            |                                          |                                          |
| Water                            |                                            |                                          |                                          |

\textsuperscript{a}PDB code 4CK8. \textsuperscript{b}PDB code 4CK9. \textsuperscript{c}PDB code 4CKA.

**Figure 5.** \textit{T. cruzi} CYP51 active site illustrating interactions with LFD (cyan carbon atoms) and LFT (green carbon atoms). (A) Superimposition of CYP51 complexes with LFD and VNI (blue); (B) Superimposition of CYP51 complexes with LFT and VNF (red).
**EXPERIMENTAL SECTION**

**Chemical Synthesis.** All reagents and solvents were of high analytical grade and were purchased from Sigma-Aldrich (Milano, Italy). Melting points were determined on a Tottoli apparatus (Buchi) and are uncorrected. Infrared spectra were recorded on a Spectrum One ATR Perkin Elmer FT-IR spectrometer. \(^1H\) and \(^13C\) NMR spectra were acquired on a Bruker AVANCE-400 spectrometer at 9.4 T, in CDCl\(_3\), CD\(_3\)OD, or DMSO-\(d_6\), at 27 °C; chemical shift values are given in \(\delta\) (ppm) relative to TMS as internal reference. Coupling constants are given in Hz. Mass analysis was carried out with a 2000 Q TRAP instrument (Applied Biosystems), a commercial hybrid triple-quadrupole linear ion-trap mass spectrometer (Q1q2QLIT), equipped with an ESI source and a syringe pump; the flow rate was 5 \(\mu\)L min\(^{-1}\). The examined compounds were dissolved in methanol (10\(^{-5}\) M), and aqueous HCl was added just before the injection. The molecular peaks (m/z) have been observed as [M + H\(^+\)]. The stereochemistry of the \((R)-\) and \((S)-\)enantiomers as well as their enantiomeric excess (e.e.) was evaluated by chiral HPLC using a 250 mm \(\times\) 4.6 mm i.d. Chiralcel OD column (Chiral Technologies Europe, Illkirch, France).\(^\text{26}\) The HPLC apparatus consisted of a PerkinElmer (Norwalk, CT, USA) 200 LC pump equipped with a Rheodyne (Cotati, CA, USA) injector, a 20 mL sample loop, an HPLC Dionex CC-100 oven (Sunnyvale, CA, USA), and a Jasco (Jasco, Tokyo, Japan) model CD 2095 Plus UV/CD detector.

### Table 3. *T. cruzi* CYP51 Substrate Binding Cavity and Inhibitor-Contacting Residues

| secondary structural elements | amino acid residues | posaconazole (3K10\(^\text{23}\)) | \((R)\)-6 [LFD] (4CK8) | \((S)\)-2 [LFT] (4CK9) | \((S)\)-3 [LFS] (4CKA) | VNF (3KSW\(^\text{23}\)) |
|------------------------------|---------------------|-------------------------------|-----------------|-----------------|-----------------|----------------|
| A' helix (substrate access channel) | 145 | 145 | 145 | 145 | 145 | 145 |
| | V46 | V46 | V46 | V46 | V46 | V46 |
| | F48 | F48 | F48 | F48 | F48 | F48 |
| | G49 | G49 | G49 | G49 | G49 | G49 |
| β1-1/β1-2 (substrate access channel) | 170 | 170 | 170 | 170 | 170 | 170 |
| | 172 | 172 | 172 | 172 | 172 | 172 |
| B' helix (substrate binding cavity) | V102 | V102 | V102 | V102 | V102 | V102 |
| | Y103 | Y103 | Y103 | Y103 | Y103 | Y103 |
| | I105 | I105 | I105 | I105 | I105 | I105 |
| | M106 | M106 | M106 | M106 | M106 | M106 |
| | F110 | F110 | F110 | F110 | F110 | F110 |
| B'/C loop (substrate binding cavity) | V114 | V114 | V114 | V114 | V114 | V114 |
| | A115 | A115 | A115 | A115 | A115 | A115 |
| | Y116 | Y116 | Y116 | Y116 | Y116 | Y116 |
| C helix (substrate binding cavity) | Q126 | Q126 | Q126 | Q126 | Q126 | Q126 |
| | L127 | L127 | L127 | L127 | L127 | L127 |
| | L130 | L130 | L130 | L130 | L130 | L130 |
| F" helix (substrate access channel) | P210 | P210 | P210 | P210 | P210 | P210 |
| | A211 | A211 | A211 | A211 | A211 | A211 |
| | V213 | V213 | V213 | V213 | V213 | V213 |
| | F214 | F214 | F214 | F214 | F214 | F214 |
| I helix (substrate binding cavity) | M284 | M284 | M284 | M284 | M284 | M284 |
| | E285 | E285 | E285 | E285 | E285 | E285 |
| | V286 | V286 | V286 | V286 | V286 | V286 |
| | A287 | A287 | A287 | A287 | A287 | A287 |
| | F290 | F290 | F290 | F290 | F290 | F290 |
| | A291 | A291 | A291 | A291 | A291 | A291 |
| | H294 | H294 | H294 | H294 | H294 | H294 |
| β4-1 (substrate binding cavity) | T295 | T295 | T295 | T295 | T295 | T295 |
| | L356 | L356 | L356 | L356 | L356 | L356 |
| | L357 | L357 | L357 | L357 | L357 | L357 |
| | M358 | M358 | M358 | M358 | M358 | M358 |
| | V359 | V359 | V359 | V359 | V359 | V359 |
| | M360 | M360 | M360 | M360 | M360 | M360 |
| | R361 | R361 | R361 | R361 | R361 | R361 |
| β4 hairpin (substrate access channel) | Y457 | Y457 | Y457 | Y457 | Y457 | Y457 |
| | H458 | H458 | H458 | H458 | H458 | H458 |
| | T459 | T459 | T459 | T459 | T459 | T459 |
| | M460 | M460 | M460 | M460 | M460 | M460 |
| | V461 | V461 | V461 | V461 | V461 | V461 |
| | V462 | V462 | V462 | V462 | V462 | V462 |
| Total no. of residues | 25 | 19(20) | 16(17) | 16 | 14 | 14 |
Figure 6. Two regions of the *T. cruzi* CYP51 molecule that strengthen the enzyme interaction with LFD (A–C) and LFT/LFS (D–F). Coloring: (yellow, cyan) complexes with LFD, molecules A and B, respectively; (green) complex with LFT; (pink) complex with LFS. (A,D) Semitransparent surface representation (P450 distal view). (B) Structural elements forming the substrate access channel entrance. (C) F’”-helix. (E) Structural elements forming the deepest portion of the CYP51 binding cavity. (F) H-bonds between Y116 and the carbamic fragments of LFT and LFS.

Figure 7. Fluorescence microscopic observation of *T. cruzi* multiplication inside cardiomyocytes treated with 4 nM of CYP51 inhibitors. Drugs were added to infected cardiomyocyte monolayers at 24 h of infection, and amastigote multiplication was observed at 72 h of infection. GFP-expressing *T. cruzi* amastigotes are green, cardiomyocyte nuclei are blue, and cardiomyocyte actin myofibrils are red.
Declaration of Purity. The purity of all the compounds, determined by combustion elemental analysis and reverse-phase HPLC, was >95%. Elemental analysis was performed using a PE 2400 (PerkinElmer) analyzer, and the analytical results were within ±0.4% of the theoretical values. The reverse-phase HPLC system was equipped with a dual-wavelength UV 2454 detector (Waters) set at 250 and 205 nm and a Symmetry C18 (5 μm, 5 μL) column. The mobile phase was 55% 0.015 M potassium phosphate (pH 7.4) and 45% acetonitrile (v/v) with an isocratic flow rate of 1.0 mL/min.

**Compound 2.** The (R)- and (S)-enantiomers of 1-(4-chlorophenyl)-2-(1H-imidazol-1-yl)ethyl-4-isopropylphenylcarbamate were synthesized using the (R)- and (S)-enantiomers of 1-(4-chlorophenyl)-2-(1H-imidazol-1-yl)ethanol suspended in 5 mL of anhydrous CH2CN. The reaction mixture was stirred for 2 h at room temperature, and then 1.5 mmol of 4-isopropylphenyl isocyanate was added and stirred for 24 h at room temperature. The solvent was evaporated under reduced pressure, and the residue was washed with MeOH (3 × 3 mL). The mother liquor was dried and purified by silica gel column chromatography using CH2Cl2/MeOH (9:1) as eluent. The white solid was obtained with 45% yield (e.e > 99%). Mp = 162–164 °C; IR: 3435 cm−1; 1H NMR (DMF-d7): δ 9.75 (s, broad, 1H), 7.55 (s, 1H), 7.43 (d, 2H, J = 8.4 Hz), 7.36 (d, 2H, J = 8.4 Hz), 7.30 (d, 2H, J = 8.4 Hz), 7.15 (s, 1H), 7.12 (d, 2H, J = 6.4 Hz), 6.84 (s, 1H), 5.94 (m, 1H, 4.41 (m, 2H), 2.80 (m, 1H), 1.14 (d, 6H, J = 6.8 Hz). 13C NMR (DMF-d7): δ 152.2, 144.3, 138.5, 135.3, 135.2, 134.2, 129.1, 127.4, 126.8, 126.4, 120.2, 119.1, 73.6, 52.4, 33.5, 24.0; Anal. Calcd for C20H15ClN2O4: C, 56.25; H, 3.87; N, 11.32. Found: (S)-form: C, 56.48; H, 3.78; N, 10.93; (R)-form: C, 65.80; H, 5.75; N, 10.91. MS-ESI* 384.8 [M + H]+.

**Compound 3.** The (R)- and (S)-enantiomers of 1-(4-fluorophenyl)-2-(1H-imidazol-1-yl)ethyl-4-isopropylphenylcarbamate were prepared using the corresponding enantiomers of 1-(4-chlorophenyl)-2-(1H-imidazol-1-yl)ethanol following the procedure described for compound (2) and (2S). The white solid was obtained with 30% yield (e.e > 99%). Mp = 200–202 °C; IR: 1715 cm−1; 1H NMR (DMSO-d6): δ 9.74 (s, broad, 1H), 7.55 (s, 1H), 7.42 (m, 2H, J = 7.21) (d, 2H, J = 8.8 Hz), 7.22 (m, 2H), 7.15 (s, 1H), 7.13 (d, 2H, J = 8.5 Hz), 6.84 (s, 1H), 5.94 (t, 1H, J = 5.4 Hz), 4.43 (m, 2H), 2.80 (m, 1H), 1.16 (d, 6H, J = 6.9 Hz); 13C NMR (CDCl3): δ 162.7 (J = 245 Hz), 153.0, 143.8, 137.8, 136.0, 133.5 (J = 3 Hz), 128.0 (J = 9 Hz), 127.5, 126.3, 120.2, 118.8, 111.5 (J = 11 Hz), 74.2, 51.3, 33.4, 23.1; Anal. Calcd for C20H15FNO4: C, 56.65; H, 6.04; N, 11.44. Found: (S)-form: C, 58.39; H, 6.05; N, 11.42; (R)-form: C, 68.61; H, 6.15; N, 11.63. MS-ESI* 366.4 [M + H]+.

**Compound 4.** The (R)- and (S)-enantiomers of 1-(4-fluorophenyl)-2-(1H-imidazol-1-yl)ethanol were also used for synthesis of (R)- and (S)-1-(4-fluorophenyl)-2-(1H-imidazol-1-yl)-ethyl 3-(1H-imidazol-1-yl)propylcarbamate. 0.5 mmol of 1-(4-fluorophenyl)-2-(1H-imidazol-1-yl)ethanol was suspended in 5 mL of anhydrous CH2CN, and then 0.25 mmol of triphosgene was added and the solution was stirred overnight at room temperature. The reaction mixture was treated with Et3O, producing a white precipitate of 1-(4-fluorophenyl)-2-(1H-imidazol-1-yl)-ethyl chloroformate. The solvent was removed by decantation; the precipitate was washed with Et3O (2 × 5 mL) and dissolved in anhydrous CH2CN. Then, 1.6 mmol of TEA and 0.8 mmol of 3-(1H-imidazol-1-yl)-propylamine were added to the solution. The reaction was stirred overnight at room temperature. The crude mixture was diluted with H2O (5 mL), and the aqueous layer was extracted with CHCl3 (3 × 10 mL). The combined organic layers were dried under anhydrous sodium sulfate, and the solvent was evaporated under reduced pressure. The crude product was purified by silica gel column chromatography using CH2Cl2/MeOH (9:5.0:5) as eluent. The white solid was obtained with 70% yield (e.e > 98%). Mp = 220–225 °C; IR: 1720 cm−1; 1H NMR (CDCl3): δ 9.75 (s, broad, 1H), 7.69 (d, 2H, J = 2.0 Hz), 7.54 (s, 1H), 7.47–7.39 (m, 2H), 7.11 (s, 1H), 6.98 (dd, 1H, J = 8.9 Hz, J = 2.8 Hz), 6.93 (d, 2H, J = 8.4 Hz), 6.86 (s, 1H), 6.11 (m, 1H), 4.46 (m, 2H), 3.38 (m, 4H), 3.36 (m, 4H); 13C NMR (CDCl3): δ 150.5, 148.7, 138.8, 135.1, 133.0, 132.9, 130.5, 129.6, 129.2, 128.1, 127.9, 125.5, 122.5, 120.4, 120.3, 117.5, 117.2, 116.3, 115.5, 70.9, 50.6, 49.8, 48.9; Anal. Calcd for C24H26Cl2N5O2: C, 50.55; H, 4.16; N, 11.57. Found: (S)-form: C, 50.55; H, 4.16; N, 11.56; (R)-form: C, 50.68; H, 4.15; N, 11.69. MS-ESI* 605.98 [M + H]+.

**CYP51 Expression and Purification.** For functional studies, including ligand binding and enzymatic activity assays, we used the full-length Tulahuen T. cruzi CYP51 (GenBank IDs AV56083) expression construct, designed and purified as previously described in ref 21. For crystallization purposes, we utilized the T. cruzi CYP51 N-terminal truncated construct where the membrane anchor sequence.
cytrate dehydrogenase, and 25 mM sodium isocitrate in 20 mM
Hepes, pH 7.4, containing 150 mM NaCl, 0.1 mM EDTA. The optical path length was 5 cm.
Ligand binding was monitored as a red shift in the Soret band
absorbance spectra upon titration with each
ligand.

CYP51 Spectral Ligand Binding Assay. Changes in the T. cruzi CYP51 Soret band absorbance spectra upon titration with each
compound were recorded at 25 °C using a dual beam Shimadzu UV-
2401PC spectrophotometer. The P450 concentration was determined from
the Soret band intensity of the reduced CO complexes using
visible absorption difference spectroscopy, ΔSoret = 91 mM−1 cm−1.12 Spectral titrations were performed at ~1 μM P450 concentration in a 20 mM K-phosphate buffer, pH 7.4, containing 200 mM NaCl and 0.1 mM EDTA. The optical path length was 5 cm. Ligand binding was monitored as a red shift in the Soret band maximum reflecting coordination of the heterocyclic nitrogen to the
P450 heme iron.13 Difference spectra were generated by recording the
P450 absorbance in a sample cuvette versus the absorbance in a
reference cuvette, both containing the same amount of the protein.
Compounds were added to the sample cuvette in the concentration
range 0.2–20 μM from 0.4 mM stock solutions in DMSO. The
titration step was 0.2 μM. At each step, the corresponding volume of
DMSO was added to the reference cuvette. The apparent dissociation
constants (Kd) and the magnitude of spectral binding (binding efficiency (ΔΔSoret)) were calculated with GraphPad Prism (Graph-
Pad Software, La Jolla, CA) using a quadratic function for tight bind-
ing ligands by fitting the data for the ligand-induced spectral change (peak to trough absorbance changes in the difference spectra (ΔΔA)) versus total
ligand concentration to the following equation:

ΔA = (ΔΔmax/2E)(L + E + Kd) − (L + E + Kd) − 4LE2

where [L] and [E] are the concentrations of the ligand and the
enzyme used for the titration, respectively.28

CYP51 Activity Assay. The enzymatic activity of T. cruzi CYP51
was reconstituted in vitro as described previously using eburicol (24-
methylene-dihydrolanosterol) as the substrate.28 Briefly, the reaction
mixture contained 1 μM CYP51, 2 μM cytochrome P450 reductase
(CPR), 100 μM dilauroyl-α-phosphatidylcholine, 0.4 mg/mL iso-
zytate dehydrogenase, and 25 mM sodium isocitrate in 20 mM
Hepes (pH 7.4), 50 mM KCl, 5 mM MgCl2, and 10% glycerol. After
addition of the 3H-labeled sterol substrate (~2,000 cpm/nmol, final
concentration 50 μM), the mixture was preincubated for 5 min at
37 °C; the reaction was initiated by addition of 100 μM NADPH and
stopped by extraction of the sterols with ethyl acetate. The reaction
products were analyzed by a reverse-phase HPLC system (Waters)
equipped with a P15-RAM detector (INUS Systems, Inc.) using a Nova
Pak C18 column. The potencies of the compounds to inhibit T. cruzi
CYP51 activity were compared as inhibition of the substrate
conversion in a 1 h reaction at molar ratio enzyme/substrate/inhibitor
= 1:50:2, as these conditions were shown previously to allow
distinguishing the most potent compounds.17,20,25

T. cruzi Cellular Growth Inhibition Assay. A cellular T. cruzi
infection assay was performed using highly invasive 20A clone of the
Tulahuen strain of T. cruzi, which was shown to infect ~98% of
exposed cardiomyocytes.33 T. cruzi trypomastigotes expressing green
fluorescent protein (GFP) were generated as described.23 Trypomasta-
tigotes were used to infect cardiomyocyte monolayers in 96-well tissue
culture plates and in 8-well LabTech tissue culture chambers in
triplicate at the ratio of 10 parasites per cell as described.24,34 Cultures
were incubated with DMEM supplemented with 10% fetal bovine
serum (FBS) as described.34 Unbound trypomastigotes were removed by
washing the cellular monolayers with DMEM; and infected
monolayers were exposed to several concentrations of the inhibitors
(from 1 nM to 25 nM) dissolved in DMEM/DMEM free of phenol
red in triplicate at 24 h of infection, and cocultured in DMEM + 10% FBS for 48 h to observe parasite multiplication. Posaconazole was used as a
control. At 72 h of infection, the cardiomyocyte monolayers were
washed with phosphate-buffered saline, and the infection was
fluorometrically quantified as Relative Fluorescence Units (RFU)
using a Synergy HT fluorometer (Biotek Instruments).23,34 For
fluorescence microscopy observation, the infection assays were
performed in 8-well LabTech tissue culture chambers in triplicate;
after 72 h of infection the cardiomyocyte monolayers were fixed with
2.5% paraformaldehyde and stained with 4′,6-diamidino-2-phenyl-
indole, to visualize DNA, and with Alexa fluor 546 phallidin
(Invitrogen) to visualize cardiomyocyte actin myofibrils as described.24

X-ray Crystallography. The initial screening of crystallization
conditions was performed using Hampton Research crystallization kits.
Twenty mM stock solutions of compounds (S)-2, (S)-3, and (R)-6 in
DMSO were added at 2-fold molar excess to the 350 μM solution of T.
cruzi CYP51 in 20 mM K-phosphate buffer, pH 7.2, containing 200 mM NaCl, 0.1 mM EDTA, 10% glycerol, and 0.048 mM m-tridecylβ-
d-maltoside. Crystals were grown at 23 °C using the hanging-drop vapor diffusion method, cryoprotected by plunging them into a drop of
reservoir solution supplemented with 20% glycerol, flash-frozen in
liquid nitrogen, and then prescreened on a Bruker Microstar
microfocus rotating-anode X-ray generator/Proteum PT135 CCD
area detector. Crystals that diffracted to ~3.0 Å resolution were
subsequently used for the data collection at the synchrotron
(Advanced Photon Source, Argonne National Laboratory, IL) on
beamline 21ID-F at 100 K. The diffraction images were integrated
using Mosflm and scaled with Aimless (CCP4 Program Suite 6.3.35)
in the P2(1)2(1)2 space group in the case of LFD (to resolution of
2.61 Å) and in the P3(1)21 space group in the case of LFT (2.60 Å) and
LFS (2.70 Å). Solvent content was estimated with a Matthews
probability calculator. The crystal structures were determined by
molecular replacement in PhaserMR using the atomic coordinates of
the posaconazole-bound T. cruzi CYP51 structure (PDB code 3K1O)
as the search model. In each case Phaser found a single solution, with
one protein molecule in the asymmetric unit in the cocrystals of T.
cruzi CYP51 with (S)-2 and (S)-3 and with two protein molecules in
the asymmetric unit in the cocrystals of T. cruzi CYP51 with (R)-6.
The models were built and refined with COOT and REFMACS,
respectively, from the CCP4 Program Suite 6.3.35 Data collection
and refinement statistics are shown in Table 2. The electron densities
for each inhibitor were well-defined in all cases, showing full
occupancy and opposite orientation of short (S)-2/(S)-3 versus long
(R)-6 (the 2Fo − Fc electron density maps for (R)-6 and (S)-3 weighted at 1.5σ are shown in the graphical abstract).

# ASSOCIATED CONTENT

Supporting Information
Two figures showing (1) surface binding subsites formed in the
complexes of T. cruzi CYP51 with LFD and with posaconazole and (2) the influence of binding of LFD and LFT on T. cruzi
CYP51 flexibility. This material is available free of charge via the
Internet at http://pubs.acs.org.

Accession Codes
The coordinates and structure factors of T. cruzi CYP51 in
complex with compounds (R)-6 (ligand PDB ID LFD), (S)-2 (LFT), and (S)-3 (LFS) have been deposited in the Protein
Data Bank, PDB codes 4CK8, 4CK9, and 4CKA, respectively
(http://www.rcsb.org).

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Notes
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ABBREVIATIONS USED

CYP, cytochrome P450; CYP51, sterol 14α-demethylyase; T. cruzi, Trypanosoma cruzi; EC50, drug concentration that gives half-maximal response in cellular growth reduction; GFP, green fluorescent protein; RuCl2(p-cymene)((SS)-Ts-DPEN), (SS)-N-(p-toluenesulfonyl)-1,2-diphenylethanediamine(chloro)(p-cymene)ruthenium(II); VNI, ((R)-(N-(1,2,4-dichlorophenyl)-2-(1H-imidazol-1-yl)ethyl)-4-(5-phenyl-1,3,4-oxadiazol-2-yl)-benzamide); VNF, ((R)-(N-(2-(1H-imidazol-1-yl)-1-phenylethyl)-4′-chlorophenyl-4-carboxamide)
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