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HIGHLIGHTS

- J54, a safe and potent inhibitor of TLK1, was generated via in silico docking studies.
- J54 leads to apoptosis of PCa cells in combination with anti-androgen administration.
- J54 has low affinity for dopamine receptor and low anti-dopaminergic activity in animals.
- TLK1B is selectively increased upon ADT thereby providing a specific target for PCa.

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Generation of Phenothiazine with Potent Anti-TLK1 Activity for Prostate Cancer Therapy

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SUMMARY
Through in vitro kinase assays and docking studies, we report the synthesis and biological evaluation of a phenothiazine analog J54 with potent TLK1 inhibitory activity for prostate cancer (PCa) therapy. Most PCa deaths result from progressive failure in standard androgen deprivation therapy (ADT), leading to metastatic castration-resistant PCa. Treatments that can suppress the conversion to mCRPC have high potential to be rapidly implemented in the clinics. ADT results in increased expression of TLK1B, a key kinase upstream of NEK1 and ATR and mediating the DNA damage response that typically results in temporary cell-cycle arrest of androgen-responsive PCa cells, whereas its abrogation leads to apoptosis. We studied J54 as a potent inhibitor of this axis and as a mediator of apoptosis in vitro and in LNCaP xenografts, which has potential for clinical investigation in combination with ADT. J54 has low affinity for the dopamine receptor in modeling and competition studies and weak detrimental behavioral effects in mice and C. elegans.

INTRODUCTION
Prostate cancer (PCa) is a leading cause of morbidity and mortality of men in the western world. The standard of care for advanced PCa after failure of localized treatments is androgen deprivation therapy (ADT) and anti-androgens (Karantanos et al., 2013; Tao et al., 2019), which provides respite from disease progression, but ultimately fails resulting in the incurable phase of the disease: mCRPC. Treatments that can suppress the conversion to mCRPC have the best potential to improve outcome and be rapidly implemented in the clinics, and this requires a clear understanding of the process of PCa cells’ mechanisms of adaptation to ADT. This process has been well studied in the LNCaP cell line model, in which we have recently elucidated some missing details (Singh et al., 2019a). Androgen deprivation of LNCaP cells results in loss of AR function with a compensatory pro-survival activation of mammalian target of rapamycin (mTOR) (Carver et al., 2011) and concomitant implementation of cell division arrest by activation of the DNA damage response (DDR) mediated by ATR-Chk1 (Chiu et al., 2012) or ATM-Chk2 (Reddy et al., 2015). The DDR is likely activated due to the role played by the AR as replication licensing factor (Litvinov et al., 2006) in combination with the mTOR-dependent increased expression of TLK1B, which is highly dependent on the mTOR-E4EBP1 pathway (Sunavala-Dossabhoy et al., 2004), and resulting activation of the Nek1>ATR > Chk1 axis (Singh et al., 2019a). Additional work from our laboratory suggested that this may be a conserved nexus in additional cell models, in the TRAMP mouse, and probably in many patients, as the specific activating phosphorylation of Nek1 by TLK1 correlates with the Gleason score (Singh et al., 2019b). The resulting cell-cycle arrest is a survival mechanism for PCa cells, which remain quiescent until they reprogram and adapt to androgen-independent (AI) growth. An attractive strategy to prevent this process would be to bypass the cell-cycle arrest via inhibition of ATM or ATR, causing the cells to undertake replication with damaged DNA that would cause mitotic catastrophe, a strategy that was in fact implemented in LNCaP treated concomitantly with bicalutamide (BIC) and ATM inhibition (Reddy et al., 2015). However, a limitation of this approach is how to make the inhibition of ATM or ATR specific to PCa cells to limit general toxicity. We have recently demonstrated that addition of a relatively specific inhibitor of TLK, thioridazine (THD) (Ronald et al., 2013), which we repurposed for the blockage of the axis, results in fact in apoptosis of LNCaP and TRAMP-C2 cells concomitantly treated with BIC. In addition, it suppresses the late re-growth of PCa in the TRAMP mouse following castration (Singh et al., 2019a, 2019b). However, THD is a known anti-...
psychotic and has undesirable side effects. Here we describe J54, a potent and safe TLK1 inhibitor, as an adjuvant to ADT for PCa.

RESULTS

A Next-Generation PTH as a Potent Inhibitor of TLK1B

A few PTH anti-psychotics were identified in a compounds library screen as good inhibitors of TLK1 (Ronald et al., 2013). To find additional inhibitors, recombinant TLK1B was purified as previously described (Bhoir et al., 2018) (Figure 1A). The enzymatic activity was determined using the ADP-Hunter reagents and a Nek1 peptide (C).

Figure 1. Kinase Assays, Inhibitor Screening, IC50 Evaluation, and Competitive Assays

(A–C) Recombinant TLK1B purified to homogeneity (A) is active in kinase assays (B) and phosphorylates a specific Nek1 peptide (C).

(D) ATP dependence.

(E) Various compounds were tested for in vitro inhibitory effects, with J54 shown.

(F and G) Inhibitory curves of J54 and J56 compared to THD were carried out at 0.1 mM ATP (F) or at different competitive concentrations (G). The data presented are the mean ± SE of n = 3 independent experiments. The DiscoverRx ADP Hunter (Eurofins DiscoverRx, Fremont, CA, USA) Kit was used to measure the generation of ADP resulting from kinase phosphorylation of substrate. Further details are explained in Section C of Supplementary Material Methods.

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A few PTH anti-psychotics were identified in a compounds library screen as good inhibitors of TLK1 (Ronald et al., 2013). To find additional inhibitors, recombinant TLK1B was purified as previously described (Bhoir et al., 2018) (Figure 1A). The enzymatic activity was determined using the ADP-Hunter reagents and a Nek1
peptide containing the T141 target site. Properties of substrate (Nek1 peptide) and ATP dependence were determined (Figures 1B–1D), and in subsequent reactions with inhibitors used at 0.1 mM, which were found to be optimal. Several structurally similar compounds were synthesized as described in Supplemental Information (see NMR and mass spectrometry data Figures S3 and S4; and also Data S2) and tested at 20 μM (Figures 1E and 1F). J3-54 (thereafter J54) and J3-56, both being PTH derivatives, were as or more inhibitory than staurosporine, which is the standard pan-kinase inhibitor. We continued our characterization of J54. The other synthesized compounds did not show much inhibition, even though several are PTH (see Table S1). We have carried out some structure-activity relationship studies and molecular modeling to explain why, and for simplicity, we show below a comparison between THD and J54.

In Silico Modeling Studies of TLK1B with Docked J54 or THD
A model of the TLK1B kinase domain was constructed by homology modeling using the ROBETTA de novo protein structure prediction server (Song et al., 2013) (see Transparent Methods in Supplemental Information) and further compared/aligned to the published crystal structure of the highly homologous TLK2 (Moruzza et al., 2018). A PDB file for the structure was included as Data S1. The model was further refined by a 1-μs molecular dynamics (MD). The backbone root-mean square deviation (RMSD) of the structure stabilized after 100 ns, although, was subject to periodic RMSD shifts until 600 ns (Figure 2A). Compounds J54 and THD were docked into the ATP site of the final MD-refined model (Figures 2B and 2C). In the docked pose of compound J54, good interactions with the hinge region residues of TLK1 were exhibited; the morpholino head forms hydrogen bonds with Asp122, with a distance of 1.68 Å from the Asp122 carboxylate O to the ligand NH group (Figure 3A). A 100-ns MD simulation of this TLK1-J54 complex indicated that the pose is stable, further evident from a favorable total binding free energy $\Delta G_{\text{bind}} = -39.7$ kcal/mol, computed by the MM/GBSA method (Genheden and Ryde, 2015) (Figure 2D). THD was docked in the same TLK1 pocket but did not form hydrogen bonds with the protein (Figures 2B and S1). MD of the complex indicated a markedly higher RMSD in ligand atoms, with greater fluctuation, than for the J45 complex (Figure 2C); the free energy of binding is 9.9 kcal/mol weaker (Figure 2D). We note that this in part could arise from the methylthio group of THD, which prevents it from entering the hinge region to the same extent as J54 (Figure S1).

J54 in Combination with Anti-androgen (Bicalutamide) Induces Apoptosis in AS PCa Cells
We previously reported that THD has some growth inhibitory effects for several PCa cells lines, either androgen sensitive (AS) or insensitive (AI) (Singh et al., 2019a). We thus tested the same panel of cells in

| Complexes   | Dock Score | $\Delta G_{\text{bind}}$ (Kcal/mol) |
|-------------|------------|-----------------------------------|
| TLK1-J54    | -9.6       | -39.7 (4.1)                       |
| TLK1-THD    | -6.9       | -28.8 (3.7)                       |
| D2-J54      | -14.2      | -29.0 (5.4)                       |
| D2-THD      | -11.2      | -30.3 (5.4)                       |
| D2-Risperidone | -19.0     | -53.4 (3.6)                       |
androgen-containing medium (FBS) by proliferation assays. The effect was a weak dose-dependent inhibition with maximal efficacy around 18 μM (Figure 4B). Note, however, that RWPE1 (normal prostate) cell line was insensitive to J54, suggesting that J54 may be a more targeted anti-cancer agent, or it is possible that the TLK1>Nek1 axis remains critical even for CRPC cells. In contrast to these mild effects, combination treatment of the AS cell lines, LNCaP, VCaP, and TRAMP-C2, with BIC and J54 resulted in 4- to 5-fold suppression in colony formation (Figure 4A, p = 0.001). Note that VCaP cells were 60% inhibited by J54 alone, consistent with their elevated replication stress-driven DDR and checkpoint activation (Figure 5E), which when suppressed with J54 could lead to apoptosis. Clonogenic assays are unable to distinguish the effects of a DNA damaging agent, which can result in either an impaired or delayed resumption of growth (cell division) or loss of viability due to increased killing of the initial population. We thus measured in LNCaP and TRAMP-C2 cells the early change in cell number (MTT assay) over 72 h with J54 alone. The results indicated an actual loss in cell counts in relation to the dose (Figures 4C and 4D, p = 0.001). These results reinforced our main thesis that inhibiting the TLK1>Nek1 axis is mostly an effective regimen for AS cells in combination with anti-androgens. Indeed, cell cycle analysis of LNCaP and TRAMP-C2 cells treated with BIC, J54, or combination for 24 h showed a strong increase in the fraction of apoptotic cells only in the combination group (Figure 4E). Note also that, unlike TRAMP-C2 treated with BIC that display an accumulation of cells at G1/S and reduction of the S and G2 populations, cells treated with both BIC and J54 do not display the arrest (particularly no loss of the G2 cells), i.e., bypass of the G1 checkpoint and consequent apoptosis (Table S2). However, note that J54 treatment of LNCaP cells resulted in a modest reduction of the S-phase cells, so we cannot exclude that in some PCa cells J54 may cause a G1 arrest. To confirm these results and the specificity of the TLK1 inhibition with a genetic approach that could obviate potential off-target effects of J54, one possibility would be to deplete TLK1, but this is problematic because of the

Figure 3. Molecular Docking and Molecular Dynamics Studies of J54 and THD with TLK1
(A) Interactions of J54 with the active site of TLK1.
(B) Interactions of THD with the active site of TLK1.
apoiesis instead of leading to AI adaptation. We have not tested the activation of ATR/Chk1 and ATM/Chk2 in these cells to better assess these central mediators of the DDR and cell cycle checkpoints. This is therefore suggested evidence that TLKs are important for implementing the DDR and cell-cycle arrest after ADT and that bypassing this role results in accumulation of incompletely replicated DNA (due to loss of AR activity). In fact, it was reported that overexpression of TLK1-B > Nek1>ATR > Chk1 axis suppresses pNek1-T141 (Singh et al., 2017). We have generated LNCaP cells overexpressing TLK1-KD and found that after treatment with BIC alone, the cells fail to form AI colonies after 2 weeks and instead die. Interestingly, cell cycle analysis of these cells during a time course after addition of BIC showed an accumulation of cells in G2 before they begin to die, and not G1 as parental LNCaP (Figure 4E). In many cell types, including LNCaP, accumulation of cells in G2 is an indication of DNA damage post-replication, consistent with our interpretation of an important role of TLKs in mediating a G1/S checkpoint and preventing accumulation of incompletely replicated DNA (due to loss of AR activity). In fact, it was reported that TLKs stabilize replication forks (Lee et al., 2018), although these authors have not reported an effect for TLKs’ knockdown in suppressing ATR activation. However, they showed that ATM was not activated in cells depleted of TLK1-2, until they added UCN-01, a specific Chk1 inhibitor (Lee et al., 2018). This is consistent with our interpretation that bypassing the Chk1-mediated arrest in cells deprived of androgen and depleted of TLK activity results in replication fork collapse and activation of ATM. The apparent difference between suppression of TLK function by overexpressing TLK1-KD versus chemical inhibition with J54 can be explained by the more rapid induction of apoptosis for the latter, thus masking the G2 accumulation. When the TLK1-KD overexpressing cells were treated with BIC + J54 for 24 h (before apoptosis) the proportion of cells arrested in G2 was greater than the proportion of cells incubated with BIC alone. However, we have not tested the activation of ATR/Chk1 and ATM/Chk2 in these cells to better assess these central mediators of the DDR and cell cycle checkpoints. This is therefore suggested evidence that TLKs are important for implementing the DDR and cell-cycle arrest after ADT and that bypassing this role results in apoptosis instead of leading to AI adaptation.

**J54 with Bicalutamide Suppresses the Checkpoint Markers and Induces Apoptotic Markers**

LNCaP, VCaP, and TRAMP-C2 are the main examples of established PCa cell lines that are initially AS, but when maintained in ADT condition, recapitulate the conversion to AI growth observed in patients, and start growing again. In the initial phase of ADT, they arrest cell division, primarily at the G1/S transition, and become quiescent, which is a protective effect from undergoing replication fork collapse upon inhibition of the licensing function of the AR (Litvinov et al., 2006). As inhibition of the TLK1B > Nek1>ATR > Chk1 axis with THD results in bypass of the DDR and apoptosis (Singh et al., 2019a), we have tested the same process with J54. First, we confirmed that J54 (with or without BIC) causes a reduction in the p-Nek1-T141, the site of activation that is phosphorylated by TLK1, in all three AS cell lines (Figures 5 A–5H); another established TLK1 substrate, Rad9-S328, also was less phosphorylated after addition of J54 (Figure S7 D). We were also able to reproduce that the expression of TLK1B was induced in LNCaP and TRAMP-C2 following the addition of BIC, in contrast to the expression of the TLK1 splice variant that remains unchanged (Figure 5F). We previously showed that the increase in TLK1B is due to an mRNA translational effect caused by the compensatory increase in mTOR activity following ADT and suppression of AR signaling (Carver et al., 2011); note that there was no change in the TLK1B mRNA with BIC (Figure S7B). J54 also resulted in a futile increase inTLK1B, and not TLK1, possibly via a selective drug-induced stabilization of TLK1B, which turns over fairly rapidly (unpublished observations). Treatment with BIC resulted in activation of the DDR, shown as an increase in p-ATR and p-Chk1 in LNCaP and TRAMP-C2. In contrast, suppression of p-Nek1 and its
activity with J54 resulted in inhibition of the BIC-induced DDR activation (Singh et al., 2017, 2019a, 2019b), as manifested by a decrease in p-ATR and p-Chk1 (Figures 5A–5H). J54 alone caused a modest increase in p-ATR but not in p-Chk1 in LNCaP and TRAMP-C2, as we previously observed with THD (Singh et al., 2017), possibly due to a mild genotoxic effect. The VCaP cells were a little different because even without any BIC, p-ATR, and p-Chk1 were already elevated. This has been observed before (Karanika et al., 2017) and attributed to a constitutive DDR activation due to the TMPRSS2-ERG fusion in these cells (Chatterjee et al., 2015). Nonetheless, the addition of J54 to VCaP cells resulted in reduction of p-Nek1 and p-Chk1, indicating that

Figure 5. J54 in Combination with Bicalutamide Suppresses the Checkpoint Activation Markers and Induces Apoptotic Markers
Western blots (WB) of cell cycle and apoptotic indicators. (A–D) LNCaP, (E) VCaP, and (F–G) TRAMP-C2 cells from the four treatment groups as indicated were analyzed by WB for several indicators of DNA damage/apoptosis and mediators of cell-cycle arrest. Representative analysis of two independent experiments for each cell line is shown.

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it can suppress the DDR checkpoint, whether activated constitutively or after BIC treatment (Figure 5E). As we previously reported for inhibition of TLK1 with THD (Singh et al., 2019a), bypass of the BIC-induced checkpoint can result in generation of DSBs (indicated by the presence of γH2AX) and leads to apoptosis (indicated by increased cleaved caspase 3 and PARP, Figures 5A–5H). Likewise, in BIC + J54-treated cells there was an increase in P21 expression (Figures 5C and S5G), an indicator of the emergency activation of the P53 > p21 pathway, and an effect previously observed for BIC + THD-treated LNCaP and TRAMP-C2 cells that resulted in pATM-S1981 activation from generation of DSBs (Singh et al., 2019a) and reproduced for J54 + BIC (Figure 5J). The induction of p21 has been likewise reported in TRAMP-C2 cells treated with imiquimod and correlated with accumulation of cells in G2 before the onset of apoptosis (Han et al., 2013).

**Bicalutamide with J54 Suppresses Growth of LNCaP Xenografts via Suppression of the TLK1B > pNek1 DDR Pathway and Promotes Apoptosis**

We sought to establish if the addition of J54 could suppress the resurgence of tumor growth of LNCaP xenografts and their later conversion to CRPC. Following formation of sizable tumors (~200 mm³), castration, or anti-androgens arrest the progression of LNCaP xenografts for some time (2–3 weeks), whereas subsequently the tumors start growing again at an accelerated rate, which is refractory to ADT (AI). Therefore, we injected LNCaP cells in Matrigel in both flanks of NOD-SCID mice and then randomly assigned them to four treatment groups (n = 5 per group × 2 independent experiments), as shown in Figures 6A–6C. The control group showed progressive exponential growth, and so did the BIC group after a 12-day lag following the beginning of BIC administration. Interestingly, treatment with J54 alone showed significant suppression of tumor growth (p = 0.01) and tumor weight (p = 0.001), whereas the combination (BIC + J54) resulted in complete suppression of tumor growth and weight (p = 0.001) and actual regression of the tumors compared with the starting size Figures 6A–6C. An immunohistochemical analysis of the available excised tumors showed that the phosphorylation of Nek1-T141 was increased in BIC-treated group (Figure 6D, p = 0.001), consistent with a corresponding increase in TLK1B expression (Singh et al., 2019a, 2019b); it was suppressed by concomitant administration of J54 (p = 0.001), which is expected to result in bypass of the DDR checkpoint and increased apoptosis and corresponding markers. In fact, the combination treatment showed a strong increase in staining for cleaved PARP and caspase 3, and γH2AX (a marker of DNA damage, p = 0.001, Figure S8). Tumors in the combination also showed a reduction in the number of proliferative cells by Ki67 staining (p = 0.001), although this was lower also for the groups singly treated with BIC and J54 (Figure S8, note that these tumors were isolated at day 30 when they were still sizable). We should also point out that we treated the mice with J54 only biweekly due to the favorable pharmacokinetics, where maximal plasma concentration of 100 ng/mL (35 µM) was reached 2 h after intraperitoneal injection, and was still present at ~6 ng/mL (6 µM) after 24 h (Figures S5A–S5C).

**J54 Has Low DR2 Binding Activity and Behavioral Effects in Animal Models**

THD has strong anti-tumor effects in combination with ADT for AS-PCa models (Singh et al., 2019a, 2019b), but after ~30 years of use, this drug was withdrawn for treatment of schizophrenia due to increased risk of cardiac arrhythmia (Wu et al., 2015) and extrapyramidal toxicity. However, we reasoned that other PTH inhibitors of TLK1 could have similar benefits for the treatment of PCa without the side effects. The repurposing of PTH anti-psychotics for cancer therapy has been proposed (Lee et al., 2016), even though their cellular targets have not been identified, as they were generally assumed to work largely through inhibition of dopamine receptors (Yin et al., 2015). Interestingly, an analysis from five independent studies of PCa incidence in individuals with schizophrenia revealed a significant decrease in standardized incidence ratio ranging from 0.49 to 0.76 (95% confidence interval) (Torrey, 2006) (Mortensen, 1989). This reasoning has led to the design and synthesis of J54. However, these observations beg the question of whether J54 has also anti-dopaminergic activity. We thus considered the potential interaction with the D2 dopamine receptor (DR2). Compounds J54, THD, and DR2 antagonist risperidone were studied by molecular docking, MD, and free energy calculations, based on the risperidone/DR2 crystal structure (PDB code 6CM4) (Wang et al., 2018). The three ligands were docked in the DR2 pocket (docking was able to reproduce the observed pose of risperidone). The three complexes were solvated and simulated via MD for 100 ns. Compound J54 binds stably in the active site of the DR2, forming a hydrogen bond with Ser159 distance of 2.24 Å (Figure S2A, related to Figure 2D). THD and risperidone also complex with the DR2 via hydrogen bonds (Figures S2A and S2C, related to Figure 2D). Interestingly, the binding free energy computed via MM/GBSA ranks J54 as the lowest affinity ligand, with a ΔGbind value of −29.0 (Figure 2D). This is followed by THD (~39.0 kcal/mol) and then risperidone (~53.4 kcal/mol). These free energy calculations suggest that J54 binds only weakly and has a lower affinity than risperidone and THD toward DR2. To verify
experimentally that J54 is a weak DR2 antagonist, we commissioned from Sekisui-Xenotech a competitive radioassay using 7-hydroxy DPAT, R-(+)-[3H] as a tracer, and two recombinant human dopamine receptors (D1 and D3). Positive control antagonists were included [R(+)-SCH-23390 and ((+)-7-Hydroxy-2-(di-n-propylamino) tetralin ([(+)-7-OH-DPAT]). In Table S3, related to Figure 2D, we show the results obtained at effective concentrations of 100 nM for J54 compared with THD, which confirmed that J54 is a very weak antagonist for dopamine receptors. Note that in these competitive studies with recombinant receptors, the antagonist (e.g., THD) is typically active in the 10–100 nM range (IC50, 20 nM for THD), whereas for DR2 antagonism for amelioration of psychotic conditions, circulating concentrations in plasma need to be ~100 μM.

We then did a general assessment of mice following injections of J54. We noticed no toxicity (even after gross inspection of organs at necropsy), no decrease in body weight, and no behavioral changes (no lethargy or extrapyramidal twitches sometime observed with THD). To get a better assessment of the possible behavioral effects of J54, we used C. elegans that has a simple but complete nervous system and has been well characterized for its responses to anti-psychotic drugs, including actions at serotonin or dopamine

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**Figure 6. Combination Bicalutamide and J54 Suppresses Growth of LNCaP Xenografts**

(A) Time course of tumor growth of LNCaP cells xenografts in 4 treatment groups. Treatment started 19 days after implantation when the tumors measure ~200 mm³. Two independent experiments with 5 mice per group were carried out. J54 and bicalutamide were dissolved in DMSO and diluted in corn oil 1:10 and administered intraperitoneally biweekly. Sectioning and processing of the tissues were carried out in the FWCC Histology Service, using automated processes and equipment to provide uniform and standardized results. Indirect labeling was with ABC Elite: RTU Vectastain Elite Reagent, Vector #PK-7100; DAB: ImmPact DAB, Vector #SK-4105. Light counterstaining was done with hematoxylin.

(B) Tumor weights were determined for all groups at end of the treatment course.

(C) Examples of tumor size at the end of the experiment.

(D) Representative sections from tumors resected from mice in the 4 treatment groups analyzed by immunohistochemistry for pNek1 (n = 3 mice; scale bar, 50 μm). Note the weak pNek1 stain in the tumor from mice treated with J54, in contrast to the increase seen with BIC.

Two-way ANOVA tests were done to compare the group for statistical significance; *p < 0.05, **p < 0.01, ***p < 0.001.
receptors (Donohoe et al., 2006, 2009). Consistent with the weak binding of J54 to DR2 (Figure 2D), in three studies, J54 had much weaker behavioral effects then TFP or risperidone, attributable to DR2 activity (Figures S6A–S6C). Note that the concentration of these drugs on agar plates need to be significantly higher than in tissue culture medium for mammalian cells because of the protective and poorly permeable cuticle in C. elegans.

DISCUSSION

Certain PTH are potent inhibitors of TLK1 (Ronald et al., 2013), and by inference TLK2, as these proteins must homo- and hetero-dimerize for full activation (Mortuza et al., 2018), in addition to being highly similar in sequence and structure, and thus similarly targetable. In fact, a Kinomescan carried out at 15 μM with J54 resulted in inhibition of TLK1 and TLK2 and little inhibition of all other kinases with the exception of TTK ("SCANMAX" file, Table S4, related to inhibition data in Figure 1). This was similar to what we previously found for THD and PPH (Ronald et al., 2013) providing additional evidence for the specificity of PTH compounds with respect to other kinases except perhaps for reported inhibition of PKC (Aftab et al., 1991) that was not tested here as it is not included in the SCANMAX list, which also does not include key PIKK (see Limitations). Two noteworthy observations are that PTHs are structurally different from many known ATP competitive inhibitors, and that J54 was carefully designed based on considerations reported previously for the THD—the Kinomescan has confirmed that J54 is specific to TLKs. We should point out that although PTH were the first reported inhibitors of TLKs (Ronald et al., 2013), other inhibitors particularly of TLK2 have since been reported (Kim et al., 2016), some based on structure docking (Mortuza et al., 2018), although neither their potency and effects in vivo nor their potential off-target effects have not been reported as they were selected from libraries of kinase inhibitors.

As not all PTH are good inhibitors of TLK1 (Ronald et al., 2013 and this study), it was critical to first sort out in silico which compounds best fit in modeled binding site. We have now shown that J54 has higher affinity than THD by MD analysis, and hence is more potent and likely more specific for TLK1. We have previously reported that a TLK1 inhibitor, THD, could synergize with ADT in promoting apoptosis on AS PCa cell lines in culture and in xenografts (Singh et al., 2019a), and in the TRAMP mouse model (Singh et al., 2019b). This rationale was derived from our previous discovery that TLK1 is an upstream activator of the Nek1>ATR > Chk1 axis [11], in conjunction with an original result that showed that TLK1B is translationally increased following AR suppression (Singh et al., 2019a) and the established compensatory mTOR activation (Carver et al., 2011). Abrogation of the TLK1>Nek1>ATR axis was expected to result in bypass of the DDR checkpoint and thus promote mitotic catastrophe. This was also consistent with the known role of AR signaling in regulation of DNA repair in PCa and synergistic killing with inhibitors of DNA repair (Karanika et al., 2017; Thompson et al., 2017).

It seems clear that certain PTH anti-psychotics correlate with a decreased risk of PCa development (Torrey, 2006), whereas the use of THD and some other PTH anti-psychotics presents some risks and side effects. Therefore, we have developed a second-generation TLK inhibitor that has lower affinity for the DR2. The potent dopamine antagonist activity of some PTH anti-psychotics has been blamed for some of the lethargic and extrapyramidal effects, as well as for the cardiac arrhythmia. In silico, J54 showed much weaker binding to the DR2 than THD or risperidone, and it appeared to have little adverse behavioral effects in mice and worms.

In this work, several PCa cell lines were studied for growth inhibition by J54, and particularly the AS cell lines, LNCaP, VCaP, and TRAMP-C2, were sensitive to apoptosis when combined with an anti-androgen (BIC). The LNCaP model was also tested in xenografts and demonstrated remarkable tumor regression. In conclusion, we suggest the use of J54 as adjuvant therapy for PCa in conjunction with anti-androgens, as a safer and more potent inhibitor of this DDR axis, which we believe is commonly activated during the initial phase of PCa cells’ adaptation to ADT. In conclusion, the regulation of the DDR by TLK1 through the Nek1>ATR > Chk1 axis, and even more importantly its upregulation after ADT, is an important finding in the field of PCa research and as a target for potential therapy (Singh et al., 2017, 2019a, 2019b). A very large amount of work in PCa therapy has been devoted to the search for better anti-androgens, whereas relatively little has been devoted to combining ADT with targeting the known role of the AR in controlling the DDR (Karanika et al., 2015; Thompson et al., 2017). Our approach, which may seem to go counter-current to the established views of standard of care for advanced PCa, is to abrogate the ADT-induced DDR checkpoint and cell-cycle arrest, thereby forcing apoptosis of PCa cells still responsive to ADT. Although it
seems clear that certain PTH anti-psychotics correlate with a decreased risk of PCa development (Torrey, 2006), the use of THD and some other PTH anti-psychotics presents some risks and side effects. In particular, an increased risk for cardiac arrhythmia (Hennessy et al., 2002) as a result of their anti-dopaminergic activity and inhibition of hERG channels could impede repositioning of some of these PTH for the treatment of PCa due to potential concerns by regulatory agencies. J54 was designed and tested to be a weaker inhibitor of DR2, and we in fact noticed it had no apparent toxicity in mice, with no extrapyramidal twitches or altered breathing after administration, and relatively weak anti-dopaminergic effects in C. elegans. Therefore, it is a bona fide, specific inhibitor of TLK1, and not the DR2, with superior efficacy and better side effects profile.

Limitations of the Study
Although we tried to ensure mechanistically that J54 works via inhibition of the ADT-activated TLK > Nek1>ATR > Chk1 DDR pathway, we cannot rule out that the tumor regression effects that we have observed could be caused by inhibition of some other target. And in fact, while this paper was under review, a similar PTH with putative lack of DR2 antagonism was described as having antitumor activity via activation of PP2A (Monta et al., 2020). In addition, key large PIKKs, like ATR, ATM, and mTOR are not included in the SCAN-MAX panel, and therefore could be potentially inhibited by J54. We have shown that ATM can become activated in cells treated with BIC + THD or J54 (Figure S7C), whereas in particular, mTOR has been reported to be inhibited (directly or indirectly) by THD (Kang et al., 2012; Park et al., 2014). Moreover, the combined activation of PP2A and inhibition of mTOR can synergistically suppress growth of pancreatic adenocarcinoma (Allen-Petersen et al., 2019). Another kinase that was inhibited by J54, TTK, may play important roles in mitotic progression and was reported to be critical in pancreatic cancer (Kaistha et al., 2014). Another potential non-kinase target is actually the AR itself (Bisson et al., 2007), which can complicate the interpretation of our experiments, although the fact that mice treated concomitantly with J54 and BIC (which is a much more powerful anti-androgen) show strong tumor regression argues against the idea that the two drugs act on the same pathway.

There are also potential negative consequences in principle to inhibiting TLKs in normal cells and tissues. We have previously reported that in a "normal" non-androgen-dependent cell line (MM3MG) overexpression of TLK1-KD, the cells are viable but a fraction shows defects in mitotic segregation and high levels of aneuploidy (Sunavala-Dossabhoy et al., 2003).

Resource Availability
Lead Contact
Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact: Arrigo De Benedetti.

Materials Availability
Plasmids that were used for this study and derived cell lines are available from the Lead Contact after fulfilling a MTA.

Data and Code Availability
All Data and Code generated or analyzed during this study and its Supplemental Information files are available upon request.

METHODS
All methods can be found in the accompanying Transparent Methods supplemental file.

SUPPLEMENTAL INFORMATION
Supplemental Information can be found online at https://doi.org/10.1016/j.isci.2020.101474.

ACKNOWLEDGMENTS
This work was supported by DoD-PCRP grant W81XWH-17-1-0417 to A.D.B. S.B. was supported by Fulbright-Nehru Doctoral Research Scholarship E0588101. S.K. and J.H. thank DRDO and SERB for funding. R.V.C. thanks the Royal Society and the Bhabha Fund for a Newton International Fellowship. R.A.B. and
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Supplemental Information

Generation of Phenothiazine

with Potent Anti-TLK1 Activity

for Prostate Cancer Therapy

Vibha Singh, Siddhant Bhoir, Rupesh V. Chikhale, Javeena Hussain, Donard Dwyer, Richard A. Bryce, Sivapriya Kirubakaran, and Arrigo De Benedetti
**TRANSPARENT METHODS:**

**Molecular Modelling, Docking, Molecular Dynamics Simulations and Free Energy:**
Details about methodology used for molecular modelling, docking, molecular dynamics simulations and free energy calculations were explained in details in section A of supplementary material and methods.

**Chemical synthesis of J54 and several different Phenothiazines:**
Details about chemical synthesis of J54 and several other Phenothiazines, and methodology were explained in section B of supplementary material and methods.

**Protein expression and purification:**
The full-length, 65KDa recombinant human TLK1B (hTLK1B) protein for the *in-vitro* experiments was expressed and purified from bacteria using the protocol referenced in main document.

**Inhibitor screening, IC\textsubscript{50} evaluation and mode of binding studies:**
The DiscoverRx ADP Hunter™ (Eurofins DiscoverRx, Fremont, CA, US) Kit was used to measure the generation of ADP resulting from kinase phosphorylation of substrate. Details about the inhibitor screening, IC\textsubscript{50} evaluation and mode of binding studies were explained in section C of supplementary material and methods.

**Cell lines:**
All prostate cancer cell lines were obtained from the ATCC, recently authenticated, and free from mycoplasma contamination, and cultured as per instructions.

**Antibodies:**
Antibodies used for present study is listed as supplementary table: Key Resources.

**Clonogenic Assay:**
One thousand cells were seeded, treated and cultured for over 2 weeks. Colonies were stained with crystal-violet and manually counted by using NIH ImageJ software

**Cell proliferation and cell viability Assay:**
Ten thousand cells were seeded, and next day treated for desired concentration of J54 and different time-point. Cell proliferation assay was done by crystal-violet staining and viability assay were done by MTT dye.

**Cell Cycle Analysis:**
We performed cell cycle analysis as described earlier (Singh et al; 2017) with LNCaP and TRAMP-C2 cells with four group of treatment i.e. Control, Bicalutamide, J54 and combination of J54 and Bicalutamide.

**Western blot analysis:**
Western blot analysis of different proteins listed in supplementary table 1 were done with Control, Bicalutamide, J54 and combination of J54 and Bicalutamide treated samples as described earlier [1] for LNCaP, TRAMP-C2, and VCaP cells.
LNCaP Xenograft Model:

All animals used in this study received humane care based on guidelines set by the American Veterinary, and approved by the Institutional Animal Care and Use Committee (IACUC) of LSU Health Sciences Center at Shreveport and also following ARRIVE (Animal Research: Reporting of In Vivo Experiments) guidelines. Male NOD/SCID mice (5 per group) were purchased from Jackson labs. 10^6 LNCaP cells were injected into the flanks of mice in 0.2 of matrigel. Mice were injected IP bi-weekly with J54: 5 mg/kg body weight (dissolved in DMSO and diluted 1:10 in corn oil). Dose estimation for J54 was derived from pharmacokinetics study. Bicalutamide was dissolved in DMSO and diluted in corn oil 1:10 and administered IP at a dose of 100 mg/kg body weight bi-weekly.

Behavioral assays in *C. elegans* to assess the behavioral effects of J54 as a potential DR2 antagonist (anti-psychotic) were carried out as described in details in Donahoe et al. 2009.

Pharmacokinetic Studies on J54: These were carried out at the preclinical testing lab of UTSW in Dallas, TX. 21 male CD-1 mice were dosed IP with 10 mg/kg J54. 0.2 ml/mouse formulated with 10 % DMSO, 90 % Corn Oil. Whole blood was collected in a syringe coated with ACD. Plasma was processed from whole blood by centrifugation at 10,000 rpm for 10 minutes.

For the standards and QC'S 98 ul & 98.8 ul of blank plasma was added to an eppendorf and spiked with 2 & 1.2 ul of initial standard. Std's, QC's & samples of 100 ul were then crashed with 200ul of methanol containing 0.15% formic acid and 12.5 ng/ml IS (final conc.). The samples were vortexed 15 sec, incubated at room temp for 10' and spun 2x 13,200 rpm in a standard microcentrifuge. The supernatant was then analyzed by LC-MS/MS. Vendor supplied plasma used in standards and QCs: Bioreclamation, LLC; lot #MSE284936; exp = 03/31/2020; ACD anticoagulant. Raw data and additional information can be obtained upon request.

Immunohistochemistry (IHC) on LNCaP Xenograft tumors:

Sectioning and processing of the tissues were carried out in the FWCC Histology Service, using automated processes and equipment to provide uniform and standardized results. Indirect labeling was with ABC Elite: RTU Vectastain Elite Reagent, Vector #PK-7100; DAB: ImmPact DAB, Vector #SK-4105. Light counterstaining was done with hematoxylin.

Statistics:

Significance between treatment groups (mean values and SDs) was performed by a one-way ANOVA followed by Tukey's multiple comparisons tests using GraphPad Prism6 software (GraphPad Software USA). p-value <0.05 (*p < 0.05; **p < 0.01; ***p < 0.001) was considered to indicating a statistically significant difference.
Supplementary Material Methods and Tables

Additional information was included in the SI for the chemical synthesis and physical authentication of the compounds synthesized.

A. Molecular Modelling, Docking, Molecular Dynamics Simulations and Free Energy Calculations:

I. TLK1B

1. TLK1B- Modelling

In the absence of a crystal structure of the TLK1B kinase domain, we constructed a 3D model of its 333 amino acid residues using the ROBETTA protein structure prediction software (Raman et al., 2009) using the hierarchical Ginzu screening method (Song et al., 2013). The modelled TLK1B structure for its stereochemical quality using the MOE software package: the majority of residues were found in the favored and allowed region, with only three residues (Val221, Glu119, Asp233) as outliers (Ramachandran et al., 1963). Good validation scores were obtained for the model using the MolProbity server (Chen et al., 2010). The MolProbity score was 1.52, with a clash score for all atoms to be 0.85 and one outlier rotamer. Further, Qualitative Model Energy analysis (QMEAN) (Benkert et al., 2011) using the QMEAN web server was carried out for the TLK1B model, which gave a QMEAN value of -1.26 which is a good score for modelled proteins, the TLK2 crystal structure (PDB 5O0I) has QMEAN value of -0.85.

1.2. Molecular dynamics simulations of TLK1B protein

MD simulations were performed using the AMBER16 software package (Case et al., 2016). The TLK1B model was immersed in a truncated octahedron of TIP3P water (Mark et al., 2001) giving a total of 17257 water molecules. Na⁺ and Cl⁻ counter ions were added to neutralize the system and provide an ionic strength of 0.1 M. The ff14SB force field (Maier et al., 2015) was used to model the protein.

Simulations were run at 300 K using the Langevin thermostat (Bussi et al., 2008) with a collision frequency of 2 ps⁻¹; and 1 atm using a Monte Carlo barostat (Jorgensen et al., 1996) with volume exchange attempts every 100 fs. A 2 fs integration step was employed. Covalent bonds involving hydrogen were constrained using SHAKE (Debolt et al., 1993). A cutoff of 8 Å was used for short range nonbonded interactions whilst long range electrostatics were treated...
using the particle mesh Ewald method (Essmann et al., 1995). Equilibration consisted of rounds of NVT and NPT equilibration for 10 ns in total. Production MD run was performed for 1 μs. Interactions were analysed using the cpptraj (Roe et al., 2013) module, taking configurations every 4 ps. For MD simulations of docked ligand complexes of TLK1B and D2 receptor (see Methods below), each system was immersed in a truncated octahedron of TIP3P water and simulated for 100 ns. Each trajectory was subjected to MM/GBSA analysis (Case et al., 2014; Genheden et al., 2015) for 10,000 equispaced frames.

1.3. Molecular Docking

Molecular docking studies were performed using FRED 3.3.0.3 (McGann et al., 2011 - OpenEye Scientific Software) with the Chemgauss4 score and energy functions (McGann et al., 2003). The TLK1B model was investigated for active sites using SiteFinder tool in MOE software (Paul and Santavy, 2007) The site with highest the PLB score was selected for docking exercise, which was the ATP binding site in the kinase hinge region. The OMEGA module (Hawkins et al., 2010) of OpenEye was used to generate all the possible conformers of the ligands for docking studies. Docking of J54 and THD to the ATP binding site of TLK1B was performed. For docking to the D2 receptor, the crystal structure of D2 receptor bound to risperidone was obtained from the Protein Data Bank (PDB 6CM4) (Wang et al., 2018). The crystal structure was analysed for any inconsistencies like breaks in the sequence and protonation states using the MOE software. The protein was further prepared by energy minimisation and addition of hydrogens to reduce the strain energy from crystal formation. Conformers of ligands J54, THD and risperidone were docked using FRED.

B. Chemical synthesis of J54 and several different Phenothiazine:

Chemical reagents and Instrumentation:
All the solvents were of HPLC grade and procured from Sigma, Finar and SD Fine. And all other reagents were of analytical grade and were used without further purification. The reaction was performed in a round-bottomed flask with magnetic stirring bar under N₂ atmosphere. Completion of the reaction was monitored by the Thin-layer chromatography carried on Merck TLC silica gel 60 F254 and visualized by ultraviolet irradiation. The purification of all the synthesized compounds was done by the column chromatography using silica gel 100-200 mesh supplied by Merck. The synthesized compounds were characterized by NMR spectra,
recorded by Bruker AVANCE III 500 (1H NMR: 500 MHz, 13C NMR: 125 MHz). The splitting patterns are referred as follows: s, singlet; d, doublet; t, triplet; m, multiplet; dd, doublet of doublets; and td, triplet of doublets. NMR spectra were comparable with those previously published (J3-56). Melting points (mp) were recorded in an open glass capillary using LABINDIA Visual Melting Range Apparatus (MR-Vis) instrument. The mass spectrum and the infrared spectra (wave numbers, cm⁻¹) were recorded by Waters Synapt-G2S ESI-Q-TOF (Positive mode) and Perkin Elmer FTIR Spectrometer (Spectrum 2) respectively.

**S1 : Synthesis of 4-(2-(10H-phenothiazin-10-yl) ethyl) morpholine, J-54**

The compound J-54 was initially tried to synthesized in 1949 (Dahlabom et al., 1949) but it was not isolated or purified in a pure state. We are the first one to report J3-54, crystalline solid, which is in pure state.

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\begin{align*}
\text{S} & \quad \text{N} \quad \text{O} \\
\text{H} & \quad \text{Cl} \\
\end{align*}
\]

**Synthetic procedure**

To a solution of phenothiazine (200 mg, 1.0 mmol) in dry DMF in a round-bottomed flask, sodium hydride (96.4 mg, 4 mmol) was added slowly under inert condition at 0 – 5 °C. The reaction mixture was stirred at room temperature for 15 – 20 min followed by the addition of 4-(2-chloroethyl) morpholine hydrochloride (186 mg, 1 mmol). The above reaction mixture was stirred at room temperature for 10 – 12 h. The reaction towards completion was monitored by TLC. Then the solution was diluted with ethyl acetate and washed with brine solution. The organic layer was dried over Na₂SO₄, evaporated on a rotary evaporator under high pressure. The crude product was purified by silica gel column chromatography using 4 % ethyl acetate/Hexane, yielding the desired product as a yellowish orange solid. Yield (185 mg, ~ 59 %); mp: 78 – 80 °C; IR (v, cm⁻¹): 1117(C–O), 1035 (C–N); 1H NMR (500 MHz, CDCl₃): δ 7.11 (m, 4H), 6.9 (m, 4H), 4.0 (t, 2H, J = 7 Hz), 3.71 (t, 4H, J = 4.5 Hz), 2.72 (t, 2H, J = 7 Hz), 2.51 (t, 4H, J = 4.5 Hz); 13C NMR (125 MHz, CDCl₃): δ 145.0, 127.5, 127.2, 124.9, 122.6, 115.4, 66.9, 56.0, 54.0, 45.7; HRMS (m/z): [M+H]⁺ calculated for C₁₈H₂₁N₂O₅, 313.1330; found, 313.1336.
S2: Synthesis of 10-methyl-10H-phenothiazine, J-56

The procedure was analogous to that used for the synthesis of J-54 except that methyl iodide was used instead of 4-(2-chloroethyl) morpholine hydrochloride. The crude product was purified by silica gel column chromatography using 0.1% ethyl acetate/Hexane, yielding the desired product as a white solid. Yield (189 mg, ~88%); mp: 100 – 102 °C; IR (ν, cm⁻¹): 1330 (C=N); ¹H NMR (500 MHz, CDCl₃): δ 7.12 (m, 2H), 6.91 (t, 2H, J = 6.5 Hz), 6.79 (d, 2H, J = 8.5 Hz), 3.3 (s, 3H); ¹³C NMR (125 MHz, CDCl₃): δ 145.8, 127.4, 127.1, 123.4, 122.4, 114.0, 35.3; HRMS (m/z): [M+H]⁺ calculated for C₁₃H₁₂NS, 214.0690; found, 214.0714.

Acetylation (J-55), benzylaition (J-50) (Burger et al., 1954), Methylation (J-56) (Das et al., 2009) and benzoylation (J-66) (Darvesh et al., 2007) of the phenothiazine molecule were performed by the previously reported procedure (with considerable modification) and named as follows:

S3: General Synthetic Scheme for the compounds having same parent scaffold, phenothiazine
Supplementary Table 1. List of the chemical structure of J-50, J-51, J-55, J-65 and J-66 compounds:

| S.I. | Chemical structure with batch number | S.I. | Chemical structure with batch number |
|------|--------------------------------------|------|--------------------------------------|
| 1    | ![Chemical structure](image1) J-50    | 5    | ![Chemical structure](image2) J-56   |
| 2    | ![Chemical structure](image3) J-51    | 6    | ![Chemical structure](image4) J-65   |
| 3    | ![Chemical structure](image5) J-54    | 7    | ![Chemical structure](image6) J-66   |
| 4    | ![Chemical structure](image7) J-55    |      |                                      |

C. Inhibitor screening, IC₅₀ evaluation and mode of binding studies:

I. Inhibitor screening, IC₅₀ evaluation and mode of binding studies:

The DiscoverRx ADP Hunter™ (Eurofins DiscoverRx, Fremont, CA, US) assay was performed to measure the generation of ADP resulting from kinase phosphorylation of substrate. The enzyme-coupled reaction produces a positive fluorescent signal that is directly proportional to the ADP in the solution and hence, the activity of the kinase. The assay was carried out in triplicates in a solid-black, 96-well plate in a volume of 40 µl solvent containing
protein kinase (e.g. hTLK1B in this case), kinase substrate (e.g. Nek1 peptide in this case), ATP and 1X Kinase Reaction Buffer (KRB). The 1X KRB comprised of 15 mM Hepes, 20 mM NaCl, 1 mM EGTA, 0.02 % Tween 20, 10 mM MgCl2, and 0.1 % bovine gamma globulins, at pH 7.4. Reactions in each well were started immediately by adding ATP and kept going for half an hour at room temperature. 40 µl of proprietary Reagent A followed by 20 µl of proprietary Reagent B were then added to each well to deplete the unconsumed ATP, terminate the kinase reaction and convert ADP to ATP. In the end, 10 µl of a stop solution was added to each well to allow the newly synthesised ATP to be measured using a detectable resorufin fluorescence signal. The amounts of kinase, substrate, ATP, reaction temperature and incubation times were optimised for the optimal performance of the assay. For the inhibitor screening experiments, 2 µl of Staurosporine, THD, and the test compounds (J3-50, J3-51, J3-54, J3-55, J3-56, J3-57, J3-59, J3-60, J3-65, and J3-66) were used at a final concentration of 20 µM in the reaction. DMSO was used as a suitable control vehicle. For the dose-response curves and the mode of binding studies, the control and the test inhibitors, and the ATP concentrations were varied to get a twelve-point data set. The fluorescence signal generated was measured using a Tecan Infinite® M1000 Pro Plate Reader (Tecan Group Ltd., Mannedorf, Switzerland) using excitation/emission wavelengths of 530/590 nm with an integration time of 0.1 s. Statistical analyses were performed using GraphPad Prism 6 statistical software program (GraphPad Software Inc., CA, USA), and the fitting of the kinetic and dose-response data was done globally using a multiresponse non-linear regression model to get a best-fit value that applies to all data sets. Suitable negative control wells without the protein kinase, substrate and ATP were also included in the kinase assay.

**D. Quantitation of human TLK1 transcript variant 3 (qRT-PCR):**

We performed RNA isolation and RT-PCR analysis as described earlier (Singh et al., 2019) with LNCaP cells. TLK1 (NM_12290) primers from IDT (PrimeTime qPCR Primer Assays) were used and transcript levels were normalized by GAPDH levels.

TLK1B - Primer 1: GCAACTCCAGTAAATCTAGCTTCC.
TLK1B - Primer 2: GTCCCTGCTGAATCACACG.

**E. Behavioral assays in C. elegans** to assess the behavioral effects of J54 as a potential DR2 antagonist (anti-psychotic) were carried out as described in details in (Donohoe et al., 2006; Donohoe et al., 2009).
**F. Pharmacokinetic Studies on J54:** These were carried out at the preclinical testing lab of UTSW in Dallas, TX. 21 male CD-1 mice were dosed IP with 10 mg/kg J54. 0.2 ml/mouse formulated with 10% DMSO, 90% Corn Oil. Whole blood was collected in a syringe coated with ACD. Plasma was processed from whole blood by centrifugation at 10,000 rpm for 10 minutes. For the standards and QC's 98 ul & 98.8 ul of blank plasma was added to an eppendorf and spiked with 2 & 1.2 ul of initial standard. Std's, QC's & samples of 100 ul were then crashed with 200ul of methanol containing 0.15% formic acid and 12.5 ng/ml IS (final conc.). The samples were vortexed 15 sec, incubated at room temp for 10' and spun 2x 13,200 rpm in a standard microcentrifuge. The supernatant was then analyzed by LC-MS/MS. Vendor supplied plasma used in standards and QCs: Bioreclamation, LLC; lot #MSE284936; exp = 03/31/2020; ACD anticoagulant. Raw data and additional information can be obtained upon request.

### Supplementary Table 2 – related to Fig. 4

| Cell line     | Sub-G1 | G1   | S    | G2/M |
|---------------|--------|------|------|------|
| LNCaP Control | 0%     | 93%  | 4%   | 3%   |
| +BIC          | 2%     | 94%  | 2%   | 2%   |
| +J54          | 1%     | 94%  | 3%   | 2%   |
| +BIC+J54      | 33%    | 53%  | 4%   | 10%  |
| TRAMP-C2 Control | 3%   | 42%  | 26%  | 29%  |
| + BIC         | 7%     | 58%  | 21%  | 14%  |
| +J54          | 11%    | 46%  | 29%  | 14%  |
| +BIC+J54      | 26%    | 27%  | 21%  | 26%  |
Sup Table 3  Inhibitory Effect of Test Substances on Radioligand Binding to two recombinant Human Dopamine Receptors (DR2)

| Assay system        | J54 1×10^{-7} mol/L | THD 1×10^{-7} mol/L | Positive substance 1×10^{-5} mol/L |
|---------------------|----------------------|----------------------|-----------------------------------|
| Dopamine D1 (Human) | 12.17                | 67.69                | R(+-)-SCH-23390                   |
| Dopamine D3 (Human) | 19.77                | 95.57                | (±)-7-OH-DPAT                     |

**Receptor:** Human recombinant, PKI, Cat No. 6110513

50 mmol/L Tris-HCl (pH 7.4) containing 120 mmol/L NaCl, 5 mmol/L KCl, 2 mmol/L MgCl₂ and 1 mmol/L CaCl₂

Tracer: SCH23390, [N-methyl-³H]-, PKI, Cat No. NET930

**Receptor:** Human recombinant, PKI, Cat No. ES-173-M 50 mmol/L Tris-HCl (pH 7.4) containing 5 mmol/L MgCl₂

Tracer: 7-Hydroxy DPAT, R(+-)[³H]-, PKI, Cat No. NET1169

Duplicate samples were analyzed and averaged results are shown.

**Data Processing**

Inhibition ratios (%) were calculated from “100 – binding ratio”.

**Binding ratio:** [(B–N)/(B₀–N)]×100 (%)

B: Bound radioactivity in the presence of the test article (individual value)

B₀: Total bound radioactivity in the absence of the test article (mean value)

N: Non-specific bound radioactivity (mean value)
Key Resources:

| S. No. | Antibody Name                                | Catalog number | Company                  |
|--------|----------------------------------------------|----------------|--------------------------|
| 1      | TLK1 antibody [N2C2]                         | GTX102891      | Genetex                  |
| 2      | Rabbit NEK1 Antibody                         | A304–570A      | Bethyl Lab               |
| 3      | Purified custom pNek1 antibody               | custom         | Thermofisher             |
| 4      | H2A.X (Ser139)                               | 05-636         | Millipore                |
| 5      | Cleaved Caspase-3                            | 9579           | Cell signaling technology|
| 6      | Cleaved Caspase-3                            | 9661           | Cell signaling technology|
| 6      | GAPDH                                        | 2118S          | Cell signaling technology|
| 7      | Cleaved PARP (Asp214)(D64E10                 | 5625           | Cell signaling technology|
| 8      | Anti-PCNA Antibody                           | MAB424         | Millipore                |
| 9      | Phospho-Chk1 (Ser317)                        | 2344           | Cell signaling technology|
| 10     | Phospho-ATR (Thr1989)                        | 5801S          | Cell signaling technology|
| 11     | Alpha mouse HRP                              | 7076           | Cell signaling technology|
| 12     | Ki-67 (D3B5)                                 | 12202          | Cell signaling technology|
| 13     | p21 Waf1/Cip1                                | 2946           | Cell signaling technology|
| 14     | Alpha rabbit IgG-HRP                         | 7074S          | Cell signaling technology|
| 15     | Anti-beta Tubulin antibody Loading Control (HRP) | Ab218         | Abcam                    |
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Fig S1. Overlay of J54 and THD ligands within the active site of TLK1 – related to Fig. 2, from various viewpoints.

Fig S2. Docked pose of ligands to active site of D2 dopamine receptor – related to Fig. 2D, after 100 ns of MD simulation: (a) J54, (b) THD and (c) risperidone.

Fig. S3 H\textsuperscript{3}NMR and MS of J54 – related to Fig. 1, “inhibitory compounds” figure.

Fig. S4 H\textsuperscript{3}NMR and MS of J3-56 - related to Fig. 1, “inhibitory compounds” figure.

Fig. S5 Pharmacokinetics plasma distribution of J54 – related to Fig. 6. These were carried out by Noelle Williams, Director of Preclinical Pharmacology Core at UTSW, and additional information on the study and data/analyses is available upon request.

Fig. S6 Behavioral assays in C. elegans - Related to Fig. 2D – reduced of DR2 binding activity. A For the pharyngeal pumping assay, wild-type animals (N2 strain) were transferred to plates with diluted DMSO (Control) or drug (J-54 or trifluoperazine) at 160 µM final concentration for 90 min before quantifying pharyngeal pumping see SI methods). We counted visible movement of the grinder (pharyngeal contractions) for 30 sec to obtain the pumping rate (N = 45). Note that in pharyngeal pumping, J54 reduced the pumping rate but to a lesser extent than a much lower concentration of TFP. Inhibition of pharyngeal pumping by trifluoperazine (TFP), a typical PTH antipsychotic, is attributed to its known activity as a dopamine and calmodulin antagonist (40).

B. Foraging, regulated by serotonin and dopamine, was studied in wild-type animals evaluated in the presence of DMSO (Control) or J54 at a high dose of 160 µM vs. TFP at 40 µM. The number of omega turns (head touches body) and reversals were then counted over the next 3 min to quantify search behavior for each group (N = 13 per group).

C. Reduction of dopamine-induced immobility. Dopamine produces immobility in C. elegans after 2-3 hr of exposure. Antipsychotic drugs, such as haloperidol, that potently block D2 dopamine receptors largely counteract the effects of dopamine, i.e., more animals continue to
move on plates that include dopamine plus drug. Wild-type animals were incubated on 60 agar plates with bacteria and diluted DMSO (Control) or J54 or haloperidol at 160 µM final concentrations. After 1 hr on these plates, they were transferred to 60 mm agar plates with bacteria that also contained dopamine (final concentration 25 mM) plus diluted DMSO (Control) or else J-54 or haloperidol at final concentrations of 160 µM. After 3 hr on the dopamine plates, we examined movement and counted animals as moving if they traversed half their body length in either the forward or backward direction during a 5-sec observation period. We then calculated the percentage of animals moving and repeated this experiment three times to confirm the effects of drug. J54 was not toxic to worms even after prolonged exposure.

Fig. S7 A. H&E stain of representative LNCaP tumors (from 3 mice per group) - Related to Fig. 6. Note that in the BIC+J54 tumor, there is lesser cancer cellularity, with “empty” spaces interspersed with fibrous tissue, clearly suggesting the combination is having an effect in reducing tumor growth or killing it. B) qRT-PCR quantitation of TLK1B mRNA expression (from 3 LNCaP samples per group in triplicate reactions). C) Activation of pATM-S1981 following 1 day incubation of LNCaP and TRAMP-C2 cells with BIC and J54. D) Reduction of Rad9 phosphorylation by J54 and THD – related to Fig. 5. LNCaP cells were treated for 12h with 10 µM J54 or THD, and pRad9-S328 or total Rad9 were determined by WB.

Fig. S8. IHC analysis of markers of proliferation (Ki67 - A), apoptosis (Cleaved PARP and Caspase 3 – B, C), and presence of γH2AX (an indicator of DSBs – related to Fig. 6. Quantitation of the stained sections is shown on the side. Representative section from 3 mice per group are shown. Two way-Anova tests were done to compare the groups for statistical significance. Related to Fig. 6.
A.

B.

J54 10mpk IP PK Plasma

Concentration (ng/ml)

Time (mins)

C.

J54 10mpk IP PK Plasma

Rsq = 0.5308  Rsq_adjusted = 0.2962  HL_Lambda z = 287.1297

4 points used in calculation

Predicted (ng/ml), Mean_Conc (ng/ml)

Time (min)
A. Effect of J54 on Pharyngeal Pumping

B. Effect of J54 on Foraging

C. Effect of J54 on DA-induced Immobility
A. Control, BIC, J54, Combination

B. Relative TLK1B mRNA Expression

C. LNCaP

D. CNT, J54, THD

TRAMP-C2
