Quantitative Systems Pharmacology Analysis of KRAS G12C Covalent Inhibitors

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KRAS has proven difficult to target pharmacologically. Two strategies have recently been described for covalently targeting the most common KRAS mutant in lung cancer, KRAS G12C. Previously, we developed a computational model of the processes that regulate Ras activation. Here, we use this model to investigate KRAS G12C covalent inhibitors. We updated the model to include Ras protein turnover, and validation demonstrates that our model performs well in areas of G12C targeting where conventional wisdom struggles. We then used the model to investigate possible strategies to improve KRAS G12C inhibitors and identified GEF loading as a mechanism that could improve efficacy. Our simulations also found resistance-promoting mutations may reverse which class of KRAS G12C inhibitor inhibits the system better, suggesting that there may be value to pursuing both types of KRAS G12C inhibitors. Overall, this work demonstrates areas in which systems biology approaches can inform Ras drug development.

Study Highlights

WHAT IS THE CURRENT KNOWLEDGE ON THE TOPIC?
☑️ The individual mechanisms of Ras signal regulation and of covalent KRAS G12C inhibitors are known. How the combined actions of the different processes that regulate Ras signals influence the efficacy of covalent inhibitors is neither well known nor easily inferred.

WHAT QUESTION DID THIS STUDY ADDRESS?
☑️ Can mathematical modeling account for the observed, nonintuitive, behaviors of Ras inhibitors? How can covalent inhibitor strategies be improved?

WHAT DOES THIS STUDY ADD TO OUR KNOWLEDGE?
☑️ This study finds that observed, nonintuitive, behaviors are fully consistent with known Ras biology and could have been predicted a priori. The study also highlights the role of Ras GEFs in oncogenic Ras regulation and targeting.

HOW MIGHT THIS CHANGE DRUG DISCOVERY, DEVELOPMENT, AND/OR THERAPEUTICS?
☑️ The identification of properties that could make NPIs efficacious suggests new assays for screening drug candidates. The identification that which of the two strategies is better can reverse upon resistance promoting mutations suggests benefit to continued pursuit of both strategies.

Human cancer cells, such as those from pancreatic cancer, colorectal cancer, and lung cancer, commonly include somatically acquired KRAS mutations.1–3 The Ras proteins (KRAS, NRAS, and HRAS) bind to guanine nucleotides GDP and guanosine-triphosphate (GTP) with high affinity.4 The Ras proteins can hydrolyze GTP to GDP, which is why they are commonly referred to as “GTPases”. The GTP-bound form of Ras is considered the active form, and downstream signaling effectors specifically bind to the GTP-bound form of Ras.5 The cancer-promoting KRAS mutations most commonly occur at codon 12, 13, or 61, and result in increased levels of GTP-bound KRAS, which in turn promotes downstream signaling.6 It has long been believed that drugs with the ability to block aberrant KRAS signaling would benefit patients with cancer.4

A major advance in the targeting of KRAS has been made with the development of molecules that covalently and irreversibly bind to the cysteine residue of the KRAS G12C mutant.7–10 The KRAS G12C mutant is particularly common in lung cancers. Selectivity of a drug for the cysteine in this mutant protein should result in specific targeting of KRAS G12C mutant-containing cancer cells. Compounds that covalently interact with the codon 12 cysteine can be grouped into two different classes based upon which pocket the molecule settles into after bonding with cysteine. The compounds described by the Gray laboratory rest in the nucleotide pocket (NP) in place of a guanine nucleotide.8 The G12C inhibitors that reside in the NP will be referred to as NP-inhibitors (or NPIs). The compounds initially described by the Shokat laboratory rest in a distinct pocket, the “switch two pocket” that was previously unappreciated.7 Compounds using this strategy will be referred to as SIIP-inhibitors (or SIIPIs). The originally described NPIs and SIIPIs were not able to reliably target KRAS G12C in cell culture. A new SIIP, ARS-853, has been described that has much improved binding properties and displays KRAS G12C inhibition in cell culture.11
Table 1 Parameters for the G12C model

| Parameter                | Value       | Units | Description                                      |
|--------------------------|-------------|-------|-------------------------------------------------|
| $k_{	ext{deg}}$         | $8 \times 10^{-6}$ | /s    | Ras degradation rate                            |
| $k_{\text{production}}$  | $3.2 \times 10^{-12}$ | /Ms   | Total Ras production rate                       |
| $k_{\text{on,NPI}}$      | $2 \times 10^{6}$ | /Ms   | On rate for NPI                                 |
| $k_{\text{on,SIIPI}}$    | 0.12        | /Ms   | On rate for SIIPI (compound 12)                 |
| $k_{\text{on,SIIPI,ARS}}$| 76          | /Ms   | On rate for SIIPI (ARS-853)                     |
| Percent_KRAS             | 50          | %     | Percentage of total Ras that is KRAS           |
| Percent_Mut              | 50          | %     | Percentage of KRAS that is mutant              |
| Scalingfactor$_{\text{GTPase,G12C}}$| 72 | %         | Intrinsic GTPase rate of G12C relative to that of wild-type Ras |
| Scalingfactor$_{\text{GDP,G12C}}$ | 120 | %     | Rate of effector dissociation of G12C relative to that of wild-type Ras |
| GTP                      | $1.8 \times 10^{-4}$ | M     | Cellular abundance of GTP                       |
| GDP                      | $1.8 \times 10^{-5}$ | M     | Cellular abundance of GDP                       |

GTPase, guanosine triphosphate; NPI, nucleotide pocket inhibitor; SIIPI, switch II pocket inhibitor.

 Previously, we developed a computational model of the processes that together regulate Ras signaling. The model has made multiple predictions about Ras signaling that went against conventional wisdom but have now been experimentally confirmed. For example, our model predicted that oncogenic Ras leads to wild-type Ras activation, which has since been experimentally demonstrated by multiple laboratories. More recently, we predicted that mutations to Ras GAP NF1 should frequently co-occur with noncanonical cancer driver genes in the Ras pathway, which was subsequently detected in melanoma genomes. Here, we applied the model to the problem of targeting KRAS G12C. To do this, the model was updated to include protein turnover, the specific biochemical properties of the G12C mutant, and the described mechanisms of interaction for the two classes of inhibitors. Simulations of the updated model yield patterns of Ras activation that match well with previous experimental observations. Application of the model to NPIs finds that GEF loading is a property that could be optimized to make NPIs more effective. The model also suggests that mutations that promote resistance to SIIPIs may increase sensitivity to NPIs. Overall, this work demonstrates the potential of mechanistically accurate models of oncogenic signal regulation to contribute to quantitative systems pharmacology.

METHODS

The details of the Ras model have been previously published and described in detail. The model is specified as a set of coupled, nonlinear, ordinary differential equations. Simulations in MATLAB were used to find steady-state levels of RasGTP and RasGTP-Effector complex, both of which were considered as measures of Ras pathway signaling. Mutant Ras proteins are specified in terms of how each parameter describing a Ras biochemical reaction differs from the wild-type Ras value. We specify these as multiplicative “scaling factors” that are equal to the ratio between the mutant parameter and the wild-type parameter found within the same experimental study. Table 1 provides parameters unique to this model, or that are particularly relevant to understanding the present study.

RESULTS

Model development and analysis

Model extension to include protein turnover and the G12C mutant. To study the G12C mutant targeted by a covalent inhibitor, the model was updated in two key ways. First, protein synthesis and degradation were added to the model. When the model was originally developed, we assumed that these processes would be slow relative to the processes that regulate steady-state Ras nucleotide binding and could, therefore, be ignored when studying oncogenic mutant activation. Because covalent inhibitors target G12C irreversibly (and could potentially increase degradation), we added a degradative process to the model. A schematic with the updates to the original model is shown in Figure 1a. Briefly, the model now allows for degradation of Ras in any form, and production of Ras results in nucleotide-free Ras, which will rapidly bind to available cytoplasmic nucleotides upon modeled production. Protein degradation was modeled as a first order process with a half-life of ~24 hours ($k_{\text{deg}} = 8 \times 10^{-6}$/s). Production was modeled to occur at a constant rate. This rate was set so the steady-state level of total Ras remains at the level of our original estimate for total cellular Ras. Values for these parameters are provided in Table 1.

We also updated the model to include the G12C mutant. Of note, oncogenic mutants at codons 12, 13, and 61 are biochemically similar in that they are insensitive to inactivation by Ras GAPs. However, there are subtle differences with other parameters between different oncogenic mutants that, in some cases, can have large effects. For example, patients with colon cancer with the oncogenic G13D KRAS mutation seem to benefit from cetuximab, whereas patients with colon cancer with the oncogenic G12D or G12V KRAS mutations seem to not benefit from cetuximab. Therefore, we conclude that it is most prudent to model the G12C KRAS mutant by incorporating recently published data. Specifically, the intrinsic GTPase rate and the effector binding constant are two parameters of the G12C mutant that vary from wild type. The values used in our model are presented in Table 1.
Tests of the extended model with turnover and the G12C mutant. We first wanted to compare the behavior of the model extended to include protein turnover with the previous version of the model that did not include turnover. For test cases, we considered cells with all wild-type Ras, that contain the G12C Ras mutant, and that include the G12D, G12V, or F28L Ras mutants studied in our original Ras model. We first evaluated the behavior of the updated model by examining the predicted level of RasGTP and RasGTP-effector complex for these different cases. For these conditions, we modeled 25% of total RAS as mutant and 75% as wild type, such as might occur if 50% of the RAS protein in the cell was KRAS, the other 50% NRAS and HRAS, and one of the KRAS alleles was mutated. Model predictions for levels of RasGTP and RasGTP-effector were essentially the same whether or not Ras turnover was included (Figure 1b). This confirmed the validity of our original assumption that turnover could be ignored.
for predicting behavior of oncogenic mutants in the absence of covalent inhibitors.

We also investigated the behavior of the updated model by considering the loss of the tumor suppressor protein neurofibromin (NF1). The NF1 is a Ras GTPase activating protein (Ras-GAP). Ras-GAPs act as negative regulators of RasGTP by promoting the conversion of RasGTP to RasGDP. Germline absence of a single copy of NF1 results in increased RasGTP, and loss of both copies of NF1 results in further increases in RasGTP. NF1 is also one of the most commonly mutated driver genes in human cancer, in which one or both copies can be mutated, deleted, and/or silenced.

Modeling 100% and 50% loss of total Ras-GAP in both the model with turnover and without turnover resulted in similar levels of RasGTP and RasGTP-effector complex (Figure 1c). We also considered fractions of total basal GAP loss other than 50% and 100% to consider conditions in which other GAPs contribute to RasGTP homeostasis. We considered 25% loss and 75% loss to span the range of levels of GAP activity, and found similar levels of RasGTP and RasGTP-effector when the model with turnover was compared to the model without turnover. Overall, these simulations found that updating our model to include turnover results in essentially identical predictions for steady-state levels of Ras signal, whether caused by oncogenes or by loss of tumor suppressor genes.

Extension of the model to covalent G12C inhibitors. We next extended the model to include the two types of KRAS G12C targeted covalent inhibitors (Figure 2a,b). Our turnover model posits that NPIs bind to nucleotide-free G12C mutant irreversibly, consistent with the manner in which NPIs have been described to date. The SIIPIs are reported to bind to GDP bound G12C mutant (and presumably nucleotide-free G12C mutant, as well), to bind irreversibly, and to prevent the inhibitor bound protein from exchanging GDP for GTP. We modeled SIIPIs with these same activities. Table 1 includes the reaction parameters for NPIs and SIIPIs. Of note, the reported difference between the original SIIPI referred to as compound 12 and the new SIIPI ARS-853 are manifested exclusively in the on-rate of drug binding.

Model validation: Testing the model’s ability to reproduce the behaviors of G12C inhibitors. We then simulated dose responses for NPI and SIIPI (Figure 2c). The NPI and SIIPI would seem similarly effective at steady-state if turnover was not modeled. However, when turnover was included in the model, the SIIPI (ARS-853 compound) was clearly superior to the other SIIPI and to the NPI. In this more physiological case, the maximal amount of drug that will bind to KRAS G12C is limited by the rate of each compound binding to KRAS G12C, which in the case of the NPI is restricted by the infrequency of nucleotide-free mutated KRAS. This highlights the well-recognized importance of considering turnover when evaluating covalent inhibitors.

Simulated behaviors of these drugs are consistent with their described abilities to inhibit KRAS G12C containing cancer cell lines, suggesting that our model includes the aspects of Ras biology needed to study KRAS G12C targeted covalent inhibitors. Experimentally, only the ARS-853...
The Ras model predicts how receptor tyrosine kinase activation and inhibition influence ARS-853 inhibition. Simulations of Ras G12C network inhibition by the switch II pocket inhibitor (SIIPI) ARS-853 were performed, including conditions of upstream activation and upstream inhibition. (a) Simulations of the kinetics of the inhibition found increased GEF activity (upstream activation) delays the kinetics of inhibition (green), whereas decreased GEF activity results in faster inhibition (red). Kinetics for the basal model (no change in GEF activity) is shown in black. (b) The steady-state dose responses of ARS-853 in these same conditions of increased (dotted line) or decreased (dashed line) GEF activity were also found through model simulations. Basal GEF activity is shown with the solid line. For both (a) and (b), the plot on the left is normalized to the total amount of Ras signal in each case, whereas the plot on the right shows total RasGTP in each case.
KRAS mutations that had both the G12C mutation as well as another mutation at a separate KRAS residue to introduce additional biochemical defects. They considered several different experiments that tested the inhibition of Ras signaling with the newly developed G12C inhibitors. We found that the model could readily reproduce experimentally observed behaviors, suggesting that our model is valid for problems involving G12C inhibitors. Additionally, it is worth noting that multiple experimental behaviors that were interpreted as contrary to conventional wisdom could have been predicted a priori had our readily available model first been applied to these problems.

Model validation: Conclusion. We considered several different experiments that tested the inhibition of Ras signaling with the newly developed G12C inhibitors. We found that the model could readily reproduce experimentally observed behaviors, suggesting that our model is valid for problems involving G12C inhibitors. Additionally, it is worth noting that multiple experimental behaviors that were interpreted as contrary to conventional wisdom could have been predicted a priori had our readily available model first been applied to these problems.

Analysis of NPIs

We next used the model to investigate NPIs. We considered why NPIs are less effective than SIIPIs (Figure 5). A key step for the drug, as described, is binding to a nonoccupied nucleotide binding pocket. A nonoccupied nucleotide binding pocket is likely to be rarely encountered within the cell due both to the high affinity of the pocket for guanine nucleotides and also the high concentration of guanine nucleotides within the cell.

We initially modeled NPI to bind the Ras NP with an on-rate similar to that for nucleotides (k_{on} = 2 \times 10^{7}/Ms). We next considered on-rates that were one, two, and three orders of magnitude faster to evaluate how much better inhibition would be if the on-rate could be increased. Computational simulations suggested that a significant enhancement of the on-rate would result in a much improved dose response, even when considering that nucleotide-free Ras protein is very limited within the cell (Figure 5a). However these levels are approaching the theoretical diffusion limit, suggesting that it would not be possible to engineer a better NPI by only optimizing the forward reaction rate constant.

We considered whether GEFs could facilitate the loading of an NPI. The authors of the NPI studies evaluated the ability of their compound to compete with GTP and GDP. Their assay included recombinant Ras protein, but did not include RasGEFs, Ras-GAPs, nor Ras-effectors. Thus, it did not assess how the network of reactions that together influence nucleotide binding would impact drug binding. As this compound cannot pass through the cell membrane, a cell-based assay is not a simple option for investigating this problem. Mathematical modeling provides one mechanism by which we can extrapolate to the relevant, physiological, network and investigate this problem.

We were particularly interested in whether or not Ras GEFs could facilitate NPI loading. Our previous modeling
studies of Ras suggest that a small level of basal GEF activity is needed to explain the experimental data of basal nucleotide exchange and that GEF inhibition should result in less oncogenic Ras signaling. We, therefore, considered the potential effects of GEF loading on NPIs.

Our simulations suggest that GEF loading could be an important variable for NPIs. If GEF loads the G12C inhibitor as well as it loads the nucleotide (e.g., it loads the G12C inhibitor like GTP and GDP, and at a rate that is proportional to the abundances of drug and nucleotides), our simulations suggest that there would be a modest increase in the ability of NPIs to target the G12C mutant (Figure 5b, solid blue line). If GEF loading could favor drug loading by a factor of 10 to 100 over nucleotide, the effectiveness of this class of drug is predicted to be much higher (Figure 5b, blue dotted and dashed lines). This computational analysis, therefore, suggests that efforts to optimize NPIs should evaluate candidates for whether or not GEFs facilitate their loading. Of note, the predicted levels of optimized (GEF-facilitated) NPI (oNPI) needed to inhibit 50% of the Ras signal were on the order of 1 μM, much smaller than the near mM levels of nucleotide that are found within the cell.

Analysis of oNPI on resistance-promoting mutations

We next considered how oNPI would respond to the mechanisms described to promote resistance to SIIPI. Our simulations found that increased intrinsic nucleotide cycling, such as what occurs with Y40A, N116H, and A146V compound mutations with G12C, could actually result in an increased sensitivity to NPI (Figure 6a). This suggests that there may be value in the continued development of oNPIs. In contrast, our simulations found that secondary mutations to KRAS that impair GTPase activity (Figure 6b) or mutations that increase RTK activation (Figure 6c) would both result in less sensitivity to NPI. This suggests that an oNPI would not be able to combat all potential forms of G12C-acquired resistance. It is worth noting that although the oNPI was less effective than the SIIPI for a G12C mutant, the oNPI was more effective than the SIIPI for all three resistance-promoting mutations.

DISCUSSION

There is great interest in the covalent targeting of KRAS G12C, and multiple inhibitors exist at different stages of development. Few of these compounds seem capable of passing through the cell membrane. Within the cell, Ras does not function in isolation but is rather a part of a complex signaling network. Modeling provides an approach for extrapolating to cellular conditions to evaluate how candidates in development may function in more physiological conditions. Modeling may also provide an approach for identifying which properties could be optimized within these physiological conditions, even before a derivative capable of being reliably delivered into the cell is developed. For investigating how a mutant will influence a signaling network, it is important that the model includes the proteins with which Ras directly interacts. Many models have been developed of the signaling networks that include Ras, and these typically include at least one representative GEF, GAP, and effector. It is much less common for pathway models that include Ras to dive into the details of Ras nucleotide exchange at the level needed to study oncogenic Ras activation, although a few models at this level of detail have been developed. Systems pharmacology may be an area in which this additional detail becomes critical.

One interesting finding from the present systems pharmacology study is that it suggests NPIs could be more effective if GEFs can promote their loading into the NP. This possibility seems consistent with known GEF biology and needs to be considered experimentally. However, the ability of Ras GEFs to load NPIs has not, to the best of our knowledge, been addressed to date. The reason it has not been addressed previously may be because it is widely believed that the increased activation of oncogenic Ras is independent of GEFs. Our work suggests that efforts to screen derivatives of the known NPIs and new NPI candidates should include the evaluation of how well Ras GEFs can load the compound. This could be done experimentally
Figure 6 Simulations find switch II pocket inhibitor (SIPI) resistant promoting mutations can make nucleotide pocket inhibitors (NPIs) the more effective KRAS G12C inhibitor. Potential SIPI resistance promoting mutations were studied for both an optimized NPI (oNPI) and an SIPI. The oNPI was assumed to have its loading facilitated by GEFs by a factor of four over the proportional abundance of NPI to total guanine nucleotides. (a) For the G12C mutant, the SIPI ARS-853 (black solid) was predicted to be superior to the oNPI (red solid), but once a secondary mutation that caused increased nucleotide exchange was modeled (100× over G12C) simulations found a decreased effectiveness of the SIPI (black dashed), but a higher effectiveness for the oNPI (red dashed). (b) When a secondary mutation that causes decreased intrinsic GTPase activity was modeled (100× relative to G12C), both the SIPI (black dashed) and oNPI (red dashed) were less effective. In the conditions of the secondary mutation, the oNPI became relatively more effective than the SIPI. (c) When a secondary mutation increased RAS activation through increased GEF activation (e.g., as might happen with an RTK mutation), the oNPI (red dashed line) became more effective than the SIPI (black dashed), although both classes of inhibitor are predicted to be overall less effective against mutations that result in increased GEF activation.
by adapting the recombinant protein based assay used by Hunter et al.⁹ to also include a recombinant GEF and determining how NPI binding to RAS is enhanced by GEFs. Our work further suggests that compounds that are loaded by GEFs preferentially over nucleotides would be most beneficial.

Another interesting finding was that, theoretically, the better inhibitor could change as secondary resistance mutations are acquired. For example, the modeled GEF-facilitated NPI is inferior to the SIIPI with respect to inhibiting KRAS G12C signaling. However, the model suggests that the GEF-facilitated NPI should be superior to the SIIPI if a G12C mutant picks up a second mutation to the same allele that result in faster nucleotide dissociation.

A major challenge in developing targeted therapies for cancer is that it can be difficult to anticipate the response of a biological network to an inhibitor. Indeed, multiple experiments using KRAS G12C inhibitors have been interpreted as unexpected and contradictory to conventional wisdom. Computational systems biology models can provide an alternative viewpoint to expert opinion. It is possible to use a computational model to find the systems level behaviors that naturally emerge from the constituent reactions when the model is based upon the fundamental reactions of the network. We have done that here, and we demonstrate that multiple experimental observations that were interpreted as unexpected were foreseeable by computational modeling. As there are ongoing efforts to advance both strategies,²⁶⁻³⁹ it may be valuable to integrate modeling into the drug development process.

Mathematical models have previously contributed to the study of G12C inhibitors. Modeling has been used to help analyze and interpret kinetic data.¹¹ Modeling has also been used to extrapolate from data on reversible inhibitors to irreversible inhibitors. That study also included an argument that Ras GEFs may accelerate binding kinetics.⁴⁹ Our model differs from these other models in that it includes a more complete set of Ras regulatory reactions and also protein turnover. Although this detail may be more than is needed for the questions addressed by the other models, this additional information is needed to evaluate signal inhibition in the cellular context (Figure 2). We anticipate that models like these will be increasingly utilized to inform efforts to develop and use targeted therapies.

Our consideration of methods to improve covalent Ras inhibitors made the assumption that increased Ras inhibition would be beneficial. Currently, most of these inhibitors display limited activity and limited selectivity, so increasing the level of Ras inhibition seems like a valid goal. However, it is possible that partial inhibition is sufficient to kill cancer cells and/or that modest changes smaller in magnitude than those studied here may have large clinical effects. At this early stage of development, we believe that our assumption that “more inhibition is better” is reasonable. As this class of drugs matures, it will be worthwhile to revisit this assumption.

Altogether, we have applied a computational systems biology approach to the analysis of targeted covalent inhibitors. Specifically, we used a model of the biochemical reaction network that regulates Ras signals to study how two specific KRAS G12C oncogenic mutant inhibition strategies alter Ras signaling. Our model finds the behaviors that logically follow from what is known about Ras signaling and about these inhibitors. We find that our model naturally reproduces the experimentally observed behaviors of both NPIs and SIIPIs, including results that were widely perceived as unexpected and inconsistent with known Ras biology. Our model also suggests strategies to improve the effectiveness of one of these classes of inhibitors.

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