

A novel pan-RAS inhibitor for Malignant Peripheral Nerve Sheath Tumors

Howard Donninger^{1,3}, Rachel Ferril², Destine Ede², Becca vonBaby², Raphael Jigo²,
Joe Burlison^{1,3}, Mike Sabo^{1,3}, John Trent^{1,3} and Geoffrey J. Clark^{1,2}

¹Molecular Targets Group, James Graham Brown Cancer Center, Departments of ²Pharmacology & Toxicology and ³Medicine, School of Medicine, University of Louisville, Louisville, KY, USA

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Introduction

Neurofibromatosis type 1 is a genetic disease that results from either heritable or spontaneous autosomal dominant mutations in the NF1 gene. Neurofibromatosis type 1 individuals frequently suffer benign tumors known as Plexiform Neurofibromas which develop from cranial and peripheral nerve sheaths. Plexiform Neurofibromas have the potential to develop into malignant peripheral nerve sheath tumor (MPNST). MPNST exhibit a low overall 5 year survival rate of less than 40%, and there is no effective treatment or cure.

The NF1 gene encodes the protein Neurofibromin. Neurofibromin is a negative regulator (a GAP) for the notorious RAS oncoprotein. Although RAS is frequently activated by mutation in many cancers, this is not the case with NF1 disease. Here, the wild type RAS protein is stabilized in the active configuration due to the loss of NF1 function (Figure 1). This is a transforming event that drives the disease. Currently, there are no targeted inhibitors of wild type RAS that are effective in the clinic.

In an attempt to combat the problem of a lack of a therapeutic treatment for Neurofibromatosis Type 1, and indeed, RAS driven cancer in general, we have performed *in silico* screening of two million compounds followed by bioassay to identify a small molecule, referred to as F3, that binds and inhibits active RAS by blocking its ability to interact with its effectors. We have subsequently used a medicinal chemistry approach to identify more effective derivatives of F3. Our current lead is designated F3-8-60, which exhibits enhanced anti-RAS biological activity *in vitro* and enhanced RAS binding.

In vivo, F3-8-60 inhibits the metastasis of an MPNST cell line and suppresses the growth of MPNST pdx. We observe no toxicity associated with the drug. We are currently continuing chemical optimization of the agent and appear to have identified variants with further improved wild type RAS binding activity. We propose this approach may lead to novel therapeutics for NF1 disease.

Discussion

NF1 disease is largely caused by deregulation of RAS due to loss of function of NF1. We have been working on developing a small molecule that binds the wild type form of RAS and blocks its ability to interact with its effectors. Our current lead, designated F3-8-60, binds to K-RAS with low uM affinity. It suppresses RAS signaling in MPNST cell lines and is active *in vitro* and *in vivo*. Moreover, it is active against MPNST pdx, suggesting there is real clinical potential in these compounds. We are continuing to optimize the drug via Medicinal Chemistry and iterative screening. ADME/PK studies are ongoing. As at least some of the cognitive issues associated with NF1 patients also appear to be due to aberrant RAS activity, and as our compounds can pass the blood brain barrier, we also hypothesize they may have utility in treating neurological defects caused by excess RAS activity.

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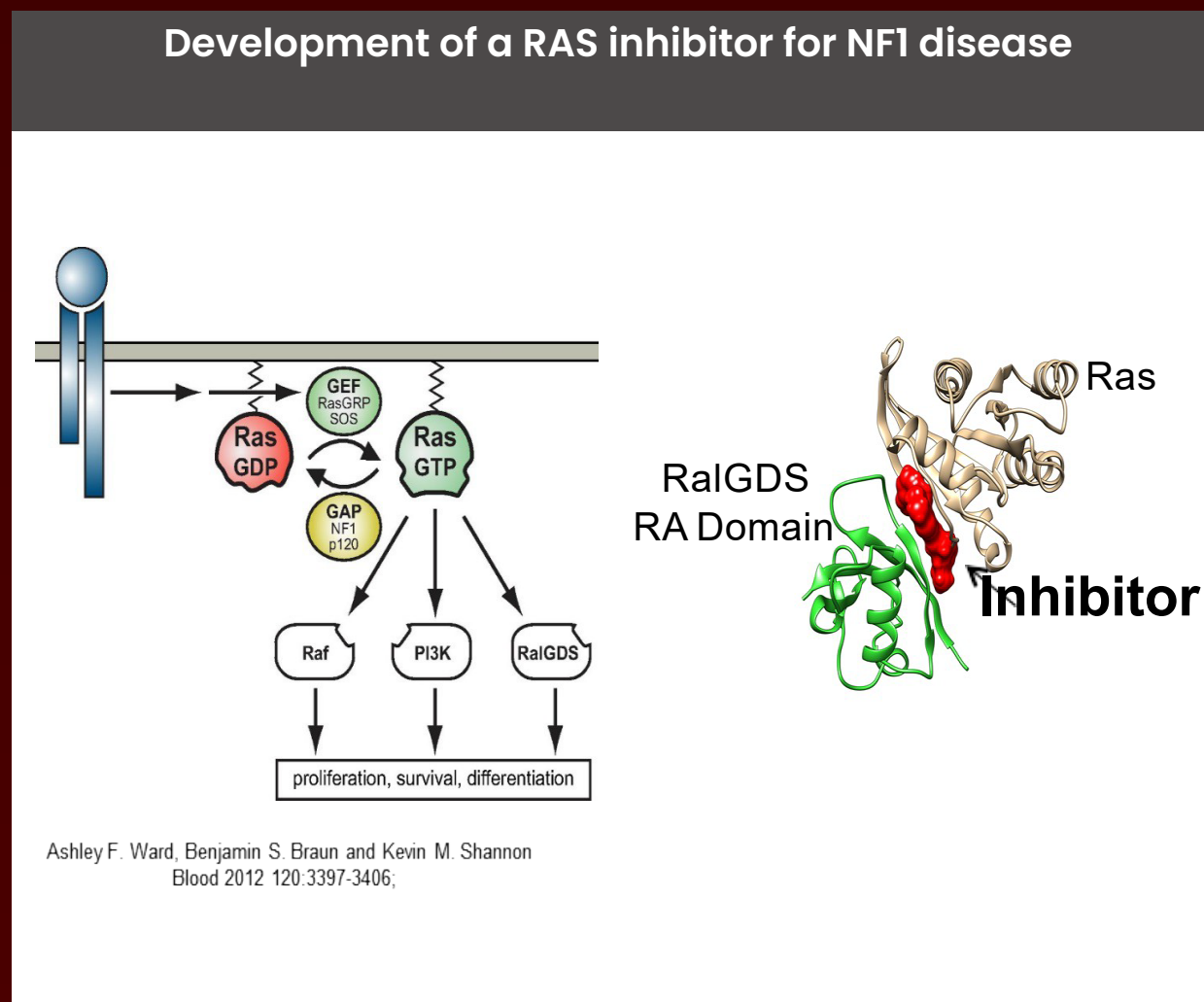


Figure 1. Development of a RAS inhibitor for NF1 disease.

LEFT: Ras activity is controlled by positive and negative regulators. NF1 is negative regulator, so inactivation of NF1 removes a "brake" on RAS resulting in an elevation of RAS activity. RIGHT: using the known crystal structure of RAS in complex with its effector RalGDS, we used *in silico* screening to predict compounds which might bind to RAS and block its effector interactions. *In silico* positives were then screened *in vitro* in 3D culture assays to identify "hits".

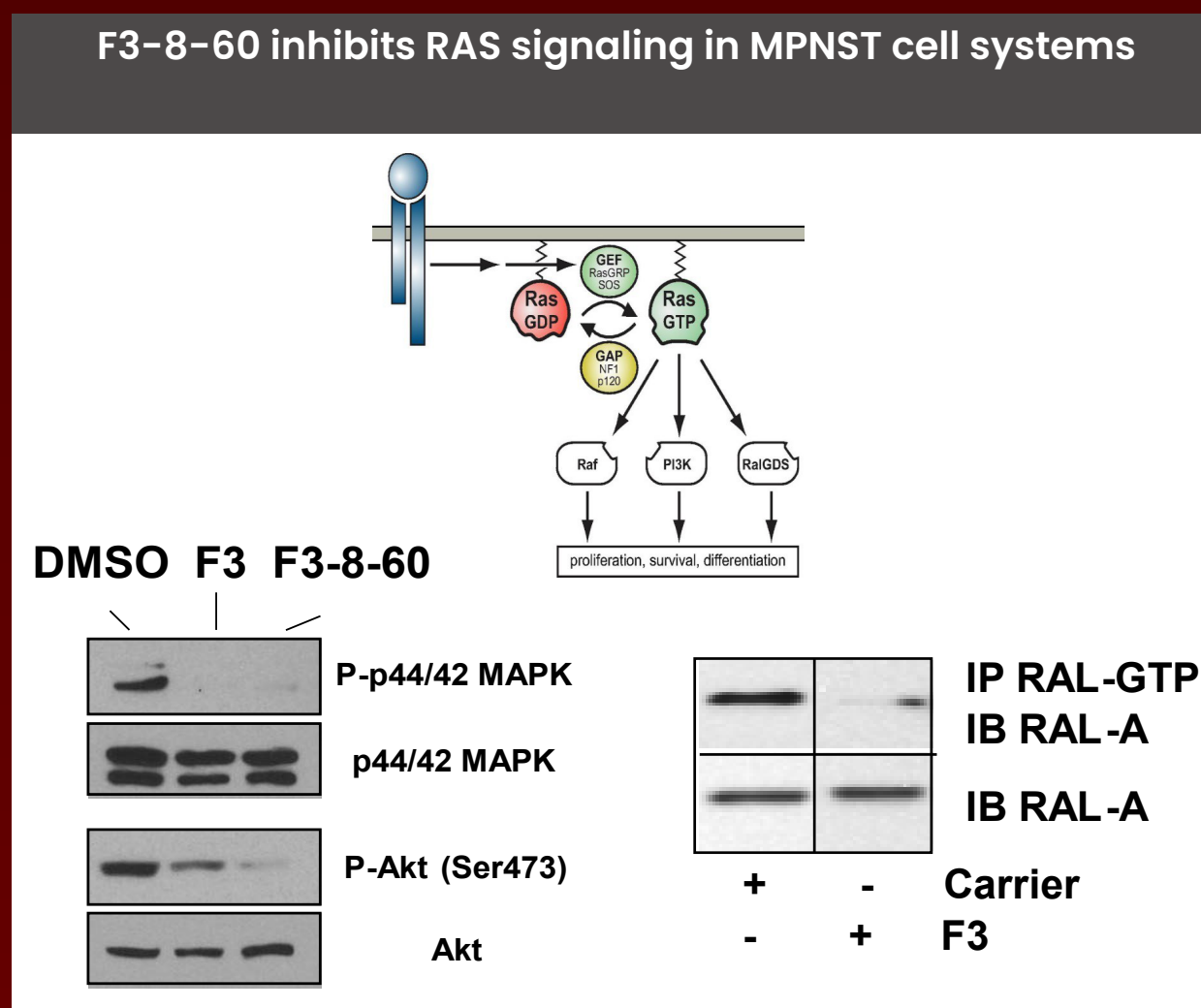


Figure 5. F3 class compounds block RAS signaling.

Top panel: cartoon of RAS signaling pathways. Left Panel: F3 and F3-8-60 inhibit the levels of phospho-ERK (a measure of MAPK pathway signaling) and phospho AKT (a measure of PI3K signaling) in MPNST cells (S46.2TY). Right Panel: F3 suppresses RAL signaling in S46.2TY MPNST cells. Cells were treated with 20uM drug for 1 hour.

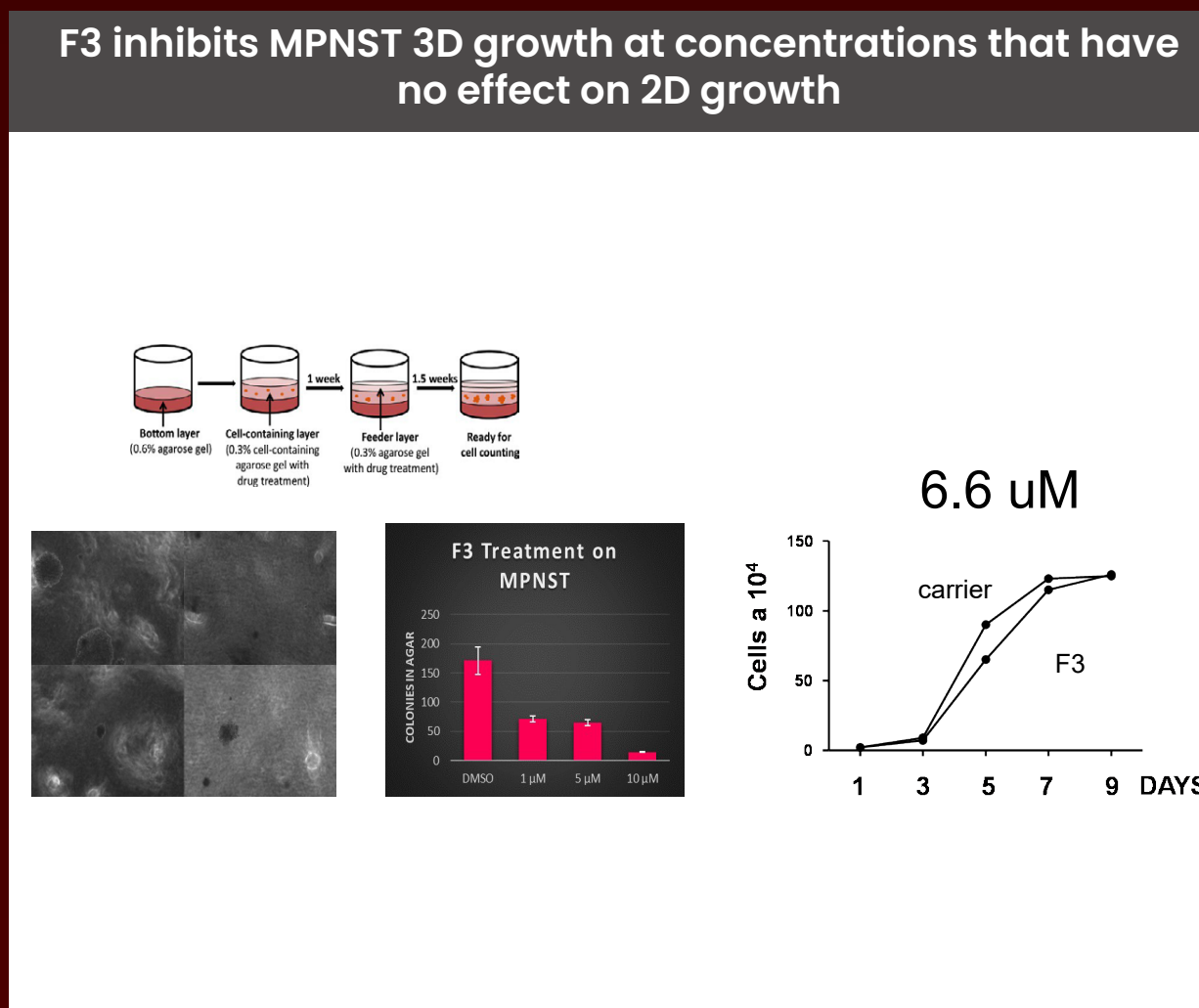


Figure 2. Anti-RAS compound F3 potently suppresses 3D growth of MPNST cells.

MPNST cells (S462.TY) were plated in soft agar in the presence or absence of test drug. Colony formation was scored after 2 weeks. Compound F3 scored as one of the most positive. An IC50 of ~1uM could be obtained. Left panels: 3D soft agar assays of F3 against an MPNST tumor cell line. Right panel: F3 had no effect on 2D cell growth.

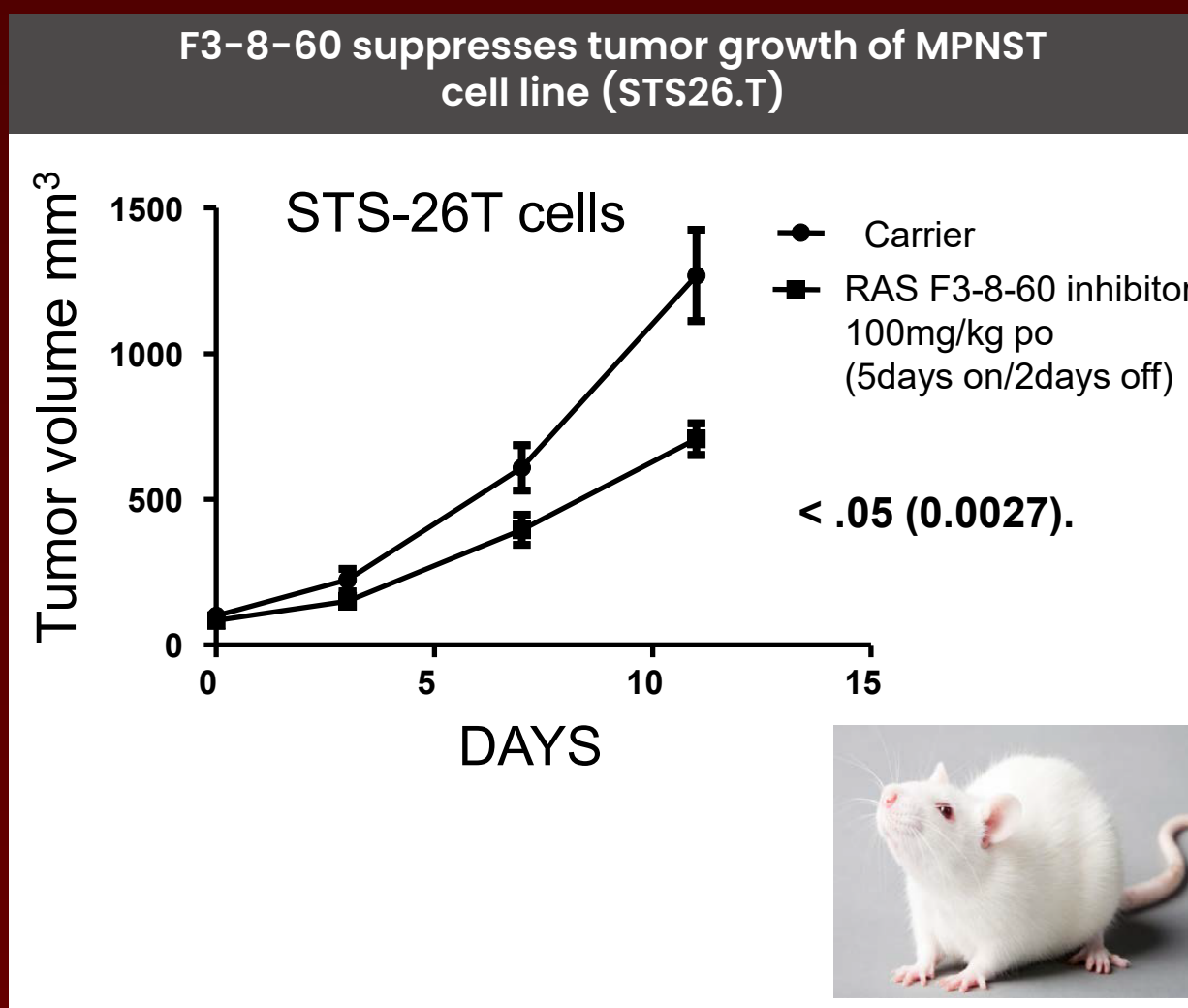


Figure 6. F3-8-60 Suppresses MPNST cell line tumor growth

STS-26T cell are an NF1 deficient human MPNST cell line. Cells were implanted into the flank of NSG mice (n=6). When tumors arose to between 50-100mm³ the mouse was randomly assigned to an experimental group and gavaged with drug. Average tumor volume is plotted.

Results

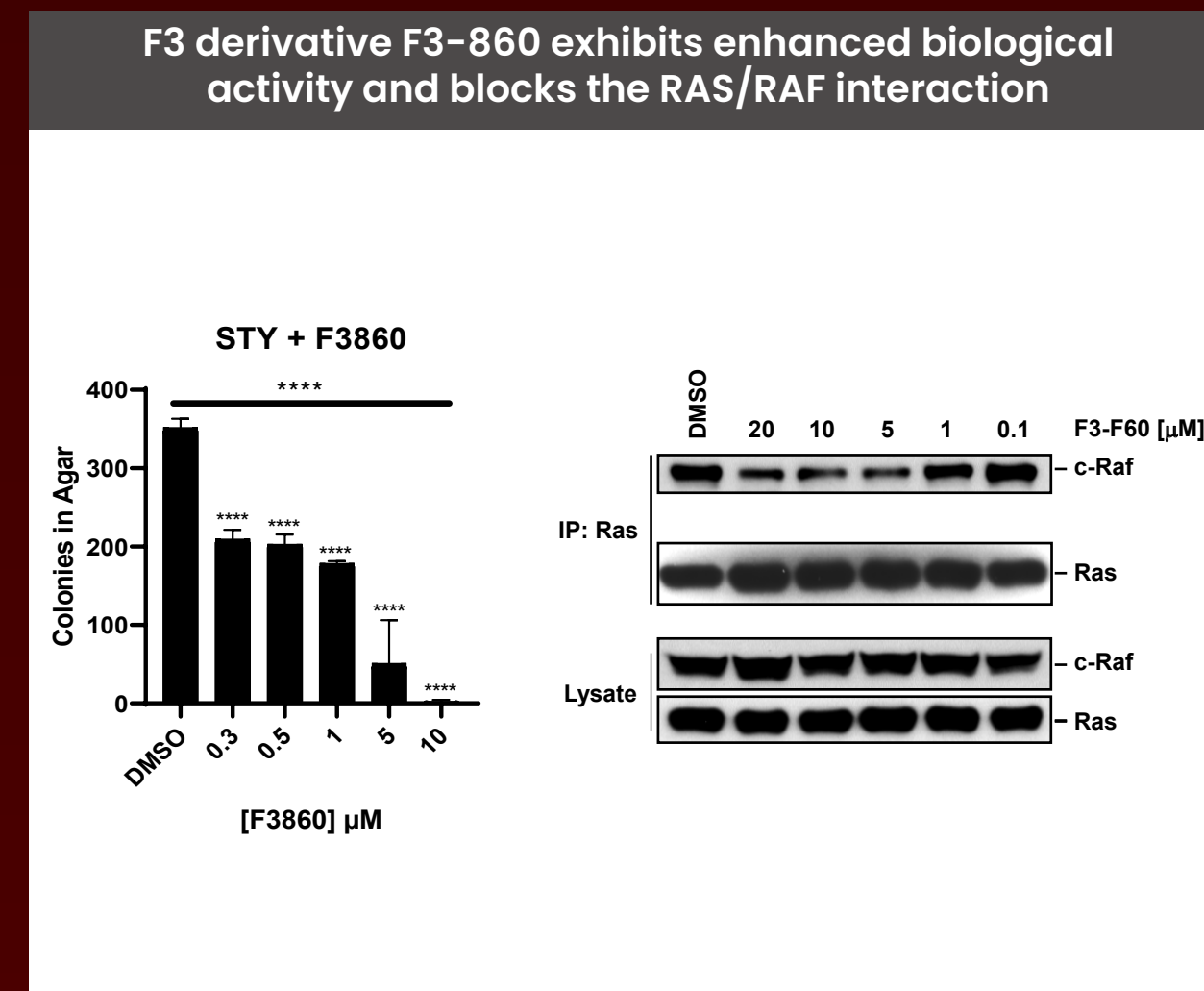


Figure 3. Inhibition of RAS effector interactions and signaling by enhanced activity derivative F3-8-60.

LEFT: F3 derivative F3-8-60 is more active against MPNST cell lines (S462.TY). RIGHT: Endogenous, mutant K-RAS was immunoprecipitated (IP) in the presence or absence of various concentrations of F3-860. The complexes were immunoblotted (IB) for RAS and c-RAF to determine the effects on RAS/RAF binding.

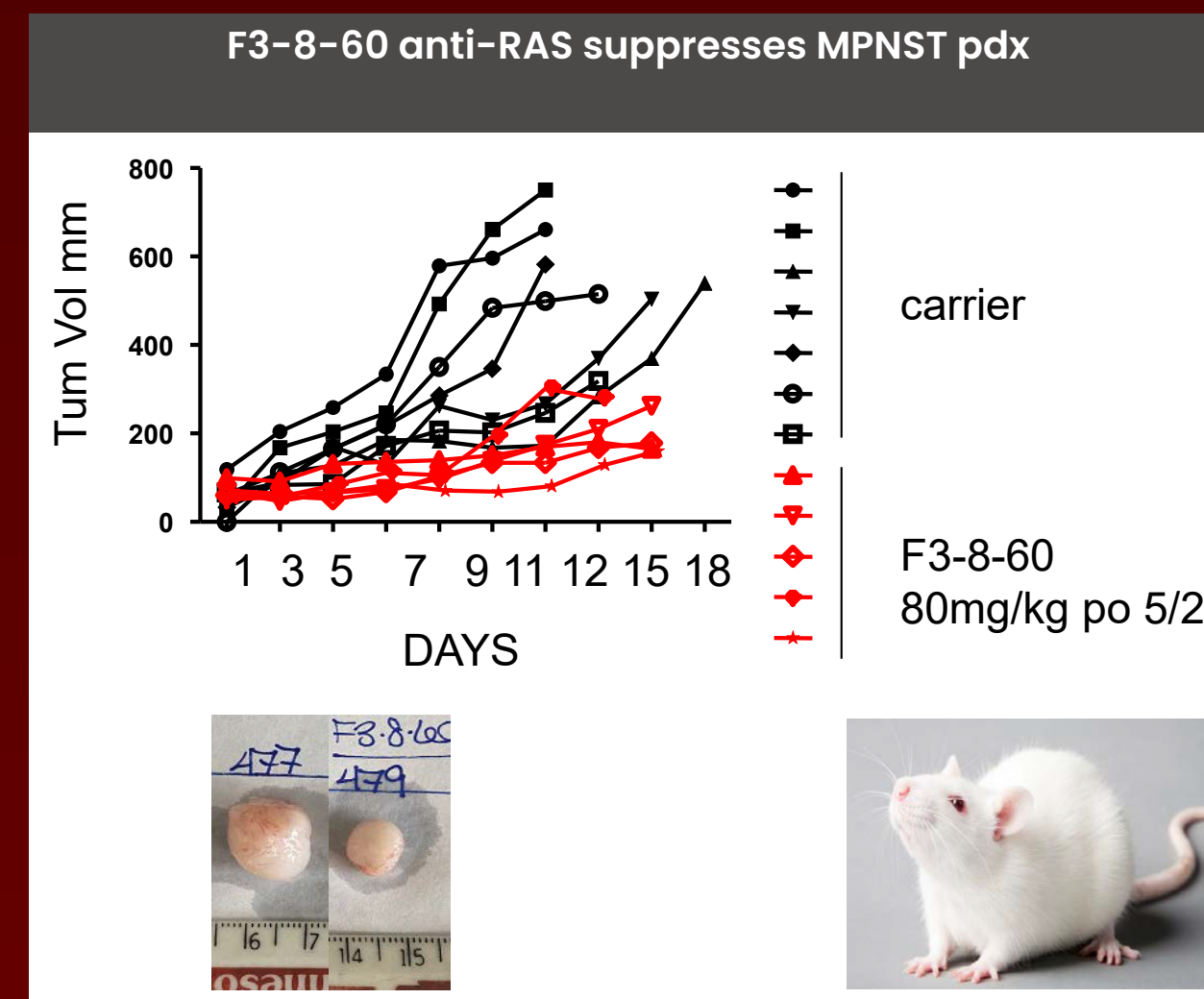


Figure 7. F3-8-60 suppresses MPNST pdx growth

To date, the best model for human tumor drug response is the Primary tumor graft system. Here, an MPNST primary tumor (gracious gift, Johns Hopkins NF1 Biospecimen repository) was grafted into a cohort of NSG mice. Graftees were randomized into control and experimental pools. When a tumor graft reached 50-100 mm³, it was treated with carrier or F3-8-60 at 80mg/kg 5 days on/2 days off by gavage for three weeks.

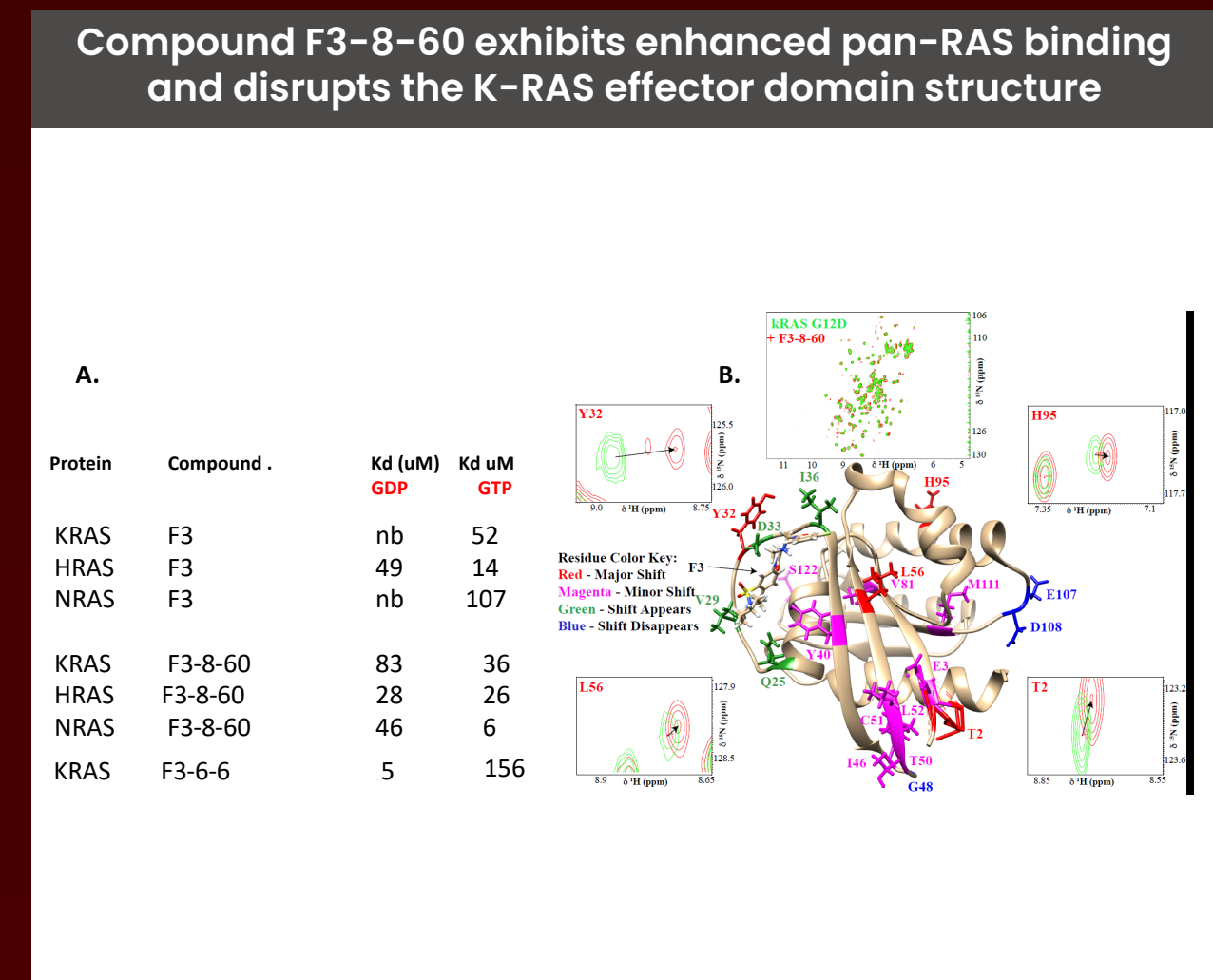


Figure 4. Compound F3 and enhanced activity derivative F3-8-60 directly bind to K-RAS. Activated K-RAS protein was prepared and used in direct compound binding assays.

A. Microscale Thermophoresis was performed in order to obtain Kd values for F3 and F3-8-60. Newer variants are being tested that exhibit higher affinities, e.g. F3-6-6. B. NMR analysis of F3-8-60 bound to K-RAS12D shows significant alterations in effector loop structure

CONCLUSIONS

1. We have identified a series of direct Novel RAS inhibitors.
2. The current lead compound is active *in vivo* against wild type RAS driven MPNST tumor systems, including primary tumor grafts.
3. They exhibit no apparent *in vivo* toxicity.
4. These agents may be developed into novel targeted therapy for NF1 disease.