Targeting primary and metastatic uveal melanoma with a G protein inhibitor

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Recent evidence suggests that mutant constitutively active Gq/11 could be targeted therapeutically in UM and other diseases (14). Oncogenic Gq/11 α-subunits are constitutively active due to defects in hydrolyzing GTP to GDP, the rate-limiting step for G protein deactivation. Nevertheless, oncogenic Gq/11 can be inhibited by FR900359 (FR) or YM-254890 (YM) (15–20), a pair of closely related, bioavailable cyclic depsipeptides that allosterically inhibit GDP release by the Gq/11/14 subfamily (16, 21). Inhibition is thought to occur because oncogenic Gq/11 α-subunits can spontaneously release GTP, stochastically bind GDP, bind YM or FR (YM/FR) and Gβγ subunits (17), forming complexes resistant to activation by G-protein-coupled receptors (14, 16) and diminishing downstream oncogenic signaling (17–20). In response to YM/FR, UM cell lines driven by oncogenic Gq/11 arrest the cell cycle and can undergo apoptosis and/or redifferentiation into melanocytic-like cells (17–20). In contrast, BRAF-driven UM cell lines, which express wild-type Gq/11, are unaffected (17–20).

The therapeutic potential of YM/FR in UM is an important opportunity based on this evidence. Critical issues and questions remain to be addressed, however. YM/FR may exert deleterious as well as therapeutically beneficial effects, because they do not discriminate between wild-type and oncogenic Gq/11 (14), and target Gq/11-dependent physiological systems (21–24) essential for homeostasis and viability (25–28). Indeed, YM/FR administered systemically at levels that reduce Gq/11 activity in host systems greater than 50% is likely to be lethal, as suggested by gene dosage studies of Gq/11 knockout mice (29). Indeed, how strongly oncogenic Gq/11 signaling in UM tumors must be inhibited for therapeutic effect remains unknown. Furthermore, although recent studies have shown that YM/FR can inhibit growth of UM tumors in mouse xenograft models (19, 30), questions left unanswered are whether YM/FR can target primary and/or metastatic UM tumor cells from patients, whether YM/FR responsiveness is tumor class or metastatic potential. FR administered systemically below its LD50 strongly inhibited growth of, but did not efficiently kill, UM tumor cells from biopsies of primary or metastatic tumors. FR evoked melanocytic redifferentiation of UM tumor cells with low (class 1), but not high (class 2), metastatic potential. FR administered systemically below its LD50 strongly inhibited growth of PDX-derived class 1 and class 2 UM tumors in mouse xenograft models and reduced blood pressure transiently. FR did not regress xenografted UM tumors or significantly affect heart rate, liver function, hematopoiesis, or behavior. These results indicated the existence of a therapeutic window in which FR can be explored for treating UM and potentially other diseases caused by constitutively active Gq/11.

**Uveal melanoma (UM)** is the most common intraocular tumor in adults. Nearly half of UM patients develop metastatic disease and often succumb within months because effective therapy is lacking. A novel therapeutic approach has been suggested by the discovery that UM cell lines driven by mutant constitutively active Gq or G11 can be targeted by FR900359 (FR) or YM-254890, which are bioavailable, selective inhibitors of the Gq/11/14 subfamily of heterotrimeric G proteins. Here, we have addressed the therapeutic potential of FR for UM. We found that FR inhibited all oncogenic Gq/11 mutants reported in UM. FR arrested growth of all Gq/11-driven UM cell lines tested, but induced apoptosis only in a few. Similarly, FR inhibited growth of, but did not efficiently kill, UM tumor cells from biopsies of primary or metastatic tumors. FR evoked melanocytic redifferentiation of UM tumor cells with low (class 1), but not high (class 2), metastatic potential. FR administered systemically below its LD50 strongly inhibited growth of PDX-derived class 1 and class 2 UM tumors in mouse xenograft models and reduced blood pressure transiently. FR did not regress xenografted UM tumors or significantly affect heart rate, liver function, hematopoiesis, or behavior. These results indicated the existence of a therapeutic window in which FR can be explored for treating UM and potentially other diseases caused by constitutively active Gq/11.

**Mutant constitutively active forms of Gq or G11 (Gq/11) α-subunits of heterotrimeric G proteins cause uveal melanoma (UM)** (1–3) and several other diseases and disorders (4–11). UM is particularly devastating because patients with metastatic disease often succumb within months due to lack of effective therapy (12). Clinical trials of metastatic UM have shown little benefit of cytotoxic chemotherapeutics, immune checkpoint inhibitors, or small-molecule inhibitors of oncogenic signaling proteins including protein kinase C and MEK (13). New therapeutic strategies for metastatic UM clearly are warranted.

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deleterious effects on viability or host physiological systems. Here, we have addressed all of these important issues.

Results

FR inhibits all oncogenic Gq and G11 mutants that drive uveal melanoma

At least ten different constitutively active Gq/11 mutants have been identified in UM (2, 31, 32). Only four had been studied previously and shown to be inhibited by FR or YM (17–20). To extend this analysis, we transiently expressed each of ten different Gq/11 mutants that drive UM (Fig. 1A) with a Gq/11-driven transcriptional reporter in a common cellular context (HEK293 cells) to exclude extrinsic factors that might affect the results. We found that each mutant G protein induced reporter expression and was inhibited by FR with similar potency (IC$_{50}$ 1.9–3.8 nM) and efficacy (Fig. 1A).

In contrast, we found that four UM cell lines (Fig. 1B) driven by three different oncogenic Gq/11 mutants responded to FR in quantitatively or qualitatively different ways. FR inhibited growth of all four Gq/11-driven UM cell lines, but with potencies differing by as much as ~80-fold (IC$_{50}$ 0.5 nM (MP41 cells), 38 nM (Mel270 cells); Fig. 1B). Whereas FR arrested the cell cycle of all Gq/11-driven UM cell lines, as indicated by reduction of S- and G2/M-phase cells (Figs. 1C and S1), it induced significant apoptosis only in two of them (92.1 and Mel270 cells), as indicated by accumulation of cells with sub-G0/G1 DNA content (Figs. 1C and S1). FR did not affect proliferation or survival of a BRAF(V600E)-driven UM cell line (Fig. 1B), as shown

Figure 1. Effects of FR on constitutively active Gq/11 mutants and UM cell lines. A, effects of FR on expression of an SRE.L reporter driven by transfection of oncogenic Gq/11 mutants in HEK293 cells. FR inhibited all ten Gq/11 mutants reported in UM, with IC$_{50}$s between 1.9 nM (Gq-Q209 L) and 3.8 nM (Gq-R183Q). Representative means and SEMs were calculated from triplicate values from one of at least three independent experiments for each mutant. B, effects of FR on UM cell lines. Growth of UM cell lines driven by constitutively active Gq or G11 mutants was inhibited by FR with IC$_{50}$s between 0.5 nM (MP41) and 38 nM (Mel270). A UM cell line (OCM-1A) driven by constitutively active BRAF(V600E) was unaffected by FR. Representative means and SEMs were calculated from triplicate values from one of at least three independent experiments for each cell line. C, effect of FR on apoptosis and cell cycle progression of UM cell lines analyzed by flow cytometry. Induction of apoptosis was indicated by increased proportion of cells with sub-G1/G0 DNA content. Cell cycle arrest was indicated by decreased proportion of cells with S- and G2/M-phase DNA content. Representative means and SEMs calculated from triplicate values from one of at least three independent experiments for each cell line.
before (17–19), demonstrating the selectivity of FR for oncogenic Gq/11 in UM.

**FR redifferentiates class 1 but not class 2 UM cells**

Class 1 and class 2 primary UM tumors have low (~5%) and high (~90%) probability of metastasis, respectively (33, 34). Class 1 UM tumors express the metastasis suppressor, BAP1, encoding an enzyme that regulates gene expression epigenetically by deubiquitinating histone H2A. Class 2 UM tumors are BAP1-deficient (35) and possess additional genetic defects rendering them prone to metastasis (32). Almost all UM cell lines are derived from class 1 tumors (36), raising the crucial question of whether both classes of primary UM tumor cells can be targeted by FR.

We addressed this question initially by analyzing MP41 (class 1; BAP1+) and MP46 (class 2; BAP1-deficient) cell lines, which were established originally from patient-derived xenografts (PDX) of primary UM tumors (37). We found that FR arrested growth and cell cycle progression of both of these cell lines without triggering significant apoptosis (Fig. 1, B and C).

Next, we used transcriptional analysis to determine whether FR could re differentiate MP41 (class 1) and MP46 (class 2) cells into melanocytic-like cells, as we had shown previously with two other class 1 UM cell lines (92.1 and Mel202) (17). MP41 and MP46 cells both responded transcriptionally to FR, as indicated by multidimensional scaling analysis of RNAseq data (Fig. 2A). However, a cluster of genes was downregulated preferentially in class 1 MP41 cells (circled in Fig. 2B), as
revealed by analyzing transcriptional responses on a gene-by-gene basis. We defined this cluster as genes with log2 fold change in MP41 cells < -2.5 in response to FR, which identified 365 genes that were more significantly downregulated in MP41 cells (mean p = 0.04) than MP46 cells (mean p = 0.17). Many genes in this cluster are targets of epigenetic regulation by polycomb repressive complex 2 (PRC2), based on gene set analysis (H3K27trimethylation, EZH2; Fig. 2C). This PRC2-targeted gene cluster was coordinately repressed by FR in class 1 MP41 cells as compared with class 2 MP46 cells (Fig. 2D).

FR targets UM cells from primary and metastatic tumors

Because UM tumor cell lines do not fully recapitulate important properties of UM tumors, we determined whether FR can target UM cells obtained from patient biopsies of primary and metastatic tumors. We began by preparing...
separate self-renewing cultures of UM tumor cells isolated from fine-needle biopsies of class 1 primary tumors from six patients and core biopsies of liver metastases from two patients. We found that all eight individually cultured-cell samples responded to FR, as indicated by lower cell numbers relative to vehicle controls (Fig. 3A), and we noted, as shown before (36, 37), that class 2 primary UM tumor samples proliferated too poorly in vitro to establish self-renewing cultures. Accordingly, each time thereafter we obtained biopsy samples from additional patients, we prepared short-term cultures and analyzed the effects of FR. A total of ten biopsy samples (Table S1), including class 1 and class 2 primary tumors and a liver metastasis, were analyzed. Cultured cells from nine of these patient samples responded to FR relative to vehicle controls, as indicated by decreased yields of total RNA as a marker of cell number (Fig. 3B). The single nonresponding sample (UM032; Fig. 3B) expressed wild-type Gq and G11 (Table S1) and thus was driven by other oncogenes whose identities were not determined. For the nine FR-responsive UM tumor samples, which included class 1 and class 2 primary tumors and a liver metastasis, we found that FR caused similar directional effects on gene expression, as revealed by unsupervised principal component analysis (PCA) (Fig. 3C).

Interestingly, the principal component that distinguished class 1 from class 2 and metastatic tumor cells (Fig. 3D; y-axis) (33) was relatively unaffected by FR (Fig. 3D), suggesting that FR did not cause class 2 UM tumor samples to lose their class identity and become more class 1-like. The one UM tumor sample that expressed wild-type Gq and G11 did not show significant directional effects of FR on gene expression (Fig. 3C). Together, these results demonstrated that, regardless of tumor class, primary and metastatic UM tumor samples driven by mutant Gq/11 responded ex vivo to FR.

Because patient-derived UM tumor biopsies include cancer and stromal cells, we determined which cell types were targeted by FR, as indicated by their relative abundance and transcriptional responses revealed by single-cell RNAseq (scRNAseq). Primary tumor samples from three UM patients undergoing enucleation were used to obtain sufficient quantity of cells for these experiments. Dissociated UM tumor and stromal cells were treated 7 days with FR at a concentration (100 nM) that strongly inhibited all oncogenic Gq/11 mutants (Fig. 1A). Transcriptional signatures readily distinguished UM cells from stromal cell types (Fig. 3E). In all three biopsy samples, FR modestly reduced the abundance of UM cells and had little effect on the abundance of non-UM cells (Fig. 3E). FR therefore did not cause extensive death of UM cells.

Further analysis of scRNAseq data indicated that UM cells, but not stromal cells, displayed striking transcriptional responses to FR. As visualized by t-SNE plots of scRNAseq data (Fig. 3E), FR dramatically shifted the transcriptional profile of UM cells in all three biopsy samples. These changes in UM gene expression in tumor samples from enucleated patients were analogous to those observed previously by bulk RNAseq of small biopsy samples (Fig. 3D). In contrast, FR did not significantly affect the transcriptional profiles of stromal cells in all three biopsy samples (Fig. 3E). UM tumor cells therefore were targeted transcriptionally by FR.

Transcriptional analysis also indicated that FR inhibited cell cycle progression of patient-derived UM tumor cells. FR-evoked changes in gene expression detected by scRNAseq in UM cells from enucleated tumors and bulk RNAseq of small biopsies were similar, as shown by hallmark enrichment analysis (Fig. 4A). In both cases, targets of E2F transcription factors (arrows in Fig. 4, A and B) were among the gene sets prominently downregulated by FR. Moreover, FR treatment of class 1 and class 2 biopsy samples downregulated expression of the E2F transcription factors E2F1, E2F2, and E2F8 (Fig. 4C) and upregulated expression of the cell cycle inhibitors RB1 and p130 (38). In sum, inhibition of oncogenic Gq/11 by FR upregulated Rb expression and downregulated E2F expression, thereby contributing to inhibition of UM cell cycle progression (Fig. 4D).

Next, we investigated whether tumor class, which predicts metastatic potential and patient prognosis, determines whether FR evokes melanocytic redifferentiation of UM tumor cells in patient biopsies, as we had observed above with UM cell lines (Fig. 2). When we examined transcriptional responses of patient-derived class 1 UM tumor cells to FR, we found strong enrichment of gene sets targeted by PRC2 (Fig. 4E). Nearly all members of the FR-responsive PRC2-targeted gene cluster that were identified in UM cell lines (Fig. 2) were also downregulated in class 1 UM tumor biopsies (Fig. 4F). In contrast, many of these genes were upregulated, rather than downregulated by FR in class 2 UM tumor biopsies (Fig. 4F). Furthermore, melanocyte differentiation genes, including MITF, TYR, and TYRP1, were upregulated significantly by FR in class 1 relative to class 2 tumor biopsy samples (Fig. 4G). Therefore, FR treatment potentially could inhibit progression of class 1 primary UM tumors by promoting melanocytic redifferentiation.

**FR up- and downregulates YAP1-driven genes**

Prior studies of UM have shown that oncogenic Gq/11 signaling leads to activation of the YAP1 transcription factor (19, 20). We therefore determined whether YAP-targeted genes are coordinately downregulated by FR in biopsy samples of class 1 and class 2 primary UM tumors. In contrast to this expectation, we found that FR upregulated and downregulated YAP-targeted genes in nearly equal proportion (Fig. S2). The mechanisms responsible for the complex effects of FR on YAP-targeted genes are unknown, but potentially involve the combinatorial actions of other transcription factors with YAP1 on the promoters of these genes.

**A therapeutic window for targeting UM tumors with FR**

Because inhibition of Gq/11 activity by more than 50% is likely to be lethal, as indicated by gene dosage studies in knockout mice (29), we sought to define protocols and conditions under which chronic, systemic administration of FR is physiologically tolerated, so we could assess the therapeutic potential of FR in mouse xenograft models of UM. Based on

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**EDITORS’ PICK: FR900359 targets primary and metastatic uveal melanoma**

**FR up- and downregulates YAP1-driven genes**

**A therapeutic window for targeting UM tumors with FR**
Figure 4. Pathway analysis of FR-induced transcriptional responses in UM. A, hallmark gene-set enrichment analysis of differential gene expression in response to FR. Left panel shows gene sets affected by FR in enucleation-derived UM cells identified by scRNAseq. Right panel shows gene sets affected by FR in nine biopsy samples described in Figure 3 as detected by RNAseq. Red arrow indicates that E2F target genes were significantly downregulated in both scRNAseq and bulk RNAseq datasets. B, enrichment plot of the E2F target hallmark gene set detected by bulk RNAseq. Heatmap below shows insignificant difference in response of this gene set to FR in class 1 versus class 2 biopsy samples. C, effects of FR on expression of Rb/E2F family members in class 1 and class 2 biopsy samples detected by RNAseq. Asterisks indicate significant effect (p < 0.01; paired t-test) relative to vehicle controls. Differences between class 1 and class 2 samples were insignificant. D, proposed model for cell cycle regulation by constitutively active Gq/11 and FR. Signaling by constitutively active Gq/11 leads to repression of Rb expression and induction of E2F expression that drives UM cell proliferation. FR attenuates these effects to arrest cell cycle progression. E, differential responses of class 1 and class 2 tumor biopsy samples to FR. Gene-set enrichment analysis identifies genes downregulated preferentially in class 1 relative to class 2 tumor biopsy samples (log-rank t-test). Red arrows indicate PRC2-specific gene sets. F, heat map showing effects of FR on PRC2-targeted genes in class 1 versus class 2 tumor samples. G, responses of melanocytic differentiation genes to FR in class 1 and class 2 tumor biopsy samples. Asterisks indicate statistically significant (p < 0.01; paired t-test) effects of FR on the indicated melanocytic differentiation genes in class 1 and class 2 biopsy samples.
prior studies examining the acute effects of FR or the closely related inhibitor YM-254890 (YM) (22, 24, 39), we administered FR by subcutaneous injection at 0.1–3.0 mg/kg on alternate days for 30 days to define an LD\textsubscript{50} of ~0.6 mg/kg in NOD-scid-gamma (NSG) mice used as xenograft recipients (Fig. 5A). We then determined the effect of FR administered chronically below its LD\textsubscript{50} (0.1 or 0.3 mg/kg by subcutaneous injection on alternate days) on several physiological parameters of NSG mice. Blood pressure, a sensitive indicator of Gq/11 activity (40), decreased significantly in conscious mice upon each injection of FR relative to vehicle controls (Fig. 5B), but rebounded to baseline within 24 h (Fig. 5C). Heart rate was affected insignificantly by FR (Fig. 5B). No evidence of anemia, monocyte or platelet deficiency (Table S2), or liver dysfunction (Fig. 5D) was observed after treating NSG mice 30 days with FR at either dose relative to vehicle controls. Lastly, when administered for 1 week at the higher dose, FR had insignificant effect on the activity, exploratory behavior, or sensorimotor function of NSG mice (Table 1 and Fig. S3), even though Gq/11 are important regulators of nervous system functions.

**Figure 5. Effects of systemically administered FR on mouse viability and physiology.** A, survival analysis. NSG mice were treated up to 30 days by s.c. injection of vehicle or FR at the indicated doses on alternate days (vehicle, n = 34 mice; FR 0.1 mg/kg, n = 22 mice; FR 0.3 mg/kg, n = 22 mice; FR 1.0 mg/kg, n = 10 mice; FR 3.0 mg/kg, n = 13 mice). Kaplan–Meier (upper panel) and survival (lower panel) curves indicate an LD\textsubscript{50} of ~0.6 mg/kg. B, effect of FR on blood pressure as an indicator of Gq/11 inhibition in host systems. Adult male NSG mice received subcutaneous administration of vehicle (2 μl DMSO in 200 μl of 5% dextrose, saline; n = 7 mice) or FR (0.3 mg/kg, n = 9 mice) once every other day for 14 days. Blood pressure and heart rate were monitored telemetrically 1 h before and 3 h after injection. Injection times are indicated by green arrows. Days without radiotelemetry recording or injection are indicated by gray columns. C, twenty-four-hour continuous radiotelemetry recordings of blood pressure and heart rate are indicated by gray columns. Circles, twenty-four-hour continuous radiotelemetry recordings of blood pressure and heart rate are indicated by gray columns. D, Effect of FR liver function. Blood samples were collected from mice after 30 days of treatment with vehicle or FR (s.c. on alternate days; n = 10 mice per group) and assayed for liver enzyme activities. No significant changes were seen with FR treatment. DBP, diastolic blood pressure; HR, heart rate; MBP, mean blood pressure; SBP, systolic blood pressure.
development and activity (40). Chronic systemic administration of FR below its LD50 therefore had modest or undetectable effects on physiological systems that potentially could limit dosing regimens in UM clinical trials.

Next, we determined whether FR delivered systemically at physiologically tolerated doses could inhibit growth of xenografted UM tumors driven specifically by oncogenic Gq/11. These experiments used MP41 and MP46 cells, for several reasons. These UM cell lines were isolated originally from patient-derived xenografts, provide models of class 1 and class 2 tumors, and, of the UM cell lines tested, responded to FR (cell cycle arrest, with little or no apoptosis) most like UM tumor cells from patient biopsies. We also used a UM cell line (OCM-1A) driven by BRAF(V600E) to determine whether effects of FR were specific for Gq/11-driven tumors. Accordingly, subcutaneous UM tumor xenografts of MP41, MP46, and OCM-1A cells were established, and vehicle or FR was administered systemically at low (0.1 mg/kg) or high (0.3 mg/kg) dose on alternate days by subcutaneous injection on the contralateral side. Tumor size was measured over time and tumor weight was determined at the end of the experiment. Relative to control vehicles, overall survival of tumor-bearing mice treated ~30 days with FR at low or high dose was respectively 85% and 71%. Under these conditions, FR inhibited growth of MP41 tumors only at the high dose (Fig. 6A), but inhibited growth of MP46 tumors at either dose (Fig. 6A). FR had no effect on growth of BRAF(V600E)-driven OCM-1A tumors (Fig. 6A), demonstrating that FR did not affect processes such as tumor vascularization or perfusion that generically support tumor growth. Thus, FR specifically targeted both class 1 and 2 UM tumor xenografts, but only those driven specifically by oncogenic Gq/11.

To determine the extent to which FR delivered systemically at a therapeutically effective dose inhibited oncogenic G protein signaling in vivo, we compared the effects of FR on Erk activity, one of several downstream pathways activated by oncogenic Gq or G11 (Fig. 1A), Erk phosphorylation in MP41 cells was reduced in vitro by 85% (Figs. 6B and S4). In contrast, FR administered in vivo at a therapeutically effective dose (0.3 mg/kg on alternate days) for 26 days reduced Erk phosphorylation in G11-driven MP41 tumors in vivo by only ~30% (Figs. 6B and S4). Thus, modulation rather than complete suppression of oncogenic G protein signaling in MP41 tumor xenografts was sufficient for FR to have therapeutic effect.

Lastly, we investigated whether FR caused durable arrest or regression of class 1 UM tumor xenografts. In this experiment, mice were implanted with MP41 cells and allowed to form tumors. MP46 tumors were not studied because their slow growth made it difficult to determine whether FR potentially caused durable arrest or regression. We arrested MP41 tumor growth by treating animals with FR at an effective dose (0.3 mg/kg s.c. on alternate days) for 26 days reduced Erk phosphorylation in G11-driven MP41 tumors in vivo by only ~30% (Figs. 6B and S4). In contrast, FR appeared to cause reversible rather than durable arrest or regression of MP41 tumors. Taken together, these results indicated the existence of a therapeutic window in which FR effectively targeted class 1 or class 2 UM tumor xenografts, while preserving sufficient function of Gq/11 in host tissues required to support essential physiological processes and viability.

Discussion
Targeting UM tumors with FR

Here we have provided several new lines of evidence that support the therapeutic potential of Gq/11 inhibitors such as FR for UM. Our findings indicate that the vast majority of primary and metastatic UM tumors potentially can be targeted by FR. We found that all constitutively active mutant forms of Gq/11 found in UM, which collectively drive tumorigenesis in >90% of patients, are sensitive to FR. Moreover, FR can target tumors. As a reference for in vivo studies, we first determined the maximal extent that Erk phosphorylation in MP41 cells was inhibited by FR in vitro. At concentration of FR (1 μM) that maximally suppressed reporter activity driven by oncogenic Gq or G11 (Fig. 1A), Erk phosphorylation in MP41 cells was reduced in vitro by 85% (Figs. 6B and S4).

| Variable                                      | CTL mean | TX mean | Diff ±SEM       | p value |
|-----------------------------------------------|----------|---------|-----------------|---------|
| Walking initiation (s)                        | 2.54     | 1.98    | 0.96 ± 1.57     | 0.3537  |
| Ledge (/60s)                                  | 48.92    | 43.35   | 5.57 ± 1.67     | 0.4315  |
| Platform (/60s)                                | 22.97    | 25.74   | 2.77 ± 2.15     | 0.7294  |
| Pole (/120s)                                  | 25.92    | 22.16   | 3.76 ± 2.65     | 0.5844  |
| Turn (/60s)                                   | 23.73    | 15.97   | −7.76 ± 5.52    | 0.5844  |
| 60 degree screen (/60s)                       | 11.43    | 10.68   | 0.75 ± 0.45     | 0.634   |
| 90 degree screen (/60s)                       | 16.98    | 16.49   | 0.49 ± 0.36     | 0.9049  |
| Inverted screen (/60s)                        | 49.92    | 41.10   | 8.82 ± 6.64     | 0.1437  |
| Fine movements (/1h)                          | 1758     | 1549.00 | 209.1 ± 115.5   | 0.0861  |
| Total ambulations (/1h)                       | 756.5    | 674.3   | −82.20 ± 64.16  | 0.2155  |
| Time rearing (/hr)                            | 688.5    | 583.3   | 105.1 ± 81.22   | 0.2111  |
| Center distance (mm/h)                        | 4288     | 3941    | 347.4 ± 506.5   | 0.5010  |
| Center time (/1h)                             | 478.3    | 557.6   | −79.30 ± 112.8  | 0.4905  |
| Center entries (/1h)                          | 263.9    | 236.1   | 27.81 ± 27.70   | 0.3281  |
| Total rest time (s/1 h)                       | 1504     | 1635    | −130.6 ± 118.7  | 0.2647  |
| Total ambulations over time                   | 4539     | 4046    | 493.2 ± 385.0   | 0.2733  |
| Rearing frequency over time                   | 116.8    | 123.5   | 6.671 ± 14.51   | 0.8508  |
Gq/11-driven UM tumor cells from patient biopsies of primary ocular tumors with low or high metastatic potential and liver metastases. And lastly, although FR does not discriminate between wild-type and oncogenic Gq/11, and host Gq/11 activity is physiologically essential, we were able to identify a therapeutic window in which FR strongly inhibited tumor growth without causing large negative effects on viability or physiology. UM tumors therefore appear to be particularly vulnerable to FR.

The vulnerability of UM tumors to FR may occur because the signaling networks downstream of oncogenic Gq/11 must be maintained above a high threshold. In support of this hypothesis, we found that a therapeutically effective dose of FR strongly inhibits UM tumor growth but only modestly reduces oncogenic G11-driven Erk phosphorylation in UM tumors. Because nearly complete inhibition of Erk phosphorylation by FR was not necessary to inhibit UM tumor growth, Erk likely cooperates with other signaling pathways downstream of oncogenic Gq/11 to sustain UM tumors. Indeed, Gq/11 and MEK inhibitors recently have been shown to synergize and regress UM tumor xenografts, suggesting that synergism might improve efficacy or safety of Gq/11 inhibitors such as FR as therapeutic agents for UM.

**Therapeutic implications**

Our results suggest several ways that Gq/11 inhibitors such as FR could be explored therapeutically to treat UM patients, either alone or in conjunction with current standards of care. Primary UM tumors usually are treated by external beam or plaque irradiation of the eye to arrest growth, induce tumor cell death, and preserve vision, although these interventions on their own do not improve prognosis for
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metastasis (13). Once metastatic UM occurs in the liver, therapy involves mechanical isolation of the hepatic vasculature by embolization and percutaneous infusion of high-dose chemotherapeutics (43) or radiotherapy (44) above what would be tolerated systemically (45). Thus, targeting delivery of FR to primary UM tumors by intravitreal injection, or hepatic metastases by chemoembolization, could be considered. Targeted delivery might be required to reduce the impact of FR on viability, blood pressure, or other dose-limiting physiological processes, or allow FR to be delivered at higher doses in an effort to promote durable arrest or tumor regression. If FR proves to have limited efficacy in monotherapy, it could be combined with standard-of-care interventions to treat primary and metastatic UM. Intravitreal injection of FR also might interfere with progression of very early-stage lesions to class 2 tumors that have high metastatic potential by promoting melanocytic differentiation, as we have shown for class 1 UM tumor cells ex vivo. Similarly, because intraocular nevi are relatively common, express constitutively active Gq/11, and can progress to early stage melanomas (46), intravitreal injection of FR might interfere with this process and reduce the incidence of UM.

UM is only one of several diseases that potentially could benefit from targeted treatment with FR. Constitutively active mutant forms of Gq/11 occur in ~5% of all cancers, including melanocytic neoplasms of the central nervous system (4), mucosal melanoma (5), choroidal hemangiomas (8, 11), and hepatic small vessel neoplasms (6), as well as in Sturge–Weber syndrome (7), certain forms of hypercalcemia and hypocalcemia (9), and autosomal dominant hypoparathyroidism (10). Treatment regimens for all of these conditions potentially could be improved by novel therapeutic approaches employing FR.

Experimental procedures

Reagents

FR900359 (FR) was purified from Ardisia crenata according to published methods (16). SRE.L assays used a firefly luciferase reporter driven by the Gq/11-dependent SRE promoter and were performed as described (47). Firefly luciferase activity was normalized to Renilla luciferase expressed from a constitutive promoter.

Cell culture assays

Cells were cultured at 37°C in 5% CO2. Human 92.1 (RRID:CVCL_8607), Mel270 (RRID:CVCL_C302), and OCM-1A (RRID:CVCL_6934) UM cells were derived by and the generous gifts of Drs. Martine Jager (Laboratory of Ophthalmology, Leiden University), Bruce Ksander (Schepens Eye Institute, Massachusetts Eye and Ear Infirmary), and June Kan-Mitchell (Biological Sciences, University of Texas at El Paso), respectively. The human PDX cell lines MP41 (ATCC Cat# CRL-3297, RRID:CVCL_4D12) and MP46 (ATCC Cat# CRL-3298, RRID:CVCL_4D13) cell lines were purchased from ATCC (Manassas, VA). Cell lines were grown in RPMI 1640 medium (Life Technologies, Carlsbad, CA) supplemented with FBS and antibiotics. Cell viability was measured using a water-soluble tetrazolium salt, WTS-8 (Bimake, Houston, TX), following the manufacturer’s protocol. Flow cytometry for analysis of cell proliferation and apoptosis was performed at the Siteman Cancer Center Flow Cytometry Core on a FACScan analyzer (BD Biosciences, San Diego CA, USA) using a standard propidium iodide staining protocol as described previously (48).

Patients and biopsy collection

Human UM biopsy or enucleation samples were obtained with patient written informed consent and with approval of the institutional review board of Washington University in St Louis. Fine-needle aspiration biopsies of primary UM tumors and ultrasound-guided core biopsies of UM liver metastases were performed as part of standard of care, which also collected biopsies for cytological evaluation of tumor cells and molecular classification (Castle Biosciences). For enucleation samples, after excision, eyes were transilluminated to localize the tumor mass and then opened opposite the tumor. Vitreous was carefully removed and tumor samples were collected through the retina from the tumor apex. All samples were collected directly into growth medium in the operating room before being transported to the laboratory. Primary uveal melanoma cells were divided equally into two wells of fibronectin-covered 6-well tissue culture plates and grown in 5% CO2 in MDMF medium, which consists of HAM’s F12 (Lonza, Walkersville MD, USA) supplemented with 1 mg/ml BSA (Sigma-Aldrich, St Louis MO, USA), 2 mM L-glutamine (Lonza), 1X SITE (Sigma-Aldrich), 1x B27 ( Gibco, Carlsbad CA, USA), 20 ng/ml bFGF (PeproTech Inc, Rocky Hill NJ, USA), and 50 μg/ml Gentamicin (Sigma-Aldrich) (49). Cells were allowed to attach to the substrate overnight before fresh medium was added to each well containing either 100 nM FR or an equal volume of vehicle (DMSO).

Bulk RNAseq

MP41 and MP46 cells were treated with 100 nM FR or vehicle (DMSO) in RPMI growth medium and collected after 3 days of treatment. FNAB samples were treated with 100 nM FR or vehicle in MDMF medium and collected after 7 days of treatment. RNA was isolated using the RNeasy Mini Kit (Qiagen) following the manufacturer’s protocol and including the optional DNase I treatment step. RNA quality was assessed on a Bioanalyzer 2100 (Agilent Technologies, Santa Clara, CA, USA). mRNA was extracted from total RNA using a Dynal mRNA Direct kit, fragmented, and reverse transcribed to double-stranded cDNA with random primers before addition of adapters for library preparation. Library preparation and HiSeq2500 sequencing were performed by the Washington University Genome Technology Access Center (gtac.wustl.edu). FastQ files were aligned to the transcriptome and the whole genome with STAR. Biologic replicates were simultaneously analyzed by edgeR and Sailfish analyses of gene-level/exon-level features. Unexpressed genes and exons were removed from the analyses. Unsupervised principal component analysis was generated in Bioconductor using edgeR.
Direct comparison of FR response in MP41 versus MP46 cells was used to identify MP41-specific genes, and this list was used as signature gene sets for Gene Set Enrichment Analysis (GSEA) (50).

**Single-cell RNAseq**

Single-cell RNAseq (scRNAseq) was performed using the 10X Genomics platform. Single-cell suspensions were counted using a hemocytometer and adjusted to 1000 cells/μl. Sample processing through 10X Genomics and NextSeq 500 (Illumina) sequencing was performed by the Washington University Genome Technology Access Center (gtac.wustl.edu). Raw base call files were analyzed using Cell Ranger v.3.0.2. The filtered counts data from all six samples were combined in Partek Flow (www.partek.com). Samples were filtered on total reads, detected features, and mitochondrial content per cell to remove cell doublets and apoptotic debris. Unexpressed features were removed, and samples were normalized and log transformed.

PCA was used to reduce dimensionality, and the first 20 principal components were further analyzed by graph-based (Louvain) clustering and t-SNE. The resulting 24 clusters were visualized in a two-dimensional t-SNE representation and were annotated to known biological cell types using canonical marker genes. The following cell types were annotated (selected markers are listed): Lymphocytes (CD3D, CD3G, IL7R); macrophages (C1QA, C1QB, C1QC); dendritic cells (CD1C and lack of expression of C1QA, C1QB and C1QC); retinal cells (SYNGR1, NPTX1); photoreceptors (GNAT1, GNGT1); RPE(PAX6, KRT8, KRT18); melanoma (DCT, TYRP1, PMEL).

**Mice**

All experiments were performed using the NOD scid gamma (NOD.Cg-Prkdc<sup>scid</sup> I2<sup>tm1Wjl/Szl</sup>; NSG) mice purchased from The Jackson Laboratory (#005557). All animal experiments were performed under protocols approved by the Animal Studies Committees of Washington University in St Louis School of Medicine and Drexel University College of Medicine. FR was prepared from a 15 mM stock solution in DMSO at doses of 0.1, 0.3, 1.0, or 3.0 mg/kg and dextrose at a final volume of 200 μl. FR was administered subcutaneously every other day. Animals were randomly grouped (n = 10/group) for placebo (1% DMSO) or FR treatments. MP41, MP46, and OCM1A cells were inoculated subcutaneously into the flanks of 5-week-old male NSG mice. Cells were prepared at a concentration of 2 x 10<sup>6</sup> cells/100 μl in a solution of iced cold PBS/Matrigel Matrix (50/50 v/v) (Fisher Scientific # CB40234 A). Primary tumors were allowed to grow to ~30 mm<sup>2</sup> basal area based on measurements taken with calipers and using the formula (L x 2W)/2 where L and W are the longest and the shortest basal diameters of the tumor, respectively, before beginning treatment with FR. Tumors reaching a maximum length of 2 cm were defined as censored end points. At censored end points or at the end of each time course, mice were euthanized, and the tumors were removed using sterile surgical technique. Excised tumors were weighed to determine end-point mass and snap-frozen in liquid nitrogen for later use.

**Immunoblotting**

For tumor samples, approximately 100 mg of tumor was homogenized on ice in 1X Cell Lysis Buffer (Cell Signaling Technology, catalog# 9803) supplemented with 1 mM PMSF. For cultured cells, cells were grown in 10 cm dishes and treated with DMSO or 1 μM FR for 18 h, and then lysed in 1X Cell Lysis Buffer. Lysates were sonicated on ice for 2 min (30 s on, 30 s off, 60% A), rotated end-over-end for 30 min, and cleared by centrifugation at 16,000 x g for 15 min. Protein concentration was determined using Bio-Rad Protein Assay Dye Reagent (Bio-Rad, catalog# 5000006). In total, 15 μg of tumor protein was resolved on 12% SDS-PAGE gels and transferred to Immobilon-FL PVDF membrane (Millipore, catalog# IPFL0010). Membranes were blocked with 5% (w/v) BSA in TBST [20 mM Tris pH 7.6, 137 mM NaCl, 0.1% v/v Tween 20] and incubated with primary antibodies [Phospho-p44/42 MAPK (Erk1/2) Cell Signaling Technology, catalog# 4370S lot#24 and p44/42 MAPK (Erk1/2) Cell Signaling Technology catalog# 9107S lot#10]. Membranes were washed with TBST at least three times and incubated with IRDye 680–coupled goat anti-rabbit (LI-COR, catalog# 926–68,071, lot# C90618–09) and IRDye 800 goat anti-mouse (LI-COR, catalog# 926–32,210, lot# C91210–09 antibodies (LI-COR Biosciences). After incubation, membranes were washed at least three times with TBST, and signals were detected using Odyssey model 9120 imaging system (LI-COR Biosciences).

**Mouse radiotelemetry**

Blood pressure and heart rate responses to chronic administration of vehicle or FR900359 were monitored by radiotelemetry in conscious mice by following previously described procedures (24, 51). Briefly, adult NSG male mice were implanted with a radiotelemetry pressure-sensing catheter in the right carotid artery under isoflurane (3% mixed with 95% oxygen) anesthesia. The body of the transmitter (HD-X10, Data Science International (DSI), St Paul, MN) was tunneled into a subcutaneous pouch on the left flank of the animal. The mice were allowed to recover from the surgery for 1 week after which baseline blood pressure and heart rate were recorded for 24 h, followed by alternating days of subcutaneous administration of vehicle or FR. After the 24-h baseline recordings, the mice were randomly assigned to two groups: one group receiving FR (0.3 mg/kg in dextrose, s.c.) and the other group receiving equal amounts of vehicle (2 μl DMSO in 200 μl of 5% dextrose, s.c.). Daily recordings were conducted from 1 to 5 PM for 14 days, and FR or vehicle was administered every other day at 3 PM. Continuous 24-h BP recordings were conducted during the first injection and seventh injection of FR. Data was acquired and analyzed using DSI Ponemah software version 6.5. Averages of continuous systolic blood pressure (SBP), diastolic blood pressure (DBP), mean blood
pressure (MBP), and heart rate (HR) recordings were plotted in 1 h intervals over 24 h.

**Mouse behavioral tests**

Mice were moved to the animal facility and allowed to acclimate to the new environment for 1 week prior to behavioral testing. All behavioral testing was conducted during the light cycle, by a female experimenter blinded to experimental group. All equipment was cleaned with 2% chlorhexidine diacetate or 70% ethanol between animals.

**Open-field activity/exploratory behavior**

General activity levels and exploratory behavior were quantified over a 60-min period in an open field (47.6 cm L x 25.4 cm W x 20.6 cm H) constructed of Plexiglas and surrounded by computerized photobeam instrumentation (Kinder Scientific, LLC, Poway, CA). General activity variables (total ambulations, rearings, time at rest) along with measures of emotionality, including time spent, distance traveled, and entries made into the central zone, were analyzed.

**Sensorimotor battery**

Walking initiation, ledge, platform, pole, and inclined and inverted screen tests were performed as previously described (52). Time in each task was manually recorded. The average of two trials was used for analyses. Test duration was 60 s, except for the pole test, which was extended to 120 s. For walking initiation, time for an animal to leave a 21 x 21 cm$^2$ square on the floor was recorded. For ledge and platform tests, the time the animal was able to balance on an acrylic ledge (0.75 cm wide and 30 cm high), and on a wooden platform (1.0 cm thick, 3.0 cm in diameter and elevated 47 cm), was recorded, respectively. The pole test was used to evaluate fine motor coordination by quantifying time to turn 180° and climb down a vertical pole. The screen tests assessed a combination of coordination and strength by quantifying time to climb up or hang onto a mesh wire grid measuring 16 squares per 10 cm, elevated 47 cm and inclined (60° or 90°) or inverted.

**Statistical analyses**

All statistical analyses were performed using IBM SPSS software (v.24) and Graphpad Prism 8. For cell line data, all experiments were performed in triplicate at least three times on different days. Means and standard errors were computed from all cell line data and t-tests were used to determine significance. For significance analysis of PCA data, correlation of each principal component with FR treatment was calculated using paired t-tests with Holm–Šidák correction for multiple comparisons. For behavioral studies, all data were screened for fit of distributions with assumptions underlying univariate analyses, which included the Shapiro–Wilk test and q-q plot investigations for normality. Means and standard errors were computed for each measure. Repeated measures analysis of variance (ANOVA) and independent samples t-tests were used to analyze behavioral data. Statistical results were confirmed with two-tailed nonparametric testing, when available, for any data sets with violations of the univariate assumptions. Probability value for all analyses was $p < 0.05$, unless otherwise stated.

**Data availability**

All RNAseq and single-cell RNAseq data have been deposited on the NCBI GEO server with the accession #GSE165552. All other data associated with this study are presented in the main text or supplementary materials.

**Supporting information**—This article contains supporting information.

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**Author contributions**—M. D. O. performed the experiments, analyzed data, wrote the first draft, and revised the final draft; C. M. M., K. K. M., J. C., and A. J. D. performed the experiments; L. H. -A., K. D. P., and P. K. R. provided the tumor biopsy material; K. N. W. provided the instruction for mouse xenograft experiments; C. M. Y. performed and analyzed the mouse behavior experiments; P. O. -O. obtained funding, and supervised and analyzed blood pressure and heart rate experiments; J. A. C. obtained funding, analyzed data, and wrote the initial draft; K. J. B. conceived and directed the study, obtained funding, analyzed data, wrote the initial draft, and revised the final draft.

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**Conflict of interest**—K. J. B. and M. D. O. are listed as coinventors on a provisional patent application on Targeted Pharmacological Therapeutics In Uveal Melanoma that is owned by Washington University in St. Louis. All other authors declare no competing interests.

**Abbreviations**—The abbreviations used are: ANOVA, analysis of variance; DBP, diastolic blood pressure; HR, heart rate; MBP, mean blood pressure; PCR2, polycomb repressive complex 2; PDX, patient-derived xenograft; UM, uveal melanoma; SBP, systolic blood pressure.

**References**

1. Van Raamsdonk, C. D., Bezrookove, V., Green, G., Bauer, J., Gaugler, L., O’Brien, J. M., Simpson, E. M., Barsh, G. S., and Bastian, B. C. (2009)
Effects of oncogenic Gqq and G11 inhibition by FR900359 in uveal melanoma. Mol. Cancer Res. 17, 963–973

Anna, S., Feng, X., Shridhar, N., Ervilimazz, F., Patt, J., Yang, J. H., Pfeil, E. M., Cervantes-Villagran, R. D., Inoue, A., Häberlein, F., Słodczyk, T., Reher, R., Kehraus, S., Monteleone, S., Schrage, R., et al. (2019) Direct targeting of Gq q and G11 oncogenes in cancer cells. Sci. Signal. 12, eaau5948

Feng, X., Arang, N., Rigiraccioli, D. C., Lee, J. S., Yeerna, H., Wang, Z., Lubrano, S., Kishore, A., Pachter, J. A., König, G. M., Maggolini, M., Kostenis, E., Schlaepfer, D. D., Tamayo, P., Chen, Q., et al. (2019) A platform of Synthetic lethal gene interaction networks reveals that the GNA14 oncogenic melanoma controls the Hippo pathway through FAK. Cancer Cell 35, 457–472.e5

Takasaki, J., Saito, T., Taniguchi, M., Kawasaki, T., Moritani, Y., Hayashi, K., and Kobori, M. (2004) A novel Galphα11-selective inhibitor. J. Biol. Chem. 279, 47438–47445

GNAQ, and GNA11 mutations occur in 9.5% of mucosal melanoma and are associated with poor prognosis. Eur. J. Cancer 65, 156–163

Joseph, N. M., Brunt, E. M., Marginean, C., Nalbantoglu, Ilk., Snover, D. C., Thung, S. N., Yeh, M. M., Umetu, S. E., Ferrell, D. L., and Gill, R. M. (2018) Frequent GNAQ and GNA14 mutations in hepatic small vessel neoplasm. Am. J. Surg. Pathol. 42, 1201–1207

Shirley, M. D., Tang, H., Gallione, C. J., Baugher, J. D., Frelin, L. P., Cohen, B., North, P. E., Marchuk, D. A., Comi, A. M., and Pevsner, J. (2013) Sturge-Weber syndrome and port-wine stains caused by somatic mutation in GNAQ. N. Engl. J. Med. 368, 1971–1979

Richards, C. A., Goss, J., Alomari, M., Alexandrescu, S., Robb, R., Smith, L. E., Hochman, M., Greene, A., and Bischoff, J. (2019) Association of somatic GNAQ mutation with Capillary Malformations in a case of GNA11 mutations occur in 9.5% of uveal melanoma.

Cranston, T., Rust, N., Hobbs, M. R., Heath, H., and Thakker, R. V. (2013) Frequent GNAQ and GNA14 mutations in hepatic small vessel neoplasm. Am. J. Surg. Pathol. 37, 5320–5324

Küsters-Vandevelde, H. V. N., Klassen, A., Küsters, B., Groenen, P. J. T. van, Angen-Van Gunsviren, I. A. C. H., Van Dijk, M. R. C. F., Reifenberger, G., Wesseling, P., and Blokx, W. A. M. (2010) Activating mutations of the GNAQ gene: a frequent event in primary melanocytic neoplasms of the central nervous system. Acta Neuropathol. 119, 317–323

Sheng, X., Kong, Y., Li, Y., Zhang, Q., Si, L., Cui, C., Chi, Z., Tang, B., Mao, L., Liu, B., Wang, X., Yan, X., Li, S., Dai, J., and Guo, J. (2016) GNAQ and GNA11 mutations occur in 9.5% of uveal melanoma.

GNAQ and GNA11 mutations occur in 9.5% of mucosal melanoma and are associated with poor prognosis. Eur. J. Cancer 65, 156–163

Joseph, N. M., Brunt, E. M., Marginean, C., Nalbantoglu, Ilk., Snover, D. C., Thung, S. N., Yeh, M. M., Umetu, S. E., Ferrell, D. L., and Gill, R. M. (2018) Frequent GNAQ and GNA14 mutations in hepatic small vessel neoplasm. Am. J. Surg. Pathol. 42, 1201–1207

Shirley, M. D., Tang, H., Gallione, C. J., Baugher, J. D., Frelin, L. P., Cohen, B., North, P. E., Marchuk, D. A., Comi, A. M., and Pevsner, J. (2013) Sturge-Weber syndrome and port-wine stains caused by somatic mutation in GNAQ. N. Engl. J. Med. 368, 1971–1979

Richards, C. A., Goss, J., Alomari, M., Alexandrescu, S., Robb, R., Smith, L. E., Hochman, M., Greene, A., and Bischoff, J. (2019) Association of somatic GNAQ mutation with Capillary Malformations in a case of choroidal hemangioma. JAMA Ophthalmol. 137, 91–95

Neha, M. S., Hannan, F. M., Hose, S. A., Babinsky, Y. N., Head, R. A., Cranston, T., Rush, N., Hobbs, M. R., Heath, H., and Thakker, R. V. (2013) Mutations affecting G-protein subunit α1 in herpetic keratitis and hypercalcaemia. N. Engl. J. Med. 368, 2476–2486

Li, D., Opas, E. E., Tuluc, F., Metzger, D. L., Hou, C., Hakonarson, H., and Levine, M. A. (2014) Autosomal dominant hypothyroidism caused by germline mutation in GNA11: Phenotypic and molecular characterisation. J. Clin. Endocrinol. Metab. 99, E1774–E1783

Francis, J. H., Milman, T., Grieskanlous, H., Albert, D., Solberg, R., Levituy, G., Coupland, S., Catalonatti, F., Rabady, D., Kandoth, C., Busam, K., and Abramson, D. (2019) GNAQ mutations in Diffuse and Solitary choroidal hemangiomas. Ophthalmology 126, 759–763

Krantz, B. A., Dave, N., Komatsubara, K. M., Marr, B. P., and Carvajal, R. D. (2017) Uveal melanoma: epidemiology, etiology, and treatment of primary disease. Clin. Ophthalmol. 11, 279–289

Carvajal, R. D., Schwartz, G. K., Tezel, T., Marr, B. F., Francis, J. H., and Nathan, P. D. (2017) Metastatic disease from uveal melanoma: treatment options and future prospects. Br. J. Ophthalmol. 101, 38–44

Kostenis, E., Pfeil, E. M., and Annala, S. (2020) Heterotrimetric Gq proteins as therapeutic targets? J. Biol. Chem. 295, 5206–5215

Fujio, C., Koda, S., Morimoto, Y,, and Biemann, K. (1988) Structure of FR900359, a cyclic depsipeptide from Ardisia crenata Sims. J. Org. Chem. 53, 2820–2825

Schrage, R., Schmitz, A. L., Gaffal, E., Annala, S., Kehraus, S., Wenzel, D., Büllesbach, K. M., Bald, T., Inoue, A., Shinjo, Y., Glandrin, S., Shridhar, N., Heise, M., Grundmann, M., Merten, N., et al. (2015) The experimental power of FR900359 to study Gq-regulated biological processes. Nat. Commun. 6, 10156

Onnen, M. D., Makepeace, C. M., Kaltenbronn, K. M., Kanai, S. M., Todd, T. D., Wang, S., Broekelmann, T. J., Rao, P. K., Cooper, J. A., and Blumer, K. J. (2018) Targeting nucleotide exchange to inhibit constitutively active G protein a subunits in cancer cells. Sci. Signal. 11, eaao6852

Lapadula, D., Farias, E., Randolph, C. E., Perwin, T. J., McGrath, D., Charpentier, T. H., Zhang, L., Wu, S., Terao, M., Sato, T., Tall, G. G., Zhou, N., Wedegaertner, P. B., Aplin, A. E., Aguirre-Ghiso, J., et al. (2019)
FR900359 targets primary and metastatic uveal melanoma