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***Basic Study***

**Free fatty acids receptors 2 and 3 control cell proliferation by regulating cellular glucose uptake**

Al Mahri S *et al.* Role of FFARs in cell proliferation

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**Abstract**

BACKGROUND

Colorectal cancer (CRC) is a worldwide problem, which has been associated with changes in diet and lifestyle pattern. As a result of colonic fermentation of dietary fibres, short chain free fatty acids are generated which activate free fatty acid receptors (FFAR) 2 and 3. *FFAR2* and *FFAR3* genes are abundantly expressed in colonic epithelium and play an important role in the metabolic homeostasis of colonic epithelial cells. Earlier studies point to the involvement of FFAR2 in colorectal carcinogenesis.

AIM

To understand the role of short chain FFARs in CRC.

METHODS

Transcriptome analysis console software was used to analyse microarray data from CRC patients and cell lines. We employed short-hairpin RNA mediated down regulation of *FFAR2* and *FFAR3* genes, which was validated using quantitative real time polymerase chain reaction. Assays for glucose uptake and cyclic adenosine monophosphate (cAMP) generation was done along with immunofluorescence studies to study the effects of FFAR2/FFAR3 knockdown. For measuring cell proliferation, we employed real time electrical impedance-based assay available from xCELLigence.

RESULTS

Microarray data analysis of CRC patient samples showed a significant down regulation of *FFAR2* gene expression. This prompted us to study the FFAR2 in CRC. Since, FFAR3 shares significant structural and functional homology with FFAR2, we knocked down both these receptors in CRC cell line HCT 116. These modified cell lines exhibited higher proliferation rate and were found to have increased glucose uptake as well as increased level of glucose transporter 1. Since, FFAR2 and FFAR3 signal through G protein subunit (Gαi), knockdown of these receptors was associated with increased cAMP. Inhibition of protein kinase A (PKA) did not alter the growth and proliferation of these cells indicating a mechanism independent of cAMP/PKA pathway.

CONCLUSION

Our results suggest role of *FFAR2*/*FFAR3* genes in increased proliferation of colon cancer cells via enhanced glucose uptake and exclude the role of PKA mediated cAMP signalling. Alternate pathways could be involved that would ultimately result in increased cell proliferation as a result of down regulated *FFAR2*/*FFAR3* genes. This study paves the way to understand the mechanism of action of short chain FFARs in CRC.

**Key words:** Cell proliferation; Glucose transporter 1; Colorectal cancer; Free fatty acids receptor 2; Free fatty acids receptor 3; Glucose uptake

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**Core tip:** Free fatty acid receptors (FFAR) have been reported to be associated with colorectal cancer (CRC). In this report, we studied short chain FFAR2 and FFAR3 and have provided preliminary evidence about the possible involvement of increased glucose uptake in FFAR2 and FFAR3 knockdown clones of CRC cell line. We generated double knock down for *FFAR2* and *FFAR3* genes in a CRC cell line (HCT116) and studied possible mechanisms of increased cell proliferation in these cells.

**INTRODUCTION**

Colorectal cancer (CRC) is a disease that is associated with the dietary patterns, metabolism and inflammation[1]. Type of diet significantly modifies the risk for development of CRC[2]. There is evidence linking high carbohydrate-low fibre diet to increased risk for CRC[3-6]. The site of this cancer is the location for processing of food aided by the gut microbiota. The effect of diet on CRC has been studied with different perspectives of associated factors. Metabolism of nutrients, role of gut microbiota and familial factors are being studied to better understand the causal factors in diet related initiation and progression of CRC[7]. Type of food intake, its digestion and metabolism are an upcoming area of research with potential to develop preventive and therapeutic strategies. Characterization of gut microbiota with newer technologies is allowing the possibility of customized probiotic treatment for the prevention of CRC[7,8]. Short Chain Fatty Acids (SCFAs) are produced in the distal gut by bacterial fermentation of macro-fibrous material that escapes digestion in the upper gastrointestinal tract and enters the colon[9,10]. SCFAs such as butyrate and propionate exert anticancer effect on colon as they have been shown to induce differentiation, growth arrest and apoptosis, mainly due to their intracellular actions, through inhibition of histone deacetylase[11,12]. This suggests that SCFAs produced in the gut could have protective properties against development of CRC. SCFAs are cognate ligands for a group of G-protein coupled receptors, free fatty acid receptor (FFAR) 2 and FFAR3 also known as GPR43 and GPR41, respectively[13,14]. *FFAR2* and *FFAR3* genes are abundantly expressed in human colon[15]. FFAR2 recognizes all three major SCFAs, but the affinities for FFAR3 are in the order of propionate > butyrate > acetate[16]. Activated FFAR2 initiate signalling through the Gαi pathway to suppress cyclic adenosine monophosphate (cAMP) levels, and through the Gαq pathway to enhance calcium mobilization[17]. Negative impact of FFAR2 on cAMP levels results in inhibition of protein kinase A (PKA) (primary downstream effector of cAMP) and its substrate, cAMP response element binding protein. Together, this leads to reduced expression of histone deacetylase[18]. Several studies on FFARs suggested their involvement in the onset and progression of colon cancer[15,19]. It has been suggested that FFAR2 plays a role in the cancer cell metabolism and growth. Loss of FFAR2 was observed using immunohistochemistry in malignant colon adenocarcinoma tissues by 80% compared to normal human colon mucosa tissue[15]. Transforming activity of loss of FFAR2 and its oncogenic potential was validated in fibroblasts[20]. However, there is no conclusive study to suggest the role of FFAR2 as well as FFAR3 in cancer cell growth and metabolism[21].

The aim of this study is to contribute towards our understanding the connection between gut microbiota and CRC. In this report, we focused on genes encoding for receptors for SCFAs in CRC patient samples. We analysed gene expression level of FFAR2 and FFAR3 in our patient cohort. Using microarray analysis of matched tumour-normal tissues, we provide evidence that gene expression of FFAR2 is significantly reduced in CRC patients. While some patients had reduced *FFAR3* gene expression, we did not observe significant differences in gene expression levels in our patient cohort. To further characterize the mechanistic effect of reduced level of expression of these receptors, we generated knockdown (KD) clones of *FFAR2* and *FFAR3* genes in HCT116 CRC cell line. While FFAR2 KD exhibited increased cell proliferation, it was further increased with subsequent KD of *FFAR3* gene. These KD clones were found to exhibit a significant increase in their glucose uptake as well as increased expression of glucose transporter 1 (GLUT1). Simultaneously, increased levels of cAMP were observed in these cells. These results provide evidence to suggest the role of FFAR2 as well as FFAR3 in progression of CRC via a previously unknown mechanism of increased glucose uptake.

**MATERIALS AND METHODS**

***Patient samples and microarray analysis***

CRC patient samples were analysed for gene expression changes using microarray analyses of the dataset we previously reported (GEO accession number: GSE 50421)[22]. For the analysis of differential gene expression between tumour and normal, we used Transcriptome Analysis Console software available from Affymetrix (Thermofisher Scientific, United States). Sample IDs and respective expression values for *FFAR2* and *FFAR3* genes in tumour and normal samples are given in Supplementary Table 1. Gene expression values were generated using Transcriptome Analysis Console and further analysed using GraphPad prism 7 software. Paired t-test was used to compare matched tumor-normal samples and detailed results are provided in Supplementary Table 2.

***Ethics, consent and permissions***

The study is approved by the Institutional Review Board at King Abdullah International Medical Research Center. Procedural and ethical consent forms were generated and approved by the Institutional Review Board office at at King Abdullah International Medical Research Center. Each patient prior to sample collection signed the procedural and ethical consent forms. The permissions and consents were obtained according to the journal's guidelines and standards.

***Cell line and culture conditions***

HCT116 cell line (ATCC® CCL-247™) was obtained from American Type Culture Collection (Manassas, VA, United States). Cells were cultured in 5%CO2 at 37 °C. Cells were grown in advanced Dulbecco’s modified Eagle’s medium (DMEM) containing 10% FBS, 100 IU/mL of penicillin, 100 μg/mL of streptomycin and 2 mmol/L L-glutamine (Gibco, Thermo Fisher Scientific, Waltham, MA, United States).

***Generation of stable FFAR2 KD HCT116 cell line***

FFAR2 short-hairpin RNA (shRNA), targeting 4 different sequences of the *FFAR2* gene, and a control scrambled shRNA plasmid were obtained from Origene Technologies, Inc. 1 × 105 HCT116 cells were plated in a 24-well plate and transfected with 0.5 μg of plasmid DNA using Lipofectamine3000 with a ratio of 1:2 (DNA:Lipo3000) according to the manufacturer’s protocol. Transfected HCT116 cells were selected in 0.8 μg/mL of puromycin (As per data available at <http://cell-lines.toku-e.com/Cell-Lines_1422.html> for 4 wk then isolated colonies were maintained in 0.8 μg/mL of puromycin antibiotic. Colonies were screened for FFAR2 KD using quantitative real time polymerase chain reaction (qRT-PCR) and one colony with most efficient KD was selected for subsequent experiments. One colony from scrambled shRNA construct was also selected to serve as control in all experiments.

***Generation of FFAR2 and FFAR3 double KD cells***

We used pre-packaged Lentiviral particles harbouring 3 different shRNA sequences against the *FFAR3* gene and a control-scrambled shRNA that were obtained from GeneTarget, Inc. To generate FFAR2/FFAR3 double KD cells. Earlier selected colony with most efficient KD of FFAR2 were transduced with the FFAR3 Lentiviral particles with a multiplicity of induction (MOI) = 5. Cells were selected in 5 μg/mL of blasticidin antibiotic (Dose was determined for HCT 116 cells using kill curve for blasticidin) for 4 wk then colonies were maintained in 5 μg/mL of blasticidin and 0.8 μg/mL of puromycin antibiotic. Colonies were screened for FFAR2 and FFAR3 KD using qRT-PCR and one colony with most efficient KD was selected for subsequent experiments.

***RNA extraction, cDNA synthesis and real-time PCR***

Total RNA from five patients’ tumour-normal paired samples was extracted using the QIAGEN RNeasy Mini Kit (Qiagen; Cat# 74104). 2 μg of RNA was used for cDNA synthesis using High-Capacity cDNA Reverse Transcription Kit (Applied Biosystems, Thermo Fisher Scientific). qRT-PCR was performed by using ABI 7900HT PCR system (Applied Biosystems, Foster City, CA, United States). GAPDH was used as housekeeping control. Sequence of primers used are GAPDH-F: 5’ ACAACTTTGGTATCGTGGAAGG 3’, GAPDH-R: 5’ GCCATCACGCCACAGTTTC 3’; FFAR2-F: 5’ TGCTACGAGAACTTCACCGAT 3’; FFAR2-R: 5’ GGAGAGCATGATCCACACAAAAC 3’; FFAR3-F: 5’ GCGTGGAGGATCTACGTGAC 3’; FFAR3-R: 5’ TGTGAGTGTTCACTGGTCTTTC 3’. PowerUp SYBR Green Master Mix (Applied Biosystems) was used. All reactions performed in triplicate and the qRT-PCR data was analysed by using the RQ method (2-ΔΔCt) by SDS RQ manager and expression Suite version 1.1. Complete details with Ct values have been provided in Supplementary Tables 3 and 4.

***Glucose uptake assay***

One × 104 HCT116cells from control-scrambled, FFAR2 KD and FFAR 2/3 double KD clones were plated in each well of a 96 well plate. Next day, cells were washed twice with phosphate buffer saline and incubated in glucose free DMEM. Glucose uptake was determined using Glucose uptake assay kit (Cayman Chemicals, MI, United States) following manufacturers protocol. Briefly, cells were incubated with 100 μg/ml fluorescent 2-N-7-Nitrobenz-2-oxa-1, 3-diazol-4-yl-Amino-2-Deoxyglucose (2-NBDG) in glucose free medium for 1h. Cells were washed with assay buffer 3 times and analysed immediately. 2-NBDG taken up by cells was detected on a Tecan Infinite 200 pro fluorimeter (Tecan Group Ltd. Männedorf, Switzerland) with fluorescent filters designed to detect fluorescein (excitation/emission = 485/535 nm). Cells without 2-NBDG incubation were used as blank control.

***cAMP assay***

Three x 104 HCT116cells from control-scrambled, FFAR2 KD and FFAR2/FFAR3 KD were plated on a 48 well plate overnight. Next day, cells were incubated with 500 μmol/L IBMX (3-isobutyl-1-methylxanthine) for 1h. cAMP in each well was measured using cAMP screen Immunoassay System (Thermo-fisher scientific) by following the manufactures instructions. Briefly, after IBMX incubation, cells were lysed in 100 μL lysis buffer (provided with the kit) by incubating at 37 °C for 30 min. 50 μL of cell lysates were added to the pre-coated 96 well plate followed by addition of cAMP-alkaline phosphatase conjugate and cAMP antibodies to each well. A cAMP standard was also prepared ranging 0.002-2000 pmol of cAMP. Plate was incubated at room temp for 2 h with constant shaking. Each well was washed 6 times with wash buffer followed by addition of CSPD/Sapphire-II™ RTU substrate/enhancer solution and incubation for 30 min. Measurements were made using a single-mode luminometer (Molecular Devices). Standard curve was made using the reading from cAMP standards and cAMP was measured in each well with reference to the standard curve. The experiment was repeated twice with 8 technical repeats for each condition.

***Cell proliferation assay using XCELLigence***

Rate of cell proliferation was measured using xCELLigence real time cell analyser-dual purpose (RTCA-DP) available from ACEA biosciences (San Diego, United States). E-16 plates were used for monitoring cell adhesion and growth. This system works on the principle of electrical impedance. A unitless parameter termed Cell Index (CI) is used to measure the relative change in electrical impedance to represent cell status. CI is a relative and dimensionless value since it represents the impedance change divided by a background value. When there are no cells present in the medium, the sensor’s electronic property will not be affected and the impedance will be small. When there are more cells on the electrodes, the impedance will be larger. CI calculation is based on the following formula: CI = (Zi – Z0)/15 ς, where Zi is the impedance at an individual point of time during the experiment and Z0 is the impedance at the start of the experiment. CI is a self-calibrated value derived from the ratio of measured impedances. For cell proliferation experiments, we plated 40000 cells from scrambled control, FFAR2 KD and FFAR 2/FFAR3 KD HCT116 cells on E-16 plate in duplicate and cell proliferation was monitored for 60 h. The experiment was repeated 3 times and slope values of the growth curve were plotted using in-built software and later analysed on Graph pad prism. H89 (PKA inhibitor) was added at a concentration of 50 μmol/L after 24 h of plating and cell growth was monitored up to 60 h.

***Immunofluorescence for GLUT1 gene expression***

Fifty thousand HCT116cells from control-scrambled, FFAR2 KD and FFAR2/FFAR3 KD were grown on cover slips were washed with PBS and fixed with 4% formaldehyde. Cells were incubated with anti-GLUT1 antibody (Catalogue #PA1-46152, Thermo Fisher Scientific, United States) in PBS containing 5% serum and 0.1% Triton for 30min at 37 °C. Cells were washed four times with PBS and incubated with fluorescein isothiocyanate (FITC) conjugated antibodies for 30 min. Cells were washed four times and immediately imaged using EVOS FL Auto imaging system (ThermoFisher Scientific). Images were quantified for GLUT1 expression using metamorph image processing software (Molecular devices).

***PKA immunoblotting***

One million HCT116cells from control-scrambled, FFAR2 KD and FFAR2/FFAR3 KD cells were lysed with Immunoprecipitation buffer (Thermo Fisher Scientific) containing protease inhibitors. The scraped cells were kept on ice for 30 min and centrifuged at 10000 *g* for 5 min and the supernatant was collected. Protein concentrations were determined by Qubit protein assay kit (Thermo Fisher Scientific). Twenty μg of lysate was boiled at 95 °C for 5 min and loaded onto ready-made gel 4%–20% Mini-PROTEAN® TGX™ Precast Protein Gels (Bio-Rad), subjected to electrophoresis and transferred to PVDF membranes (Bio-Rad). The membranes were blocked using 5% BSA (MILLIPORE) in Tris buffer saline + Tween 20 (TBST) buffer for 1 h with shaking at room temperature. The following primary antibodies were used: PKA-PAS (Cell signalling) isotype anti-rabbit at recommended dilution of 1:1000 in 5% BSA with gentle shaking overnight at 4°C. The blots were incubated with the appropriate Horse Radish Peroxidase-conjugated secondary antibody (Bio-Rad), and the signals were detected with Chemiluminiscent HRP Substrate (Bio-Rad). Images were captured and analysed using Chemidoc gel documentation system (Bio-Rad).

***Statistical analysis***

Data were presented as the mean ± SD and analysed for statistical signiﬁcance using two tailed students *t* test available in the GraphPad prism 7 software version 7.03. Each bar represents an average of at least two independent experiments with multiple technical replicates in each experiment. Significance was set for a *P* value of < 0.05.

**RESULTS**

***Reduced expression of FFARs in CRC patients***

In a cohort of 18 CRC patients, we used microarray data to compare the expression levels of *FFAR2* gene in matched tumour-normal tissue samples and found it to be significantly downregulated (Figure 1A). There was -1.388 ± 0.2065 times difference between means of normal tissue versus matched tumour for *FFAR2* gene signals (*P* < 0.0001). But there was no significant difference in expression levels of *FFAR3* gene (Figure 1B). We did qRT-PCR analysis for *FFAR2* and *FFAR3* genes on five CRC patient samples that were available from our patient cohort. In these five CRC patients, there was significant downregulation of both *FFAR2* and *FFAR3* genes. *FFAR2* gene showed 5.388 times down regulation in these samples (*P* = 0.0066) and FFAR3 was also down regulated more than two times (Figure 1C and D). A heat map visualization of FFAR2 and FFAR3 expression in each of the patient tumour-normal matched tissues reflects the inter-patient heterogeneity (Figure 1E). We analysed other members of FFARs namely FFAR1 and FFAR4 in these patients and found no significant difference in their expression levels (Supplementary Figure 1 and Supplementary Table 5).

***Loss of FFARs in HCT116 cells results in increased proliferation***

We engineered HCT116 cells to downregulate the expression of *FFAR2* and *FFAR3* genes. First, we knocked down *FFAR2* gene and obtained 63% reduced stable expression. In this cell line FFAR3 levels were also found to be slightly affected with 32% reduction in expression. Next, we down regulated FFAR3 levels in this cell line. We found stable reduced expression of FFAR2 (77%) and FFAR3 (68%) in this cell clone. These two cell lines were chosen for all further experiments (Figure 2A). We measured the rate of proliferation of these HCT116 KD cells and found increased rate of proliferation as reflected by the cell index values in FFAR2 knock down cells (2.84 times compared to scrambled control. Double KD cells showed even higher increase in cell proliferation with 8.26 times cell index as compared to scrambled control (Figure 2B).

***Loss of FFAR2 and FFAR3 leads to increased glucose uptake and GLUT1 expression***

We measured the uptake of glucose in the HCT116 KD cells. HCT116 with FFAR2 KD showed about 1.5 times more glucose uptake whereas double KD exhibited even more with 1.8 times increase over the control-scrambled HCT116 cells (Figure 3A). We subsequently measured the levels of GLUT1 expression in these cells and found it to be significantly increased as higher fluorescence intensity was observed in modified cells (Figure 3B). GLUT1 expression pattern correlated with the glucose uptake in the modified cells as reflected in coefficient of determination value of 0.9615 (Figure 3C).

***Double KD cells exhibit increased cAMP production***

In order to understand the mechanistic role of FFARs in CRC, we attempted to interrogate the involvement of cAMP pathway. We found highly significant increase in cAMP levels in HCT116 double KD cells with more than 23 times increased cAMP levels as compared to FFAR2 KD and control-scrambled HCT116 cells (Figure 4A). Interestingly, the lack of significant change in cAMP level in FFAR2 KD HCT116 cells with a significant increase in rate of cell proliferation suggests a cAMP independent effect of FFAR2. To further validate this hypothesis, we blocked the effect of cAMP using the pharmacological inhibitor H89, which blocks the function of PKA, and tested the rate of cell proliferation in the presence or absence of H89. H89 showed a saturated inhibitory effect on HCT116 cells above 1 μmol/L concentration (Figure 4B). As expected, we did not observe any difference in rate of cell proliferation in the presence or absence of H89 in HCT116 KD clones confirming our hypothesis of an PKA independent mechanism of cAMP activity in HCT116 KD cells (Figure 4C). H89 also showed sustained inhibition up to 48h, which confirmed the efficacy during the entire duration of the cell proliferation experiment (Supplementary Figure 2).

**DISCUSSION**

As the evidence is supporting the strong connection between diet and CRC, there is an increased quest to understand the underlying molecular mechanism. Both long and short chain free fatty acids have been shown to be associated with cancer and metastasis[23-27]. This becomes relevant especially in CRC where high fat diet has been strongly correlated with the initiation and progression of disease. Receptors for these fatty acids would be good targets for designing prevention strategies. But the available evidence so far is not clear on establishing their role in different types of cancer. These receptors have shown to mediate increased cancerous activity as well as reduction in growth in different types of cancer cells. Also, these receptors belong to family of G-protein coupled receptors which are favourite molecules as drug targets[28]. In the present study, we focused on understanding the role of gut microbiota derived SCFA receptors in CRC. We used both patient samples and HCT116 cell lines in this study. FFAR2 and FFAR3 are well known receptors for SCFAs. FFAR2 has been implicated in CRC[29] but there is no known evidence for FFAR3 association. There are reports where the heterodimers of these receptors have been suggested to signal the short chain fatty acids. Earlier, we had reported cytogenetic and gene expression profile of CRC patients[22,30,31]. We used this data to analyse the expression profile of short chain FFARs in patient cohort. While we found significant down regulation of *FFAR2* gene in few CRC patients, the variability of expression was high. This further supports the notion of inter-patient heterogeneity observed in cancer especially CRC[32,33] and strengthens argument in favour of personalized medicine[8]. We further validated our results using available CRC patient mRNA samples and carried out qRT-PCR based expression analysis. Both microarray and qRT-PCR analyses confirmed down regulation of *FFAR2* gene in tumour samples. Our results thus confirm previous reports of down regulated *FFAR2* gene in CRC[29]. However, our observations regarding FFAR3 are novel and this study underscores the importance of studying the two receptors together. While we observed no significant difference in expression level of *FFAR3* gene in CRC patient cohort, there was significant down regulation observed in qRT-PCR data as reported earlier[29]. This could be due to the variability observed in these selected patient samples and/or due to higher sensitivity of qRT-PCR assay and possible differences in the two techniques[34].

In order to establish the role of short chain FFARs, we engineered HCT116, a colon cancer cell line to reduce the expression of *FFAR2* and *FFAR3* genes using shRNA technology. Our hypothesis suggested increased cell proliferation in HCT116 cells with reduced expression of short chain FFARs. Two cell line clones from HCT116 were generated – One with reduced expression of *FFAR2* gene alone and another with reduced expression of both *FFAR2* and *FFAR3* genes. FFAR2 KD cells showed highly significant increase in cell proliferation whereas double KD showed comparatively enhanced effect in cell proliferation as well as glucose uptake and cAMP production. This is a clear evidence of FFAR2 and FFAR3 function as tumour suppressors via mechanism that needs to be fully understood. A recent report suggested epigenetic dysregulation of inflammation suppressors by FFAR2[19]. CRC cells are known to uptake glucose at a higher rate and Warburg effect is a hallmark of cancer cells[35]. HCT116 with reduced levels of FFAR2 and FFAR3 also displayed increased uptake of glucose in an additive manner, which could be responsible for increased cell proliferation. We suggest the involvement of FFAR2 and FFAR3 in glucose metabolism but this needs to be further studied to better understand the network affected by the reduced expression of these genes. Increased glucose uptake in the engineered cells was accompanied with an increased expression of GLUT1 – a well-known glucose transporter. Overexpression of GLUT1 has been suggested as a negative prognostic biomarker in CRC and indicator of clinically aggressive disease[36]. Our results thus suggest a previously unknown important connection between FFAR2/FFAR3 and glucose metabolism. Increased glucose metabolism has been known to be induced by short chain fatty acids[37].

To further understand the effect of FFAR2/FFAR3 on the signalling pathways, we measured the cAMP levels in engineered cells. Intracellular cAMP level has been shown to regulate cellular motility[38].cAMP has also been shown to suppress apoptosis in CRC cells[39]. There was a huge increase in cAMP levels in double KD cells which was correlated with increased cell proliferation. However, FFAR2 single KD cells showed an increased cell proliferation and glucose uptake without any changes in cAMP levels. These results suggest that impact of FFAR2/FFAR3 on cell proliferation and glucose uptake are independent of cAMP signalling pathway. To further evaluate the role cAMP pathway, we inhibited PKA, a known downstream target of cAMP signalling by H89 molecule which is a known PKA inhibitor[40]. Inhibition of PKA had no impact on the rate of cell proliferation. Some studies have shown FFAR2/FFAR3 to signal through other pathways like p38 and JNK signalling[41] and Hippo-Yap pathway[42]. These pathways may be involved in mediating FFAR2/FFAR3 effect in our study and are an interesting area of research for future projects.

As illustrated in (Figure 5), our results conclusively establish the role of FFAR2 and FFAR3 in increased proliferation of CRC cells. This study also provides evidence to suggest the involvement of GLUT1 and PKA independent cAMP signalling pathway which needs to be further studied for identifying therapeutic targets and biomarkers for CRC progression.

**ARTICLE HIGHLIGHTS**

***Research background***

Colorectal cancer (CRC) has been linked with free fatty acid receptors (FFARs). However, the mechanism of action of FFARs in CRC needs to be better studied.

***Research motivation***

To generate evidence that can better explain the role of diet in CRC and its association with gut microbiome.

***Research objectives***

To understand how FFAR2 and FFAR3 contribute to CRC cell growth and metabolism.

***Research methods***

Cell culture, RNA interference (RNAi), Transfection, quantitative real time PCR, Western blot, Glucose uptake assay, cAMP assay, real time cell proliferation assay, Immunofluorescence, statistical and computational analyses.

***Research results***

FFAR2 is downregulated in CRC patient samples. CRC cells with reduced levels of *FFAR2* and *FFAR3* genes expression show increased rate of proliferation. Increased levels of glucose transporter and subsequent increase in glucose uptake is observed alongside increased cAMP levels in cells with reduced expression of FFAR2 and FFAR3.

***Research conclusions***

Short chain FFARs FFAR2 and FFAR3 may contribute in increased cell proliferation by increased glucose uptake.

***Research perspectives***

Processing of food by gut microbiota could be associated with initiation, progression and severity of CRC. Modifying dietary profiles for high-risk individuals may be a preventive measure for CRC.

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**Footnotes**

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**Conflict-of-interest statement:** The authors declare that they have no conflict of interest.

**Data sharing statement**: This patient microarray dataset is available on Gene Expression Omnibus, National Center for Biotechnology Information, National Institute of Health, United States (Accession: GSE 50421).

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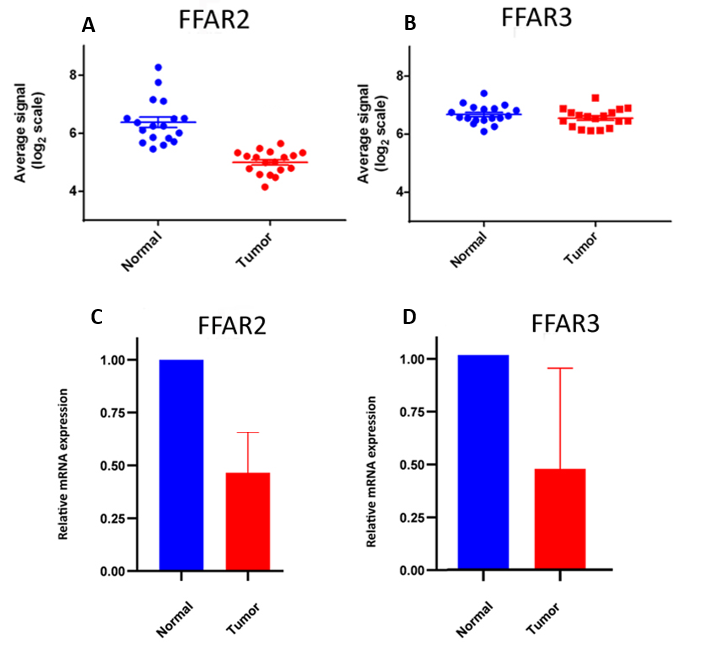
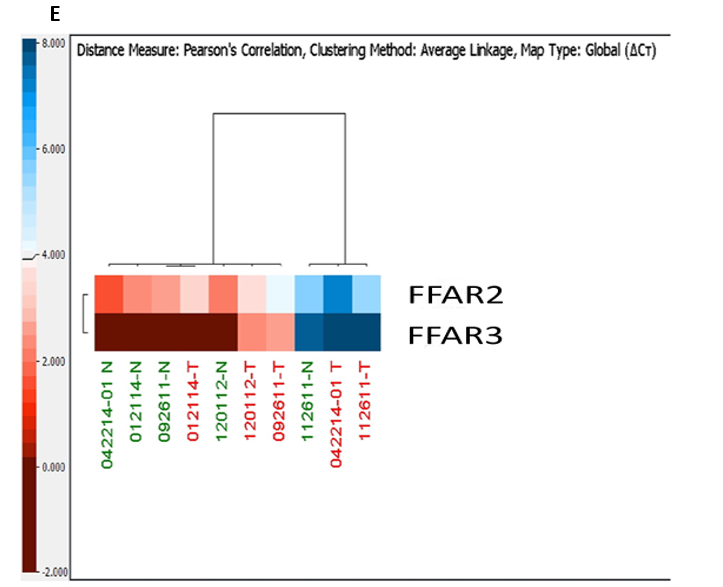
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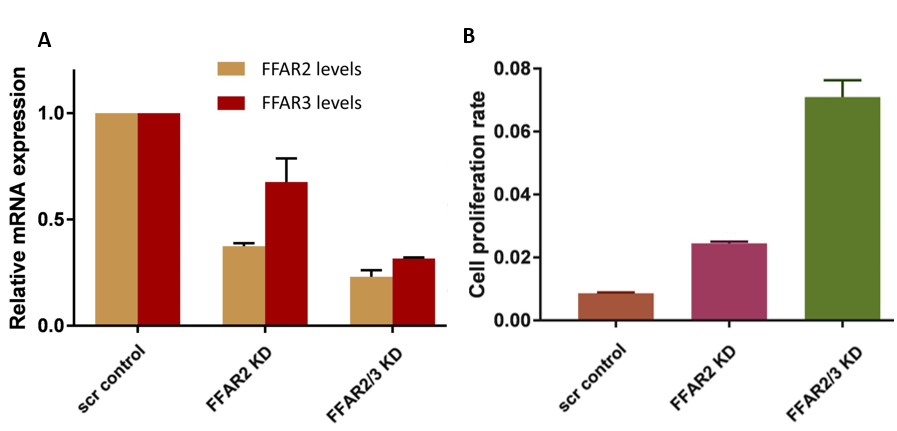
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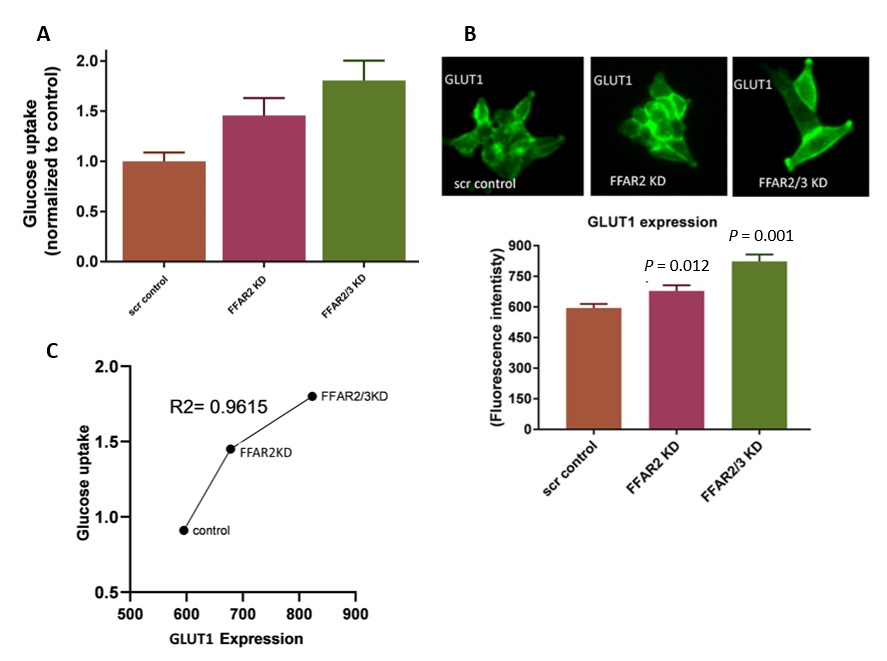
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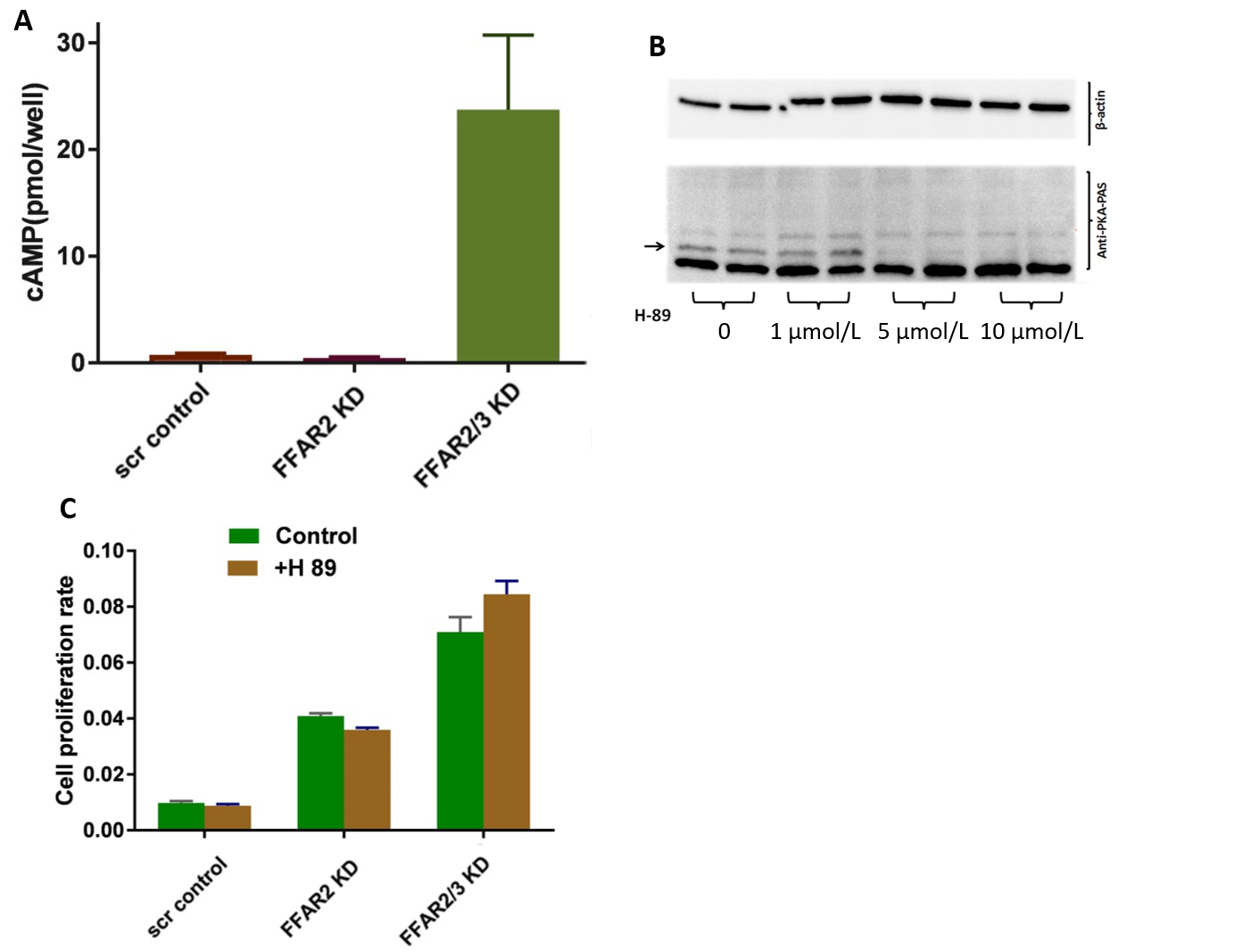
**Figure 1 Gene expression analysis of *free fatty acid receptor 2/3* genes in patients diagnosed with colorectal cancer.** A: Microarray analysis of gene expression of free fatty acid receptor (FFAR) 2 in tumor-normal paired tissues; B: Microarray analysis of gene expression of FFAR3 in tumor-normal paired tissues; C: Quantitative polymerase chain reaction (QPCR) gene expression analysis of FFAR2 in tumor-normal paired tissues; D: QPCR gene expression analysis of FFAR3 in tumor-normal paired tissues; E: Heatmap representation of *FFAR2* and *FFAR3* genes expression using QPCR in tumor-normal paired tissues. Color intensity ruler is given to represent Threshhold cycle (Ct) values with a range of -2 to 2 which is inverse of expression level. More red means less Ct and hence more expression. More blue reflects More Ct value suggesting less expression. Each bar represents an average of at least two independent experiments with multiple technical replicates in each experiment. Significance was set for a *P* value of < 0.05 (paired *t* test). FFAR: Free fatty acid receptor; Ct: Threshhold cycle.



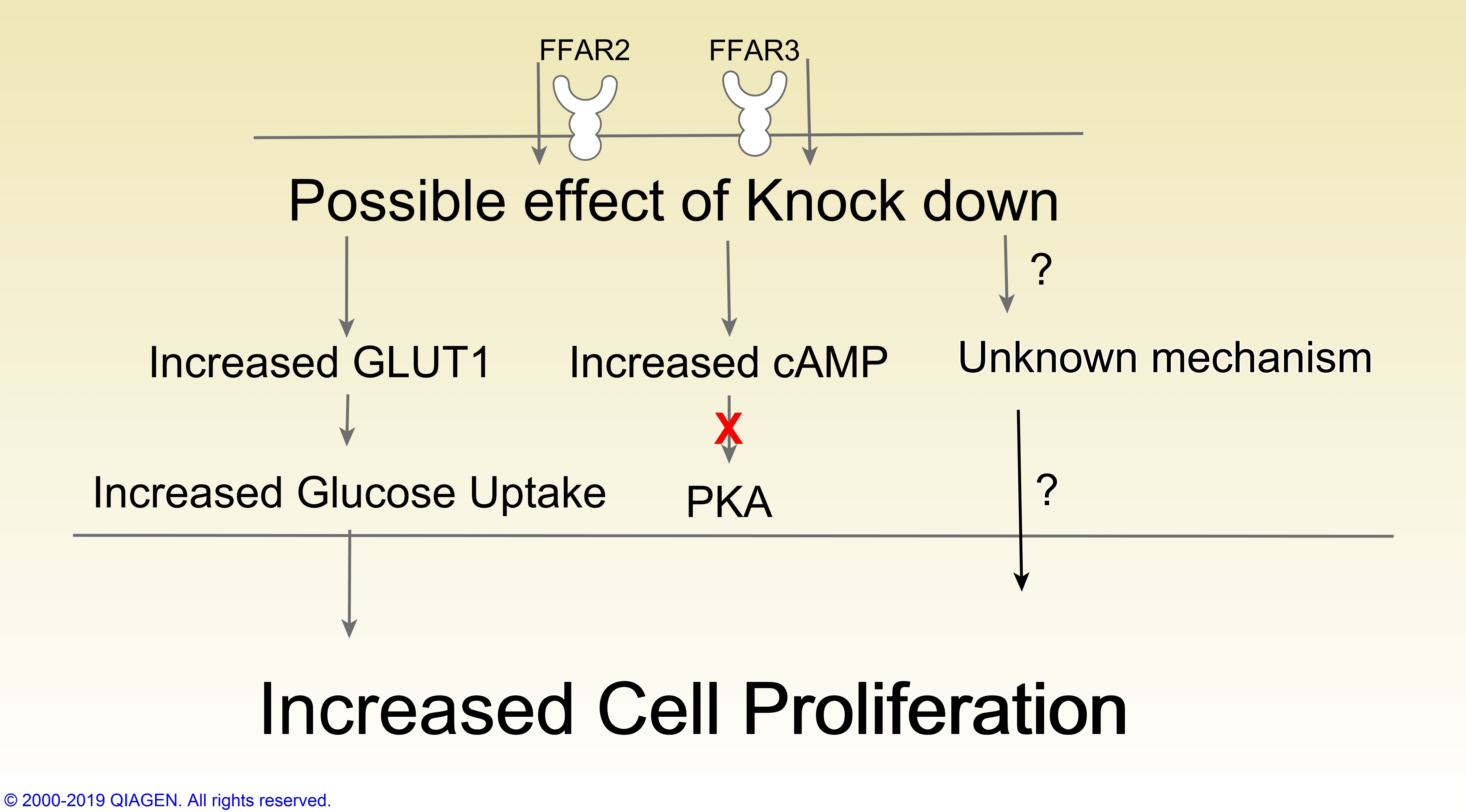
**Figure 2 Increased proliferation of colorectal cancer cells in single free fatty acid receptor 2 and double free fatty acid receptor 2/3** **knockdown colorectal cancer cells.** A: Relative mRNA expression in single free fatty acid receptor (FFAR) 2 and double FFAR2/3 knockdown clones compared to scrambled control; B: xCELLigence proliferation assay of FFAR2 and FFAR2/FFAR3 double knockdown clones. Each barrepresents an average of at least two independent experiments with 2 technical replicates in each experiment. Significance was set for a *P* value of < 0.05. FFAR: Free fatty acid receptor; KD: knockdown.



**Figure 3 Single knockdown of free fatty acid receptor 2 and double knockdown of free fatty acid receptor 2/3 increases glucose uptake and glucose transporter 1 expression in colorectal cancer cells.** A: Glucose uptake in single free fatty acid receptor (FFAR) 2 and double FFAR2/3 knockdown clones compared to control; B: Immunofluorescence for glucose transporter 1 in single FFAR2 and double FFAR2/3 knockdown clones compared to scrambled control; C: Correlation plot between glucose transporter 1 immunofluorescence and glucose uptake. *R*2 value close to 1 suggest a good correlation. Each bar represents an average of at least two independent experiments with multiple technical replicated in each experiment. Significance was set for a *P* value of < 0.05. FFAR: Free fatty acid receptor; KD: knockdown; GLUT1: Glucose transporter 1.



**Figure 4 Increased cyclic adenosine monophosphate levels in double free fatty acid receptor 2/3 knockdown colorectal cancer cells.** A: Cyclic adenosine monophosphate level in single free fatty acid receptor (FFAR) 2 and double FFAR2/3 knockdown clones compared to scrambled control; B: Western blot analysis of effect of different doses of H89 in inhibiting phosphorylation of protein kinase A. Dose of 5 μmol/L and 10 μmol/L showed complete inhibition; C: Cell proliferation of single FFAR2 and double FFAR2/3 knockdown clones compared to scrambled control with and without protein kinase A pharmacologic inhibitor. FFAR: Free fatty acid receptor; KD: knockdown; cAMP: Cyclic adenosine monophosphate; PKA: Protei kinase A.



**Figure 5 An illustration depicting the possible role of free fatty acid receptor 2 and 3 that results into increased cell proliferation.** This figure was generated through the use of IPA (QIAGEN Inc., <https://www.qiagenbioinformatics.com/products/ingenuity-pathway-analysis>). FFAR: Free fatty acid receptor; GLUT1: Glucose transporter 1; cAMP: Cyclic adenosine monophosphate; PKA: Protei kinase A.