WITHDRAWN ARTICLE

WITHDRAWN: LncRNA EGOT regulates the proliferation and apoptosis of colorectal cancer by miR-33b-5p/CROT axis

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LncRNA EGOT regulates the proliferation and apoptosis of colorectal cancer by miR-33b-5p/CROT axis

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18

19 Abstract

Accumulating researches have proved that long noncoding RNAs (lncRNAs) regulate a variety of cellular processes during cancer progression. However, the detailed function of most lncRNAs in

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colorectal cancer (CRC) remains mostly unknown. This study was aimed 23 at exploring the specific role of lncRNA EGOT in CRC. Data from this 24 study revealed that EGOT expression was obviously upregulated in CRC 25 tissues and cell lines, and high EGOT expression indicated poor overall 26 survival of CRC patients. Besides, functional assays proved that EGOT 27 knockdown inhibited cell proliferation and promoted cell apoptosis in 28 CRC. Then, subsequent molecular mechanism assays uncovered that 29 EGOT could bind with miR-33b-5p and negatively regulate miR-33b-5p 30 expression. Additionally, CROT was a downstream target of miR-33b-5p. 31 Further, rescued-function assays suggested that the suppressive influence 32 of EGOT depletion on CRC progression was reversed by miR-33b-5p 33 inhibition or CROT overexpression. In conclusion, IncRNA EGOT 34 mediates the tumor-facilitating part in CRC via miR-33b-5p/CROT 35 pathway. 36

Key words: EGOT; CRC; miR-33b-5p; CROT.

38

39 Introduction

Colorectal cancer (CRC) is a kind of frequent malignant cancer with poor clinical outcome (1). The only hope of treating precancerous CRC might be the radical surgery (2). However, even after the surgery, radiation and chemotherapy still have little effect (3). Previous researches have indicated that some molecular markers are associated with prognosis, but

Long noncoding RNAs (lncRNAs) are recognized as a group of RNA 48 molecules which are longer than 200 nucleotides and lack any open 49 reading frame to encode protein. Besides, they have been previously 50 manifested as crucial regulators in diverse tumors (5, 6). Specifically, 51 IncRNA SPRY4-IT1 contributes to the tumorigenesis of gastric cancer (7); 52 IncRNA PANDAR predicts poor prognosis and is involved in 53 tumorigenesis in hepatocellular carcinoma (8); and lncRNA LUNAR1 54 facilitates cell proliferation and high expression of LUNAR1 leads to 55 poor prognosis in diffuse large B-cell lymphoma (9). LncRNA EGOT is 56 backward-looking at the level of nucleotide and was initially found to be 57 implicated in the development of eosinophils and expressed in mature 58 eosinophils (10). In addition, EGOT has been revealed to have a high 59 expression in bone marrow and function in the development of bone 60 marrow hematopoietic stem cells (11). In recent years, the close 61 relationship between abnormally expressed EGOT and tumorigenesis has 62 been clarified in many cancers (11-15). The findings of these researches 63 imply that dysregulation of EGOT is tightly associated with prognosis 64 and that EGOT plays significant role in the occurrence and development 65 of tumors. However, the specific role of EGOT in CRC is largely 66

This study was the first attempt to investigate the underlying function and mechanism of EGOT in CRC, aiming to shed new lights on CRC treatment.

71

72 Materials and methods

73 Clinical specimens

The 70 pairs CRC tissue specimens and matched non-tumor tissue specimens were collected from CRC patients from 2013 and 2018. The ethical approval of this study was obtained from the Ethics Committee of Harbin Medical University Cancer Hospital. All the participants didn't receive any treatment and they all signed the informed consents prior to this study. All clinical samples were instantly frozen in liquid nitrogen and further stored at -80°C.

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82 Cell lines

Human normal cell line NCM460 and four CRC cell lines including
SW480, SW620, HCT15 and HT29, were cultivated continuously in
DMEM (Gibco, Grand Island, NY, USA) at 37°C and 5% CO₂. All cell
lines were purchased from the American Type Culture Collection (ATCC;
Rockville, Maryland, USA). Culture medium was added with 10% fetal
bovine serum (FBS; Gibco) and antibiotics as supplements.

Quantitative real-time polymerase chain reaction (qRT-PCR) 90 Extraction of total RNAs from SW620 and HT29 cells or tissue samples 91 were performed using TRIzol Regent (Invitrogen, Carlsbad, CA, USA). 1 92 µg of sample was reversely transcribed into cDNA for conducting qPCR 93 in the presence of SYBR[®] Premix Ex Taq[™] II (Takara, Tokyo, Japan) on 94 the Step-One Plus Detection System (Applied Biosystems, Foster City, 95 CA, USA). RNA level was calculated by the comparative $2^{-\Delta\Delta Ct}$ method. 96 The sequences of primers were listed in Supplementary Table 1. 97

98

99 Cell transfection

CRC cells (SW620 and HT29) were seeded in the 6-well and grown in an 100 incubator with CO_2 at 37°C. The two specific shRNAs to EGOT 101 (sh-EGOT#1/2), CROT (sh-CROT#1/2) and control shRNAs (sh-NC), as 102 well as pcDNA3.1/CROT (with empty pcDNA3.1 vector as control), 103 RiboBio (Guangzhou, China). Moreover, all produced at were 104 miR-33b-5p mimics and NC mimics, miR-33b-5p inhibitor and NC 105 inhibitor, were bought from Genechem (Shanghai, China). Cell 106 transfection was performed for 48 h in the presence of Lipofectamine 107 2000 reagent (Invitrogen). The shRNA sequences were shown in 108 Supplementary Table 2. 109

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111 Colony formation assay

¹¹² Transfected CRC cell lines were plated into 6-well plates $(1 \times 10^3/100 \,\mu\text{L})$ ¹¹³ at 37°C with 5% CO₂ for 2 weeks. Colonies were rinsed in ¹¹⁴ phosphate-buffered saline (PBS), fixed in 5% paraformaldehyde and ¹¹⁵ labeled with 1 mL of 0.1% crystal violet solution. After counting ¹¹⁶ manually, visible colonies were recorded.

117

118 TUNEL assay

CRC cells were transfected with indicated plasmids and harvested for
TUNEL assay. Fixed cells in 1% formaldehyde were subjected to 0.2%
Triton X-100 for permeabilization and dUTP-end labeling kit (Clontech,
Mountain View, CA, USA). Cells were counterstained with DAPI and
analyzed by fluorescence microscope (NIKON, Tokyo, Japan).

124

125 Western blot

Cells were seeded into 6-well plates prior to being collected for western blot analysis. The total proteins were isolated from the cell lines using RIPA buffer reagent (Thermo Fisher Scientific, Rochester, New York, USA) separated with 10% SDS-PAGE, and transferred onto PVDF membranes on ice overnight. Bovine serum albumin (BSA, 5%; Beyotime, Shanghai, China) was used to block membranes for 2 h at room temperature. Afterwards, membranes were subjected to primary

antibodies and secondary antibodies in succession. Primary antibodies 133 including anti-Bcl-2 (ab692), anti-Bax (ab182733), anti-cleaved 134 caspase-3 (ab2302), anti-total caspase-3 (ab2302), anti-CROT (ab175450), 135 anti-Tubulin (ab210797) and anti-GAPDH (ab22555), together with 136 HRP-labeled secondary antibodies were all acquired from Abcam 137 (Cambridge, MA, USA). Subsequent to washing in PBS, bands were 138 visualized with enhanced chemiluminescence (ECL) system (Bio-Rad lab, 139 Hercules, CA, USA). 140

141

142 Subcellular fractionation

In line with the manufacturer's recommendations of PARIS[™] Kit (Invitrogen), subcellular fractionation was assayed in SW620 and HT29 cells. The cell lysates from cell fractionation buffer were first centrifuged and then mixed with cell disruption buffer. The levels of EGOT in cytoplasmic and nuclear fractions were determined by qRT-PCR, with GADPH and U6 as cytoplasmic and nuclear, respectively. Downloaded from http://port.silverchair.com/bioscirep/article-pdf/doi/10.1042/BSR20193893/908269/bsr-2019-3893.pdf by guest on 17 April 2024

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150 **Dual-luciferase reporter vector analysis**

For EGOT and CROT 3'UTR luciferase analysis, the pmirGLO Dual-Luciferase reporter constructs containing wild type or mutant miR-33b-5p interacting sequences were formed and termed as EGOT-WT/Mut and CROT-WT/Mut. SW620 or HT29 cells in 24-well plates were co-transfected with above reporter constructs and indicated
transfection plasmids for 48 h, followed by analysis by the Dual
Luciferase Assay System (Promega, Madison, WI, USA). Firefly
luciferase activity was normalized to the corresponding Renilla luciferase
activity.

160

161 **RNA immunoprecipitation (RIP)**

Using EZ-Magna RIP Kit (Millipore, Billerica, MA, USA), RIP assay was carried out in CRC cells. SW620 and HT29 cells were washed in ice-cold PBS and lysed in RIP lysis buffer. Human anti-Ago2 antibody and nonspecific control normal IgG antibody (Millipore) were used for immunoprecipitation. RIP lysates were cultured with magnetic beads all night and then precipitated RNAs were purified for qRT-PCR analysis. Downloaded from http://port.silverchair.com/bioscirep/article-pdf/doi/10.1042/BSR20193893/908269/bsr-2019-3893.pdf by guest on 17 April 2024

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169 **RNA pull-down assay**

The wild type or mutant CROT sequences which contained the possible 170 miR-33b-5p binding sites obtained and biotinylated into 171 were Bio-CROT-WT/Mut. Cell extracts from RIPA lysis buffer were treated 172 with Bio-CROT-WT/Mut or Bio-NC. After adding magnetic beads, the 173 complex was centrifuged and evaluated by qRT-PCR. 174

175

176 Statistical analyses

Graph generation and data analyses in this study were both disposed via PRISM 6 (GraphPad, San Diego, CA, USA). Besides, data were showed as the mean \pm SD of three independent assays. Significant differences between groups were assayed through Student's t test or ANOVA, with threshold of probability (p-value) less than 0.05.

182

183 **Results**

Downregulation of EGOT inhibited cell proliferation and promoted cell apoptosis in CRC

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To make the exploration of the underlying role that EGOT played in CRC, 186 we first detected its expression status in tissue samples obtained from 187 CRC patients. And the data from qRT-PCR uncovered that EGOT was 188 highly expressed in CRC tissue specimens versus peri-tumor specimens 189 (Fig. 1A). Then clinical data revealed that high expression of EGOT 190 resulted in low survival rate of CRC patients (Fig. 1B). Next, the 191 expression of EGOT was examined in the normal NCM460 cells and four 192 CRC cell lines by qRT-PCR. The result showed that EGOT expression 193 was relatively high in CRC cell lines, particularly in SW620 and HT29 194 cells (Fig. 1C). Therefore, SW620 and HT29 cells were selected for the 195 following experiments. After knocking down EGOT in SW620 and HT29 196 cells by transfection with sh-EGOT#1/2 (Fig. 1D), cell proliferation 197 ability was weakened (Fig. 1E). Conversely, data from TUNEL assay 198

revealed that knockdown of EGOT facilitated the apoptosis of SW620 199 and HT29 cells (Fig. 1F). Besides, western blot analysis demonstrated 200 that downregulation of EGOT decreased the expression of Bcl-2 whereas 201 increased the expression of Bax and cleaved caspase-3, further validating 202 that silenced EGOT could induce the apoptosis of SW620 and HT29 cells 203 (Fig. 1G). All these data clarified that EGOT was overexpressed in CRC 204 tissues and cells, and EGOT depletion suppressed cell proliferation but 205 promoted cell apoptosis in CRC. 206

207 MiR-33b-5p combined with EGOT

Afterwards, we aimed to study the mechanism of EGOT in CRC, the 208 subcellular fractionation assay was first implemented to find out the 209 subcellular location of EGOT. Interestingly, the result showed that EGOT 210 was mostly located in the cytoplasm of SW620 and HT29 cells (Fig. 2A), 211 implying that EGOT was involved in the posttranscriptional regulation of 212 gene expression. After utilizing online starBase, 7 miRNAs which could 213 possibly bind with EGOT were predicted. Then, data from qRT-PCR 214 revealed that only the expression of miR-33b-5p was found affected most 215 by knockdown of EGOT (Fig. 2B). To further investigate the interaction 216 between miR-33b-5p and EGOT, starBase was applied again to predict 217 the binding sites between EGOT and miR-33b-5p (Fig. 2C). After 218 elevating miR-33b-5p expression in SW620 and HT29 cells by 219 transfection with miR-33b-5p mimics (Fig. 2D), the luciferase activity of 220

EGOT-WT reporter was decreased a lot. Nevertheless, when the binding 221 sequences of EGOT were mutated, the luciferase activity of EGOT-Mut 222 reporter showed no clear changes between two groups (Fig. 2E). This 223 indicated that EGOT could interact with miR-33b-5p at predicted sites. 224 Moreover, qRT-PCR analysis demonstrated that the expression of 225 miR-33b-5p was signally elevated in CRC cell lines when compared with 226 the normal cell line (Fig. 2F). Further, we observed that the proliferation 227 ability of SW620 and HT29 cells was obviously decreased after 228 miR-33b-5p was overexpressed (Fig. 2G). On the contrary, upregulation 229 of miR-33b-5p could promote the apoptosis of SW620 and HT29 cells 230 (Fig. 2H). Taken together, EGOT targeted the tumor-suppressive 231 miR-33b-5p in CRC. 232

233 CROT was the target of miR-33b-5p

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Based on the above findings, we proceeded to find the targets of 234 miR-33b-5p by utilizing starBase. With the help of five bioinformatics 235 tools (miRmap, PITA, TargetScan, PicTar and RNA22), two possible 236 candidate targets of miR-33b-5p were screened out (Fig. 3A). Then, it 237 was exhibited that under the condition of EGOT knockdown, CROT 238 expression was downregulated while NAA15 expression was not 239 influenced in SW620 and HT29 cells (Fig. 3B). Thus, CROT was chosen 240 for further analysis. Later, the binding sites between CROT and 241 miR-33b-5p were predicted by starBase (Fig. 3C). Afterwards, the 242

luciferase reporter assay suggested that elevating miR-33b-5p expression 243 decreased the luciferase activity of CROT-WT reporter but not that of 244 CROT-Mut reporter (Fig. 3D). Furthermore, RIP and RNA pull down 245 assays further proved that CROT could bind with miR-33b-5p (Fig. 246 3E-F). More importantly, it was proved that the suppressive effect of 247 silenced EGOT on the mRNA and protein levels of CROT was reversed 248 by miR-33b-5p inhibition (Fig. 3G-H). These data showed that EGOT 249 regulated the expression of CROT in CRC by competitively binding with 250 miR-33b-5p. Then, the expression of CROT was found upregulated in 251 CRC cell lines (Fig. 3I). Moreover, cell proliferation capability was 252 attenuated whereas the apoptosis capability was enhanced in SW620 and 253 HT29 cells with CROT depletion (Fig. 3J-L). Briefly, CROT was the 254 target of miR-33b-5p and CROT functioned as a tumor promoter in CRC. 255

LncRNA EGOT regulated the proliferation and apoptosis of CRC by targeting miR-33b-5p/CROT axis

After validating the existence of EGOT/miR-33b-5p/CROT network in CRC, we further studied the influence of this competing endogenous RNA (ceRNA) mechanism on CRC cell proliferation and apoptosis. Prior to rescued-function assays, we conducted qRT-PCR assay and obtained a satisfactory efficiency of CROT overexpression (Fig. 4A). Expectedly, we unveiled that miR-33b-5p inhibition or CROT upregulation could rescue the suppressive influence of EGOT depletion on cell proliferation

(Fig. 4B). Likewise, the increased cell apoptosis rate caused by EGOT 265 silence reversed by inhibiting miR-33b-5p expression was or 266 overexpressing CROT (Fig. 4C), while such phenomenon was further 267 testified by western blot data (Fig. 4D). To sum up, EGOT regulated CRC 268 cell proliferation and apoptosis via miR-33b-5p/CROT axis. 269

270

271 **Discussion**

CRC is the leading cause of cancer-related mortality worldwide. 272 Accumulating evidence has verified that lncRNA is implicated in the 273 progression of many malignant tumors, including CRC (16-19). Disorder 274 of EGOT is a common phenomenon in different kinds of cancers, such as 275 breast cancer (11), gastric cancer (14) and renal cell carcinoma (12). As 276 far as we know, few studies have focused on the specific role of lncRNA 277 EGOT in CRC. Therefore, further study of EGOT in CRC may do some 278 help for the treatment of this fatal disease. This study first revealed the 279 expression pattern of EGOT in CRC and further explored the impact of 280 EGOT on CRC cell proliferation and apoptosis, along with its underlying 281 mechanism. 282

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In our study, EGOT was expressed at high levels in CRC tissues and cells, and EGOT predicted poor prognosis of CRC patients, indicating that EGOT might be a possible biomarker for CRC diagnosis. Next, we studied the impact of EGOT on CRC cell proliferation and apoptosis.

Data from loss-of-function assays confirmed the tumor-accelerating effect 287 of EGOT on CRC tumorigenesis. Moreover, Bcl-2 and Bax are essential 288 regulators for the activity of cell apoptosis. Accumulating studies have 289 shown that lncRNAs regulate the expression of Bcl-2 and Bax in various 290 cancers. Yin Z et al uncovered that PPMD1 depletion could increase the 291 protein level of Bax and decrease the protein level of Bcl-2 expression, 292 thus inducing cell apoptosis in colon cancer (20). Wang AH et al 293 elucidated that lncRNA PCAT-1 could regulate Bax and Bcl-2 expression 294 and facilitate cell apoptosis in CRC (21). In this paper, we proved that 295 EGOT depletion decreased Bcl-2 expression and enhanced Bax 296 expression in CRC and induced cell apoptosis in CRC. Besides, several 297 previous findings unveiled that EGOT promoted tumorigenesis in gastric 298 cancer through activating Hedgehog pathway (22), which provided 299 another way for the potential downstream pathway for EGOT in 300 regulating CRC. 301

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Since ceRNA is a typical post-transcriptional pattern and EGOT was mainly distributed in cytoplasm of CRC cells, we assumed that EGOT served as a ceRNA via sponging certain miRNA to regulate CRC cells. Via a range of molecular mechanism assays and rescue tests, we verified that EGOT competitively bound with miR-33b-5p to enhance CROT expression, thus facilitating CRC cell proliferation. Interestingly, Yang X et al. revealed that miR-33b-5p could sensitize gastric cancer cells to

chemotherapy drugs via targeting HMGA2 (23). Moreover, Zhao M et al. 309 previously disclosed the existence of ceRNA network containing 310 miR-33b-5p clarified that in prostate cancer. and 311 CUL4B/miR-33b/C-MYC axis contributed to the progression of prostate 312 cancer (24). However, CROT was scarcely researched in cancers. 313 Previously, Chen X et al. suggested that FECH/CROT axis could predict 314 recurrence of prostate cancer (25). Besides, Hage-Sleiman R et al. 315 indicated that CROT was involved in oxidative stress response in 316 leukemia cells of Molt-4 (26). 317

Further, lncRNA-associated ceRNA pattern in CRC has been widely 318 reported. Yu C et al. unveiled that lncRNA AC009022.1 could promote 319 CRC cell proliferation, migration and invasion via promotion on 320 ACTR3B expression through miR-497-5p suppression (27). And Wang Y 321 et al. suggested that lncRNA TTN-AS1 enhanced KLF15 expression via 322 acting as a sponge of miR-376a-3p, therefore promoting CRC progression 323 (28). In addition, Wang Y et al. manifested that upregulation of lncRNA 324 CCAT5 could boost CRC cell growth, invasion and metastasis through 325 activation of STAT3 (29). Consistently, our study further provided the 326 evidence of the existence of ceRNA mechanism in CRC and 327 EGOT/miR-33b-5p/CROT axis was firstly confirmed to influence CRC 328 cell proliferation and apoptosis. 329

330 On the whole, present study unveiled a novel ceRNA pattern in which

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335 Ethical Standards

The ethical approval of tissue samples was obtained from the Ethics Committee of Harbin Medical University Cancer Hospital. All the participants signed the informed consents.

339

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No funding was received.

342

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343 Authors' contribution

This research was designed mainly by Yin Ni and most of the experiments were accomplished by Yin Ni and Chunbo Li; the experimental data was acquired by Chen Bo; the data was analyzed and interpreted by Bomiao Zhang; figures and tables was completed by Yanlong Liu and Xuefeng Bai; the writing and revision of the manuscript was finished by Bingbing Cui and Peng Han.

350

351 **Conflict of interest**

352 None.

353

354 Acknowledgements

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432 **Figure legends**

433 Figure 1 Down-regulation of EGOT inhibited cell proliferation and

434 promoted cell apoptosis in CRC.

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A. EGOT expression CRC tissue samples and peri-tumor samples wasdetected via qRT-PCR.

B. The association between EGOT expression and overall survival of
CRC patients was analyzed via Kaplan-Meier analysis.

C. The expression of EGOT was tested in the normal NCM460 cells andfour CRC cell lines by qRT-PCR.

D. The efficiency of EGOT knockdown in SW620 and HT29 cells wasexamined via qRT-PCR.

E. Colony formation assay was done to test the effect of EGOTknockdown on cell proliferation.

F. TUNEL assay was implemented to examine the impact of EGOT
depletion on cell apoptosis. Scale bar: 200 μm.

G. Western blot analysis was used to analyze the influence of EGOT deficiency on the expression of apoptosis-related proteins. **P < 0.01.

449 Figure 2 MiR-33b-5p combined with EGOT.

A. The subcellular fractionation assay was used to determine thesubcellular location of EGOT in SW620 and HT29 cells.

452 B. qRT-PCR was used to test the effect of silenced EGOT on the 453 expression of seven candidate miRNAs in SW620 and HT29 cells.

454 C. The binding sites between EGOT and miR-33b-5p were predicted by455 starBase.

19

E. Luciferase reporter assay was done to confirm the interaction betweenEGOT and miR-33b-5p.

F. The expression of miR-33b-5p in CRC cell lines compared to NCM460cells was examined via qRT-PCR.

G. The proliferation of SW620 and HT29 cells transfected with miR-33b-5p mimics or NC mimics was evaluated via colony formation assay.

465 H. The apoptosis of transfected cells was analyzed via TUNEL assay. 466 Scale bar: 200 μ m. *P < 0.05, **P < 0.01.

467 Figure 3 CROT was the target of miR-33b-5p.

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468 A. Target genes of miR-33b-5p were predicted by starBase.

B. The expression of CROT and NAA15 in SW620 and HT29 cells

transfected with sh-NC and sh-EGOT#1 was detected by qRT-PCR.

471 C. The binding sites between CROT and miR-33b-5p were predicted by472 starBase.

D-F. The interaction between CROT and miR-33b-5p was validated via
luciferase reporter, RIP and RNA pull down assays.

475 G. The efficiency of miR-33b-5p inhibition was examined by qRT-PCR.

476 H. The mRNA and protein levels of CROT in different groups were

477 detected via qRT-PCR and western blot.

- I. The expression of CROT in CRC cell lines and NCM460 cells weretested by qRT-PCR.
- J. The expression of CROT in SW620 and HT29 cells transfected with
 sh-NC, sh-CROT#1 and sh-CROT#2 was detected by qRT-PCR.
- 482 K. The colony formation assay was done to test the effect of CROT483 knockdown on cell proliferation.
- 484 L. TUNEL assay was implemented to examine the impact of silenced 485 CROT on cell apoptosis. Scale bar: 200 μ m. ^{**}P < 0.01.

Figure 4 LncRNA EGOT regulated the proliferation and apoptosis of
CRC by targeting miR-33b-5p/CROT axis.

- A. The efficiency of CROT overexpression was detected by qRT-PCR in
 SW620 and HT29 cells.
- B. The proliferation of transfected cells was assessed via colonyformation assay.
- C. TUNEL experiment was used to analyze the apoptosis of transfected
 cells. Scale bar: 200 μm.
- 494 D. The expression of apoptosis-associated proteins was measured via 495 western blot. $^*P < 0.01$.

496 Supplementary Table 1

497 The sequences of primers were listed.

498 **Supplementary Table 2**

499 The shRNA sequences were shown.







