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1 Deciphering colorectal cancer genetics through multi-omic analysis of 100,204
2 cases and 154,587 controls of European and East Asian ancestries

3
4 Ceres Fernandez-Rozadilla^{1,2^}, Maria Timofeeva^{3,4^}, Zhishan Chen^{5^}, Philip Law^{6^}, Minta Thomas^{7^},
5 Stephanie Schmit^{8,9^}, Virginia Díez-Obrero^{10,11,12,13^}, Li Hsu^{7,14^}, Juan Fernandez-Tajes¹, Claire
6 Palles¹⁵, Kitty Sherwood¹, Sarah Briggs¹⁶, Victoria Svinti³, Kevin Donnelly³, Susan Farrington³,
7 James Blackmur³, Peter Vaughan-Shaw³, Xiao-ou Shu⁵, Jirong Long⁵, Qiuyin Cai⁵, Xingyi Guo^{5,17},
8 Yingchang Lu⁵, Peter Broderick⁶, James Studd⁶, Jeroen Huyghe⁷, Tabitha Harrison⁷, David Conti¹⁸,
9 Christopher Dampier¹⁹, Mathew Devall¹⁹, Fredrick Schumacher^{20,21}, Marilena Melas²², Gad
10 Rennert^{23,24,25}, Mireia Obón-Santacana^{11,10,26}, Vicente Martín-Sánchez^{12,27}, Ferran Moratalla-
11 Navarro^{10,11,12,13}, Jae Hwan Oh²⁸, Jeongseon Kim²⁹, Sun Ha Jee³⁰, Keum Ji Jung³⁰, Sun-Seog
12 Kweon³¹, Min-Ho Shin³¹, Aesun Shin^{32,33}, Yoon-Ok Ahn³², Dong-Hyun Kim³⁴, Isao Oze³⁵, Wanqing
13 Wen⁵, Keitaro Matsuo^{36,37}, Koichi Matsuda³⁸, Chizu Tanikawa³⁹, Zefang Ren⁴⁰, Yu-Tang Gao⁴¹,
14 Wei-Hua Jia⁴², John Hopper^{43,44}, Mark Jenkins⁴³, Aung Ko Win⁴³, Rish Pai⁴⁵, Jane Figueiredo^{46,18},
15 Robert Haile⁴⁷, Steven Gallinger⁴⁸, Michael Woods⁴⁹, Polly Newcomb^{7,50}, David Duggan⁵¹, Jeremy
16 Cheadle⁵², Richard Kaplan⁵³, Timothy Maughan⁵⁴, Rachel Kerr⁵⁵, David Kerr⁵⁶, Iva Kirac⁵⁷, Jan
17 Böhm⁵⁸, Lukka-Pekka Mecklin⁵⁹, Pekka Jousilahti⁶⁰, Paul Knekt⁶⁰, Lauri Aaltonen^{61,62}, Harri
18 Rissanen⁶³, Eero Pukkala^{64,65}, Johan Eriksson^{66,67,68}, Tatiana Cajuso^{62,61}, Ulrika Hänninen^{62,61},
19 Johanna Kondelin^{62,61}, Kimmo Palin^{62,61}, Tomas Tanskanen^{62,61}, Laura Renkonen-Sinisalo⁶⁹, Brent
20 Zanke⁷⁰, Satu Männistö⁶³, Demetrius Albanes⁷¹, Stephanie Weinstein⁷¹, Edward Ruiz-Narvaez⁷²,
21 Julie Palmer^{73,74}, Daniel Buchanan^{75,76,77}, Elizabeth Platz⁷⁸, Kala Visvanathan⁷⁸, Cornelia Ulrich⁷⁹,
22 Erin Siegel⁸⁰, Stefanie Brezina⁸¹, Andrea Gsur⁸¹, Peter Campbell⁸², Jenny Chang-Claude^{83,84},
23 Michael Hoffmeister⁸⁵, Hermann Brenner^{85,86,87}, Martha Slattery⁸⁸, John Potter^{89,7}, Konstantinos
24 Tsilidis^{90,91}, Matthias Schulze^{92,93}, Marc Gunter⁹⁴, Neil Murphy⁹⁴, Antoni Castells⁹⁵, Sergi Castellví-
25 Bel⁹⁵, Leticia Moreira⁹⁵, Volker Arndt⁸⁵, Anna Shcherbina⁹⁶, Mariana Stern^{97,98}, Bens
26 Pardamean⁹⁹, Timothy Bishop¹⁰⁰, Graham Giles^{101,43,102}, Melissa Southey^{102,103,101}, Gregory
27 Idos¹⁰⁴, Kevin McDonnell^{104,24,25}, Zomoroda Abu-Ful²³, Joel Greenson^{105,24,25}, Katerina Shulman²³,
28 Flavio Lejbkovicz^{106,23,25}, Kenneth Offit^{107,108}, Yu-Ru Su¹⁰⁹, Robert Steinfeld⁷, Temitope Keku¹¹⁰,

29 Bethany van Guelpen^{111,112}, Thomas Hudson¹¹³, Heather Hampel¹¹⁴, Rachel Pearlman¹¹⁴, Sonja
 30 Berndt⁷¹, Richard Hayes¹¹⁵, Marie Elena Martinez^{116,117}, Sushma Thomas¹¹⁸, Douglas Corley^{119,120},
 31 Paul Pharoah¹²¹, Susanna Larsson¹²², Yun Yen¹²³, Heinz-Josef Lenz¹²⁴, Emily White^{7,125}, Li Li²¹,
 32 Kimberly Doheny¹²⁶, Elizabeth Pugh¹²⁶, Tameka Shelford¹²⁶, Andrew Chan^{127,128,129,130,131,132},
 33 Marcia Cruz-Correa¹³³, Annika Lindblom^{134,135}, David Hunter^{131,136}, Amit Joshi^{131,127}, Clemens
 34 Schafmayer¹³⁷, Peter Scacheri¹³⁸, Anshul Kundaje^{96,139}, Deborah Nickerson¹⁴⁰, Robert Schoen¹⁴¹,
 35 Jochen Hampe¹⁴², Zsofia Stadler^{143,108}, Pavel Vodicka^{144,145,146}, Ludmila Vodickova^{144,145,146},
 36 Veronika Vymetalkova^{144,145,146}, Nickolas Papadopoulos^{147,148,149}, Christopher Edlund¹⁸, William
 37 Gauderman¹⁸, Duncan Thomas¹⁸, David Shibata¹⁵⁰, Amanda Toland¹⁵¹, Sanford Markowitz¹⁵²,
 38 Andre Kim¹⁸, Stephen Chanock⁷¹, Franzel van Duijnhoven¹⁵³, Edith Feskens¹⁵⁴, Lori Sakoda^{119,7},
 39 Manuela Gago-Dominguez^{155,156}, Alicja Wolk¹²², Alessio Naccarati^{157,158}, Barbara Pardini^{157,158},
 40 Liesel FitzGerald^{159,101}, Soo Chin Lee¹⁶⁰, Shuji Ogino^{161,162,131,163}, Stephanie Bien⁷, Charles
 41 Kooperberg⁷, Christopher Li⁷, Yi Lin⁷, Ross Prentice^{7,164}, Conghui Qu⁷, Stéphane Béziau¹⁶⁵,
 42 Catherine Tangen¹⁶⁶, Elaine Mardis¹⁶⁷, Taiki Yamaji¹⁶⁸, Norie Sawada¹⁶⁹, Motoki Iwasaki^{168,169},
 43 Christopher Haiman¹⁷⁰, Loic Le Marchand¹⁷¹, Anna Wu¹⁷², Chenxu Qu¹⁷³, Caroline McNeil¹⁷³,
 44 Gerhard Coetzee¹⁷⁴, Caroline Hayward¹⁷⁵, Ian Deary¹⁷⁶, Sarah Harris¹⁷⁷, Evropi Theodoratou¹⁷⁸,
 45 Stuart Reid³, Marion Walker³, Li Yin Ooi^{179,3}, Victor Moreno^{10,11,12,13*}, Graham Casey^{19*}, Stephen
 46 Gruber^{104*}, Ian Tomlinson^{1,15*}, Wei Zheng^{5*}, Malcolm Dunlop^{3*}, Richard Houlston^{6*}, Ulrike
 47 Peters^{7,180*}

48

49 ¹Edinburgh Cancer Research Centre, Institute of Genomics and Cancer, University of Edinburgh,
 50 Edinburgh, United Kingdom, ²Genomic Medicine Group, Instituto de Investigacion Sanitaria de
 51 Santiago (IDIS), Santiago de Compostela, Spain, ³Colon Cancer Genetics Group, Medical Research
 52 Council Human Genetics Unit, Institute of Genetics and Cancer, University of Edinburgh,
 53 Edinburgh, United Kingdom, ⁴Danish Institute for Advanced Study (DIAS), Department of Public
 54 Health, University of Southern Denmark, Odense, Denmark, ⁵Division of Epidemiology,
 55 Department of Medicine, Vanderbilt-Ingram Cancer Center, Vanderbilt Epidemiology Center,
 56 Vanderbilt University Medical Center, Nashville, USA, ⁶Division of Genetics and Epidemiology,
 57 The Institute of Cancer Research, London, United Kingdom, ⁷Public Health Sciences Division, Fred

58 Hutchinson Cancer Research Center, Seattle, USA, ⁸Genomic Medicine Institute, Cleveland Clinic,
59 Cleveland, USA, ⁹Population and Cancer Prevention Program, Case Comprehensive Cancer
60 Center, Cleveland, USA, ¹⁰Colorectal Cancer Group, ONCOBELL Program, Bellvitge Biomedical
61 Research Institute (IDIBELL), Barcelona, Spain, ¹¹Oncology Data Analytics Program, Catalan
62 Institute of Oncology (ICO), Barcelona, Spain, ¹²Consortium for Biomedical Research in
63 Epidemiology and Public Health (CIBERESP). Madrid, Madrid, Spain, ¹³Department of Clinical
64 Sciences, Faculty of Medicine, University of Barcelona, Barcelona, Spain, ¹⁴Department of
65 Biostatistics, School of Public Health, University of Washington, Seattle, USA, ¹⁵Institute of Cancer
66 and Genomic Sciences, College of Medical and Dental Sciences, University of Birmingham,
67 Birmingham, United Kingdom, ¹⁶Department of Public Health, Richard Doll Building, University of
68 Oxford, Oxford, United Kingdom, ¹⁷Department of Biomedical Informatics, Vanderbilt University
69 School of Medicine, Nashville, USA, ¹⁸Department of Preventive Medicine, USC Norris
70 Comprehensive Cancer Center, Keck School of Medicine, University of Southern California, Los
71 Angeles, USA, ¹⁹Center for Public Health Genomics, Department of Public Health Sciences,
72 University of Virginia, Charlottesville, USA, ²⁰Department of Population and Quantitative Health
73 Sciences, Case Western Reserve University, Cleveland, USA, ²¹Case Comprehensive Cancer
74 Center, Case Western Reserve University, Cleveland, USA, ²²The Steve and Cindy Rasmussen
75 Institute for Genomic Medicine, Nationwide Children's Hospital, Columbus, USA, ²³Department
76 of Community Medicine and Epidemiology, Lady Davis Carmel Medical Center, Haifa, Israel,
77 ²⁴Ruth and Bruce Rappaport Faculty of Medicine, Technion-Israel Institute of Technology, Haifa,
78 Israel, ²⁵Clalit National Cancer Control Center, Haifa, Israel, ²⁶Consortium for Biomedical Research
79 in Epidemiology and Public Health (CIBERESP), Madrid, Spain, ²⁷Biomedicine Institute (IBIOMED),
80 University of León, León, Spain, ²⁸Center for Colorectal Cancer, National Cancer Center Hospital,
81 National Cancer Center, Gyeonggi-do, South Korea, ²⁹Department of Cancer Biomedical Science,
82 Graduate School of Cancer Science and Policy, National Cancer Center, Gyeonggi-do, South
83 Korea, ³⁰Department of Epidemiology and Health Promotion, Graduate School of Public Health,
84 Yonsei University, Seoul, South Korea, ³¹Department of Preventive Medicine, Chonnam National
85 University Medical School, Gwangju, South Korea, ³²Department of Preventive Medicine, Seoul
86 National University College of Medicine, Seoul, South Korea, ³³Cancer Research Institute, Seoul

87 National University, Seoul, South Korea, ³⁴Department of Social and Preventive Medicine, Hallym
88 University College of Medicine, Okcheon-dong, South Korea, ³⁵Division of Cancer Epidemiology
89 and Prevention, Aichi Cancer Center Research Institute, Nagoya, Japan, ³⁶Division of Molecular
90 and Clinical Epidemiology, Aichi Cancer Center Research Institute, Nagoya, Japan, ³⁷Department
91 of Epidemiology, Nagoya University Graduate School of Medicine, Nagoya, Japan, ³⁸Laboratory
92 of Clinical Genome Sequencing, Department of Computational Biology and Medical Sciences,
93 Graduate School of Frontier Sciences, University of Tokyo, Tokyo, Japan, ³⁹Laboratory of Genome
94 Technology, Human Genome Center, Institute of Medical Science, University of Tokyo, Tokyo,
95 Japan, ⁴⁰School of Public Health, Sun Yat-sen University, Guangzhou, China, ⁴¹State Key
96 Laboratory of Oncogenes and Related Genes and Department of Epidemiology, Shanghai Cancer
97 Institute, Renji Hospital, Shanghai Jiaotong University School of Medicine, Shanghai, China,
98 ⁴²State Key Laboratory of Oncology in South China, Cancer Center, Sun Yat-sen University,
99 Guangzhou, China, ⁴³Centre for Epidemiology and Biostatistics, Melbourne School of Population
100 and Global Health, The University of Melbourne, Melbourne, Australia, ⁴⁴Department of
101 Epidemiology, School of Public Health and Institute of Health and Environment, Seoul National
102 University, Seoul, South Korea, ⁴⁵Department of Laboratory Medicine and Pathology, Mayo Clinic
103 Arizona, Scottsdale, USA, ⁴⁶Department of Medicine, Samuel Oschin Comprehensive Cancer
104 Institute, Cedars-Sinai Medical Center, Los Angeles, USA, ⁴⁷Division of Oncology, Department of
105 Medicine, Cedars-Sinai Cancer Research Center for Health Equity, Los Angeles, USA, ⁴⁸Lunenfeld
106 Tanenbaum Research Institute, Mount Sinai Hospital, University of Toronto, Toronto, Canada,
107 ⁴⁹Memorial University of Newfoundland, Division of Biomedical Sciences, St. John's, Canada,
108 ⁵⁰School of Public Health, University of Washington, Seattle, USA, ⁵¹Translational Genomics
109 Research Institute, City of Hope National Medical Center, Phoenix, USA, ⁵²Institute of Medical
110 Genetics, Cardiff University, Cardiff, United Kingdom, ⁵³MRC Clinical Trials Unit, Medical Research
111 Council, United Kingdom, ⁵⁴MRC Institute for Radiation Oncology, University of Oxford, Oxford,
112 United Kingdom, ⁵⁵Department of Oncology, University of Oxford, Oxford, United Kingdom,
113 ⁵⁶Radcliffe Department of Medicine, University of Oxford, Oxford, United Kingdom,
114 ⁵⁷Department of Surgical Oncology, University Hospital for Tumors, Sestre milosrdnice University
115 Hospital Center, Zagreb, Croatia, ⁵⁸Department of Pathology, Central Finland Health Care District,

116 Jyväskylä, Finland, ⁵⁹Central Finland Health Care District, Central Finland Health Care District,
117 Jyväskylä, Finland, ⁶⁰Department of Health and Welfare, Finnish Institute for Health and Welfare,
118 Helsinki, Finland, ⁶¹Department of Medical and Clinical Genetics, University of Helsinki, Helsinki,
119 Finland, ⁶²Genome-Scale Biology Research Program, University of Helsinki, Helsinki, Finland,
120 ⁶³Department of Public Health and Welfare, Finnish Institute for Health and Welfare, Helsinki,
121 Finland, ⁶⁴Finnish Cancer Registry, Institute for Statistical and Epidemiological Cancer Research,
122 Helsinki, Finland, ⁶⁵Faculty of Social Sciences, Tampere University, Tampere, Finland, ⁶⁶Folkhälsan
123 Research Centre, University of Helsinki, Helsinki, Finland, ⁶⁷National University of Singapore,
124 Human Potential Translational Research Programme, Singapore, Singapore, ⁶⁸Unit of General
125 Practice and Primary Health Care, University of Helsinki and Helsinki University Hospital, Helsinki,
126 Finland, ⁶⁹Department of Surgery, Abdominal Centre, Helsinki University Hospital, Helsinki,
127 Finland, ⁷⁰University of Toronto, Department of Oncology, Toronto, Canada, ⁷¹Division of Cancer
128 Epidemiology and Genetics, National Cancer Institute, National Institutes of Health, Bethesda,
129 USA, ⁷²Department of Nutritional Sciences, School of Public Health, University of Michigan, Ann
130 Arbor, USA, ⁷³Slone Epidemiology Center at Boston University, Boston, Massachusetts, USA,
131 ⁷⁴Department of Medicine, Boston University School of Medicine, Boston, USA, ⁷⁵Colorectal
132 Oncogenomics Group, Department of Clinical Pathology, The University of Melbourne, Parkville,
133 Australia, ⁷⁶University of Melbourne Centre for Cancer Research, Victorian Comprehensive
134 Cancer Centre, Parkville, Australia, ⁷⁷Genomic Medicine and Family Cancer Clinic, The Royal
135 Melbourne Hospital, Parkville, Australia, ⁷⁸Department of Epidemiology, Johns Hopkins
136 Bloomberg School of Public Health, Baltimore, USA, ⁷⁹Huntsman Cancer Institute and
137 Department of Population Health Sciences, University of Utah, Salt Lake City, USA, ⁸⁰Cancer
138 Epidemiology Program, H. Lee Moffitt Cancer Center and Research Institute, Tampa, USA,
139 ⁸¹Institute of Cancer Research, Department of Medicine I, Medical University Vienna, Vienna,
140 Austria, ⁸²Department of Epidemiology and Population Health, Albert Einstein College of
141 Medicine, Bronx, USA, ⁸³Division of Cancer Epidemiology, German Cancer Research Center
142 (DKFZ), Heidelberg, Germany, ⁸⁴University Medical Centre Hamburg-Eppendorf, University
143 Cancer Centre Hamburg (UCCH), Hamburg, Germany, ⁸⁵Division of Clinical Epidemiology and
144 Aging Research, German Cancer Research Center (DKFZ), Heidelberg, Germany, ⁸⁶Division of

145 Preventive Oncology, German Cancer Research Center (DKFZ) and National Center for Tumor
146 Diseases (NCT), Heidelberg, Germany, ⁸⁷German Cancer Consortium (DKTK), German Cancer
147 Research Center (DKFZ), Heidelberg, Germany, ⁸⁸Department of Internal Medicine, University of
148 Utah, Salt Lake City, USA, ⁸⁹Research Centre for Hauora and Health, Massey University,
149 Wellington, New Zealand, ⁹⁰Department of Hygiene and Epidemiology, University of Ioannina
150 School of Medicine, Ioannina, Greece, ⁹¹Department of Epidemiology and Biostatistics, School of
151 Public Health, Imperial College London, London, United Kingdom, ⁹²Department of Molecular
152 Epidemiology, German Institute of Human Nutrition Potsdam-Rehbruecke, Nuthetal, Germany,
153 ⁹³Institute of Nutritional Science, University of Potsdam, Potsdam, Germany, ⁹⁴Nutrition and
154 Metabolism Branch, International Agency for Research on Cancer, World Health Organization,
155 Lyon, France, ⁹⁵Gastroenterology Department, Hospital Clínic, Institut d'Investigacions
156 Biomèdiques August Pi i Sunyer (IDIBAPS), Centro de Investigación Biomédica en Red de
157 Enfermedades Hepáticas y Digestivas (CIBEREHD), University of Barcelona, Barcelona, Spain,
158 ⁹⁶Department of Genetics, Stanford University, Stanford, USA, ⁹⁷Department of Population and
159 Public Health Sciences, USC Norris Comprehensive Cancer Center, Keck School of Medicine,
160 University of Southern California, Los Angeles, USA, ⁹⁸Jeonnam Regional Cancer Center, Chonnam
161 National University Hwasun Hospital, Hwasun, South Korea, ⁹⁹Bioinformatics and Data Science
162 Research Center, Bina Nusantara University, Jakarta, Indonesia, ¹⁰⁰Leeds Institute of Medical
163 Research at St. James's, University of Leeds, Leeds, United Kingdom, ¹⁰¹Cancer Epidemiology
164 Division, Cancer Council Victoria, Melbourne, Australia, ¹⁰²Precision Medicine, School of Clinical
165 Sciences at Monash Health, Monash University, Clayton, Australia, ¹⁰³Department of Clinical
166 Pathology, The University of Melbourne, Victoria, Australia, ¹⁰⁴Department of Medical Oncology
167 and Center For Precision Medicine, City of Hope National Medical Center, USA, ¹⁰⁵Department of
168 Pathology, University of Michigan, Ann Arbor, USA, ¹⁰⁶The Clalit Health Services, Personalized
169 Genomic Service, Lady Davis Carmel Medical Center, Haifa, Israel, ¹⁰⁷Clinical Genetics Service,
170 Department of Medicine, Memorial Sloan-Kettering Cancer Center, New York, USA,
171 ¹⁰⁸Department of Medicine, Weill Cornell Medical College, New York, USA, ¹⁰⁹Kaiser Permanente
172 Washington Health Research Institute, Seattle, USA, ¹¹⁰Center for Gastrointestinal Biology and
173 Disease, University of North Carolina, Chapel Hill, USA, ¹¹¹Department of Radiation Sciences,

174 Oncology Unit, Umeå University, Umeå, Sweden, ¹¹²Wallenberg Centre for Molecular Medicine,
175 Umeå University, Umeå, Sweden, ¹¹³Ontario Institute for Cancer Research, Toronto, Canada,
176 ¹¹⁴Division of Human Genetics, Department of Internal Medicine, The Ohio State University
177 Comprehensive Cancer Center, Columbus, USA, ¹¹⁵Division of Epidemiology, Department of
178 Population Health, New York University School of Medicine, New York, USA, ¹¹⁶Population
179 Sciences, Disparities and Community Engagement, University of California San Diego Moores
180 Cancer Center, La Jolla, USA, ¹¹⁷Department of Family Medicine and Public Health, University of
181 California San Diego, La Jolla, USA, ¹¹⁸Fred Hutchinson Cancer Research Center, Seattle, USA,
182 ¹¹⁹Division of Research, Kaiser Permanente Northern California, Oakland, USA, ¹²⁰Department of
183 Gastroenterology, Kaiser Permanente Medical Center, San Francisco, USA, ¹²¹Department of
184 Public Health and Primary Care, University of Cambridge, Cambridge, United Kingdom,
185 ¹²²Institute of Environmental Medicine, Karolinska Institutet, Stockholm, Sweden, ¹²³Taipei
186 Medical University, Taipei, Taiwan, ¹²⁴Department of Medicine, Keck School of Medicine,
187 University of Southern California, Los Angeles, USA, ¹²⁵Department of Epidemiology, University
188 of Washington School of Public Health, Seattle, USA, ¹²⁶Center for Inherited Disease Research
189 (CIDR), Department of Genetic Medicine, Johns Hopkins University School of Medicine,
190 Baltimore, USA, ¹²⁷Clinical and Translational Epidemiology Unit, Massachusetts General Hospital
191 and Harvard Medical School, Boston, USA, ¹²⁸Division of Gastroenterology, Massachusetts
192 General Hospital and Harvard Medical School, Boston, USA, ¹²⁹Channing Division of Network
193 Medicine, Brigham and Women's Hospital and Harvard Medical School, Boston, USA, ¹³⁰Broad
194 Institute of Harvard and MIT, Cambridge, USA, ¹³¹Department of Epidemiology, Harvard T.H. Chan
195 School of Public Health, Harvard University, Boston, USA, ¹³²Department of Immunology and
196 Infectious Diseases, Harvard T.H. Chan School of Public Health, Harvard University, Boston, USA,
197 ¹³³Comprehensive Cancer Center, University of Puerto Rico, San Juan, Puerto Rico, ¹³⁴Department
198 of Clinical Genetics, Karolinska University Hospital, Stockholm, Sweden, ¹³⁵Department of
199 Molecular Medicine and Surgery, Karolinska Institutet, Stockholm, Sweden, ¹³⁶Nuffield
200 Department of Population Health, University of Oxford, Oxford, United Kingdom, ¹³⁷Department
201 of General Surgery, University Hospital Rostock, Rostock, Germany, ¹³⁸Department of Genetics
202 and Genome Sciences, Case Western Reserve University, Cleveland, USA, ¹³⁹Department of

203 Computer Science, Stanford University, Stanford, USA, ¹⁴⁰Department of Genome Sciences,
204 University of Washington, Seattle, USA, ¹⁴¹Department of Medicine and Epidemiology, University
205 of Pittsburgh Medical Center, Pittsburgh, USA, ¹⁴²Department of Medicine I, University Hospital
206 Dresden, Technische Universität Dresden (TU Dresden), Dresden, Germany, ¹⁴³Department of
207 Medicine, Memorial Sloan-Kettering Cancer Center, New York, USA, ¹⁴⁴Department of Molecular
208 Biology of Cancer, Institute of Experimental Medicine of the Czech Academy of Sciences, Prague,
209 Czech Republic, ¹⁴⁵Institute of Biology and Medical Genetics, First Faculty of Medicine, Charles
210 University, Prague, Czech Republic, ¹⁴⁶Faculty of Medicine and Biomedical Center in Pilsen,
211 Charles University, Pilsen, Czech Republic, ¹⁴⁷Department of Oncology Ludwig Center at the
212 Sidney Kimmel Cancer Center, Johns Hopkins University School of Medicine, Baltimore, USA,
213 ¹⁴⁸Sol Goldman Pancreatic Cancer Research Center, Johns Hopkins University School of Medicine,
214 Baltimore, USA, ¹⁴⁹Department of Pathology, Johns Hopkins School of Medicine, Baltimore, USA,
215 ¹⁵⁰Department of Surgery, University of Tennessee Health Science Center, Memphis, USA,
216 ¹⁵¹Departments of Cancer Biology and Genetics and Internal Medicine, Comprehensive Cancer
217 Center, The Ohio State University, Columbus, USA, ¹⁵²Departments of Medicine and Genetics,
218 Case Comprehensive Cancer Center, Case Western Reserve University and University Hospitals
219 of Cleveland, Cleveland, USA, ¹⁵³Division of Human Nutrition and Health, Wageningen University
220 & Research, Wageningen, The Netherlands, ¹⁵⁴Division of Human Nutrition, Wageningen
221 University and Research, Wageningen, The Netherlands, ¹⁵⁵Genomic Medicine Group, Galician
222 Public Foundation of Genomic Medicine, Servicio Galego de Saude (SERGAS), Santiago de
223 Compostela, Spain, ¹⁵⁶Instituto de Investigación Sanitaria de Santiago de Compostela (IDIS),
224 Santiago de Compostela, Spain, ¹⁵⁷Italian Institute for Genomic Medicine (IIGM), Candiolo Cancer
225 Institute - FPO-IRCCS, Turin, Italy, ¹⁵⁸Candiolo Cancer Institute - FPO-IRCCS, Candiolo, Italy,
226 ¹⁵⁹Menzies Institute for Medical Research, University of Tasmania, Hobart, Australia, ¹⁶⁰National
227 University Cancer Institute, Singapore; Cancer Science Institute of Singapore, National University
228 of Singapore, Singapore, Singapore, ¹⁶¹Department of Pathology, Brigham and Women's Hospital,
229 Harvard Medical School, Boston, USA, ¹⁶²Cancer Immunology Program, Dana-Farber Harvard
230 Cancer Center, Boston, USA, ¹⁶³Broad Institute of MIT and Harvard, Cambridge, USA,
231 ¹⁶⁴Department of Biostatistics, University of Washington, Seattle, USA, ¹⁶⁵Service de Génétique

232 Médicale, Centre Hospitalier Universitaire (CHU) Nantes, Nantes, France, ¹⁶⁶SWOG Statistical
 233 Center, Fred Hutchinson Cancer Research Center, Seattle, USA, ¹⁶⁷Department of Pediatrics,
 234 Nationwide Children's Hospital, The Steve and Cindy Rasmussen Institute for Genomic Medicine,
 235 Columbus, USA, ¹⁶⁸Division of Epidemiology, National Cancer Center Institute for Cancer Control,
 236 National Cancer Center, Tokyo, Japan, ¹⁶⁹Division of Cohort Research, National Cancer Center
 237 Institute for Cancer Control, National Cancer Center, Tokyo, Japan, ¹⁷⁰Department of Preventive
 238 Medicine, Center for Genetic Epidemiology, University of Southern California, Los Angeles, USA,
 239 ¹⁷¹Cancer Center, University of Hawaii, Honolulu, USA, ¹⁷²Preventative Medicine, University of
 240 Southern California, Los Angeles, USA, ¹⁷³USC Norris Comprehensive Cancer Center, Keck School
 241 of Medicine, University of Southern California, Los Angeles, USA, ¹⁷⁴Van Andel Research Institute,
 242 Grand Rapids, USA, ¹⁷⁵MRC Human Genetics Unit, Institute of Genomics and Cancer, University
 243 of Edinburgh, Edinburgh, United Kingdom, ¹⁷⁶Lothian Birth Cohorts group, Department of
 244 Psychology, University of Edinburgh, Edinburgh, United Kingdom, ¹⁷⁷Lothian Birth Cohorts group,
 245 Department of Psychology, University of Edinburgh, Edinburgh EH8 9JZ, United Kingdom,
 246 ¹⁷⁸Centre for Global Health, Usher Institute, University of Edinburgh, Edinburgh, United Kingdom,
 247 ¹⁷⁹Department of Pathology, National University Hospital, National University Health System,
 248 Singapore, Singapore, ¹⁸⁰Department of Epidemiology, University of Washington, Seattle, USA

249 ^These authors contributed equally

250 *These authors jointly supervised this work

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252 Corresponding authors: Ian PM Tomlinson (Ian.Tomlinson@ed.ac.uk); Wei Zheng
 253 (wei.zheng@vumc.org); Malcolm G Dunlop (malcolm.dunlop@ed.ac.uk); Richard S Houlston
 254 (Richard.Houlston@icr.ac.uk); Ulrike Peters (upeters@fredhutch.org).

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256

257 **ABSTRACT**

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259 **Colorectal cancer (CRC) is a leading cause of mortality worldwide. We conducted a genome-**
260 **wide association study meta-analysis of 100,204 CRC cases and 154,587 controls of European**
261 **and East Asian ancestry, identifying 205 independent risk associations, of which 50 were**
262 **unreported. We performed integrative genomic, transcriptomic and methylomic analyses**
263 **across large bowel mucosa and other tissues. Transcriptome- and methylome-wide association**
264 **studies revealed an additional 53 risk associations. We identified 155 high confidence effector**
265 **genes functionally linked to CRC risk, many of which had no previously established role in CRC.**
266 **These have multiple different functions, and specifically indicate that variation in normal**
267 **colorectal homeostasis, proliferation, cell adhesion, migration, immunity and microbial**
268 **interactions determines CRC risk. Cross-tissue analyses indicated that over a third of effector**
269 **genes most likely act outside the colonic mucosa. Our findings provide insights into colorectal**
270 **oncogenesis, and highlight potential targets across tissues for new CRC treatment and**
271 **chemoprevention strategies.**

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281 **INTRODUCTION**

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283 Colorectal cancer (CRC), which affects approximately 1.9 million people worldwide annually¹, has
284 a strong heritable basis². Our understanding of CRC genetics has been informed by genome-wide
285 association studies (GWAS), which have so far identified 150 statistically independent risk
286 variants^{3,4}. To provide a comprehensive description of CRC genetics, we brought together the
287 great majority of GWAS performed to date. We complemented GWAS with transcriptome- and
288 methylome-wide association analyses (TWAS and MWAS; **Fig. 1**). Through integration of these
289 data, we investigated the genes and mechanisms underlying established and novel CRC risk loci.
290 We identified credible effector genes and the tissues in which they act, informing our
291 understanding of colorectal tumorigenesis.

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294 **RESULTS**

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296 **Genetic architecture of colorectal cancer**

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298 We performed a meta-analysis of CRC GWAS data sets, comprising 100,204 CRC cases and
299 154,587 controls (73% European and 27% East Asian ancestry) (**Supplementary Tables 1 & 2**).
300 We identified 205 associations, including 37 single-nucleotide polymorphisms (SNPs) at novel loci
301 (sentinel risk SNPs > 1 megabase (Mb) from another significant SNP), 13 independent novel risk
302 SNPs in conditional analysis (**Table 1**), and 155 previously reported SNPs or proxies **Table 1**,
303 **Supplementary Tables 3-4, Supplementary figures 1 & 2**). There was limited heterogeneity
304 ascribable to population effects (**Supplementary Table 2, Supplementary figure 3**), although four
305 risk variants (rs12078075, rs57939401, rs151127921 and rs5751474) were monomorphic in East
306 Asian participants (Table 1).

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308

309 Using linkage-disequilibrium (LD) score regression (LD hub), we estimated the heritability of CRC
310 attributable to all common genetic variants to be similar in Europeans (h^2 0.11, s.d. 0.008) and
311 East Asians (h^2 0.09, s.d. 0.006), which translates to 73% of familial CRC risk. Restricting estimates
312 to the 205 GWAS-significant SNPs explained 19.7% of this familial risk. We evaluated the
313 performance of a polygenic risk score (PRS) based on these SNPs in two cohorts independent of
314 the GWAS discovery samples^{7,8}. For Europeans and East Asians, individuals in the top PRS decile
315 exhibited odds ratios of 2.22 (95%CI: 1.92-2.57; $P = 1.80 \times 10^{-26}$) and 1.96 (95%CI: 1.64-2.34; $P =$
316 8.9×10^{-14}) compared to the remaining individuals. Corresponding areas under the receiver
317 operating characteristic curve (AUC) were 0.62 (95%CI: 0.60-0.63) and 0.60 (95%CI: 0.59-0.62).

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320 **Discovery of risk loci by TWAS and MWAS**

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322 TWAS was performed by implementing the PredictDB pipeline using mRNA expression data from
323 1,107 colorectal mucosa samples as reference (709 in house, 368 GTEx transverse colon)^{9,10}. In
324 addition to associations identified by GWAS or those previously reported by TWAS (*PYGL* and
325 *TRIM4*^{11,12}), we identified 15 novel associations at Bonferroni-corrected significance ($P_{\text{Bonferroni}}$,
326 **Table 2, Supplementary Tables 5 & 6, Supplementary figure 4**). We extended the main TWAS to
327 a transcript isoform-wide association study (TIsWAS), both to ascertain whether specific
328 transcripts could account for TWAS associations and to identify previously unreported risk
329 associations (**Supplementary Tables 7 & 8**). For a third of TWAS genes, a significant association
330 with CRC risk was found for a single mRNA isoform (**Supplementary Table 7**). The TIsWAS also
331 identified eight loci associated with CRC risk (**Table 3**). To improve power for discovery, and
332 because some CRC risk SNPs may not exert their effects in colorectal mucosa, we also conducted
333 a cross-tissue TWAS using our in-house RNA sequencing (RNAseq) data and the full GTEx and
334 Depression Genes and Networks (DGN) project data (49 tissues)¹³. We identified a further 23 risk
335 associations (**Table 4, Supplementary Tables 9-13**).

336

337 To complement the TWAS, identify further CRC risk loci and gain mechanistic insights, we
338 extended the PredictDB pipeline to perform MWAS based on quantitative methylation data from
339 histologically normal colorectal mucosa (**Supplementary Methods**). We found significant
340 associations between CRC risk and methylation of individual CpGs at 69 loci (**Supplementary**
341 **Tables 14 & 15**). This included seven novel independent risk loci (**Table 5**). Risk SNPs may
342 influence CRC risk through changes in the CpG methylation status of regulatory elements leading
343 to changes in gene expression. We therefore explored the relationship between gene expression,
344 CpG methylation and CRC risk in colorectal mucosa for 6,722 genes with both TWAS and MWAS
345 predictions. There was a strong tendency for genes to be represented in both TWAS and MWAS
346 ($P < 10^{-7}$, Fisher's exact test). Subsequently, we conditioned TWAS associations on the top MWAS-
347 significant CpG within 1Mb, finding that 67/91 (75%) genes did not retain a significant TWAS
348 association ($P_{\text{Bonferroni}} > 5.50 \times 10^{-4}$; **Supplementary Table 16**). Our data are consistent with a
349 model in which many CRC risk SNPs act through changes in DNA methylation, although formal
350 causality analysis could not be performed to exclude reverse causation or possible confounders.

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352

353 **Effector genes and biological pathways of CRC oncogenesis**

354

355 A major, largely unfulfilled aim of cancer GWAS is to identify genes and functional mechanisms
356 that may ultimately be clinically useful targets, for example in chemoprevention. The large GWAS
357 and TWAS datasets in this study address this aim by enabling a detailed functional analysis of the
358 molecular mechanisms contributing to CRC risk. Since TWAS approaches do not identify causal
359 genes directly, we used our data to compile a set of 155 credible effector genes from the
360 independent associations identified through GWAS, TWAS, TisWAS and MWAS (details in
361 **Supplementary Table 17** and **Supplementary Methods**).

362

363 We identified molecular pathways enriched in effector genes using Enrichr
364 (<https://maayanlab.cloud/Enrichr/>) (**Supplementary Table 18**). This analysis was complemented
365 with DEPICT based on the GWAS SNPs (<https://data.broadinstitute.org/mpg/depict/>)

366 **(Supplementary Table 19)**. CRC effectors were principally enriched in genes regulating TGF-
 367 β /BMP, Wnt WNT and Hippo pathways. A number of the credible effector genes that map to
 368 these pathways have no established role in CRC, including the intestinal stem cell regulator
 369 *ZNRF3*¹⁴, the TGF repressor *LEMD3*¹⁵, and the EMT regulator *RREB1*¹⁶.

370
 371 To complement the pathway analysis, we performed gene-level functional annotation based on
 372 the principal cellular function of each effector gene as reported in the literature (**Figure 2**,
 373 **Supplementary Table 20**). Thirty-six genes (mostly Wnt and BMP family members) were
 374 annotated to colorectal homeostasis (i.e. cellular stemness/differentiation). Intriguingly, 16
 375 genes (including *ARHGEF19*, *ARHGEF4*, *GNA12*, *RHOG*, *TAGLN*, *TSPAN8*, *STARD13* and *LLGL1*)
 376 were linked to cell migration through RhoA/ROCK signaling. We found eight genes (*SPSB1*,
 377 *PIK3C2B*, *DUSP1*, *LRIG1*, *GAB1*, *RREB1*, *MAPKAPK5-AS1* and *PDGFB*) to act within the Ras/Raf
 378 growth factor signaling pathway. In addition to the previously reported association at *FUT2*, the
 379 novel fucosyltransferase effector genes *FUT3* and *FUT6* supported a relationship between the gut
 380 microbiome and CRC risk¹⁷. Inflammation is important in CRC¹⁸, and the TWAS association at the
 381 *FADS* gene cluster and *PTGES3*, specifically highlighted the role of prostaglandin metabolism in
 382 CRC risk. Finally, our data also indicated several effector genes with roles in ion transport and
 383 cytoskeletal components (**Fig. 2, Supplementary Table 20**).

384
 385 Although our pathway analysis and functional annotation indicated that the colorectum was the
 386 likely target tissue of many effector genes (**Supplementary Tables 19 & 20**), some genes were
 387 associated with principal roles in other tissue types, for example neuronal cells (*LINGO4*, *TULP1*
 388 and *CNIH2*) and leukocytes (*TOX*, *TOX4* and *MAF*, plus many candidate genes within the MHC
 389 region) (**Supplementary Table 20**). We therefore performed a systematic analysis of effector
 390 gene tissue specificity, based on the premise that TWAS associations tend to be present in tissues
 391 in which a gene functionally affects CRC risk. Cross-tissue analysis showed that all but one
 392 effector gene exhibited a TWAS association ($FDR_{TWAS} < 0.05$) in at least one tissue and 52 (34%)
 393 genes showed an association in multiple tissues (**Supplementary figure 5**). For 26 (17%) genes,
 394 associations were confined to the colorectal mucosa (P_{TWAS} Bonferroni-significant in mucosa,

395 $P_{\text{TWAS}} > \text{FDR}$ elsewhere). In contrast, 67 genes (43%) showed no evidence of a TWAS association
396 in colorectal mucosa ($\text{FDR}_{\text{TWAS}} > 0.05$). Notably, 12 (8%) gene associations were present only in
397 immune cells (**Supplementary figure 5, Supplementary Table 11**) and four (3%) were restricted
398 to mesenchymal cells (**Supplementary figure 5, Supplementary Table 12**).

399

400 **Linking colorectal cancer risk to other traits**

401

402 To gain insight into the role of potentially modifiable risk factors in CRC genetics, we performed
403 cross-trait LD score regression analyses¹⁹ using publicly available GWAS summary statistics for
404 171 phenotypes. Twelve genetic correlations remained significant (two-sided Z-test, Bonferroni-
405 corrected $P < 2.93 \times 10^{-4}$). Notably, positive associations with CRC risk (**Supplementary Table 21**)
406 included insulin resistance (raised fasting insulin and glucose), smoking, and obesity (body mass
407 index - BMI, waist-to-hip ratio - WHR, waist circumference), traits that have previously
408 been reported in observational epidemiological studies to be associated with CRC risk^{3,20,21}. These
409 associations not only highlight shared biology, but also suggest that public health interventions
410 to reduce cardiometabolic disease will additionally lower CRC burden.

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412

413 **DISCUSSION**

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415 We report a comprehensive genetic analysis of CRC risk in the general population. To identify the
416 most credible effector genes for each risk variant, we performed detailed annotation using tissue-
417 specific gene expression and other relevant data types. Our study is twice as large as previous
418 CRC GWAS, and also includes participants of both European and East Asian ancestries,
419 demonstrating that most loci are shared across these ancestral groups. This increased power for
420 GWAS, coupled with complementary analyses, including TWAS and MWAS, identified 103
421 previously unreported risk associations and identified 155 effector genes. These data
422 substantially expand our existing knowledge regarding the impact of common genetic variation
423 on the heritable risk of CRC.

424

425 The availability of large, multi-omic data sets has allowed us to assign the most likely
426 target/effector genes of GWAS and TWAS associations (**Fig. 3**), and confidence in these
427 assignments will increase as additional functional data are reported in the literature. It is clear
428 that pathways (*e.g.*, Wnt , BMP, Hippo) involved in normal intestinal homeostasis play
429 important roles in CRC risk, suggesting that modulation of normal mucosal dynamics has the
430 potential to prevent colorectal neoplasia. The gut flora is intimately involved in normal bowel
431 homeostasis, and effector genes are likely to be involved in microbial interactions. By contrast,
432 Ras pathway activity is thought to be more important during repair or tumorigenesis, and the Ras
433 effector genes we have found may act after tumor initiation. Our finding of multiple risk genes
434 involved in cell adhesion and migration naturally suggests roles in malignant progression,
435 although effects earlier in tumorigenesis also remain plausible. Similarly, immune pathway
436 effector genes could, in principle, have their effects on normal cell function or at any stage of
437 tumorigenesis, from mediating day-to-day microbial interactions to killing of cells in early
438 neoplastic transformation or established tumors.

439

440 Cross-tissue analyses indicated that the colorectal mucosa was the most likely site of action of
441 many effector genes, but some genes are more likely to act in different tissue types. For example,
442 it is highly likely that genes such as *HIVEP1*, *LIF*, *SH2B3*, *TOX* and *TOX4* (and probably genes in the
443 MHC region) influence the development of CRC through immune cell variation, and that *EDNRB*
444 influences risk through effects on blood vessels. An unexpected finding was that several credible
445 effector genes have primary roles in neurogenesis, raising the intriguing possibility that the
446 enteric nervous system is involved in CRC risk.

447

448 While germline genetics has guided the development of drugs to prevent cardiovascular disease
449 (*e.g.* statins and PCSK9 inhibitors), such a paradigm has yet to be realized for cancer. Since almost
450 all CRCs develop from colonic polyps, and up to 40% of the screened population will be diagnosed
451 with one or more polyps, CRC is particularly well-suited to evaluate novel chemopreventive
452 agents. Our findings highlight candidate targets for chemoprevention, such as gut microbiota,

453 prostaglandin metabolism, and signaling through the Wnt ~~WNT~~, BMP and Hippo pathways.
454 Specific potential targets in the near term include CDK6, which is targeted by drugs in clinical use
455 for cancer therapy, such as palbociclib and ribociclib. Similarly, Wnt ~~WNT~~ pathway activity can
456 be targeted indirectly using porcupine inhibitors (e.g. LGK974, ETC159, CGX-1321 and RXC004)
457 that prevent Wnt ~~WNT~~ ligand palmitoylation²², although future approaches may more specifically
458 target effector genes such as *WNT4* and *ZNRF3*. Hence, adapted forms of these drugs or modified
459 dosing regimens could be repurposed for chemoprevention, possibly initially for high-risk groups,
460 such as those with in the top PRS percentiles or Lynch Syndrome cases. Based on our data, we
461 speculate that in the longer term, targeted approaches based on demethylation of specific CpG
462 sites from MWAS could be effective means of prevention with minimal toxicity.

463
464 The identification of additional risk associations has the potential to provide further biological
465 insights into CRC. However, cohort numbers required in European and East Asian populations to
466 identify additional risk SNPs through GWAS are likely to be prohibitive. Indeed, to identify SNPs
467 explaining 80% of the heritable risk of CRC risk loci, thus providing comprehensive biological
468 insights, will require sample sizes in excess of 500,000 cases and at least that number of controls
469 (**Supplementary figure 6**). This is far higher than a previous estimate²³, which was based on a
470 small subset of the GWAS included herein. Extending GWAS to African and other populations
471 may detect further risk SNPs, including population specific ones. Complementary approaches
472 such as TWAS and MWAS are demonstrably useful for the discovery of further risk loci, especially
473 if, and when, reference data sets from multiple populations are made available.

474
475 Overall, our findings demonstrate the power of multi-omics to provide new insights into the
476 biological basis of CRC, including both the identification of candidate effector genes and support
477 for previously unsuspected functional mechanisms. Importantly, several of the genes and
478 pathways we have identified are potential targets for CRC treatment or chemoprevention.

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528 **Author contributions**

529 Study design: CFR, MNT, PJJ, VM, GC, SBG, IT, WZ, MGD, RSH, UP; Patient recruitment and sample
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549

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 562 The remaining authors declare no competing interests.

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565

567 TABLES

568 **Table 1. Previously unreported colorectal cancer risk associations identified by genome-wide association study analysis.** *P*-values
 569 calculated from a fixed-effects meta-analysis; *, conditional SNP association, with *P*-values and ORs derived from analysis conditional
 570 on known risk loci within 1Mb; RAF, risk allele frequency; EUR, European ancestry population; EAS, East Asian ancestry population;
 571 OR, odds ratio; I^2 , fraction of variance attributable to between study heterogeneity; bp, base pairs. Association statistics for European
 572 and East Asian populations are detailed in Supplementary Table 3.

SNP	Cytoband	Position (bp, GRCh37)	Risk/Alt Allele	RAF (EUR)	RAF (EAS)	OR (95% CI)	<i>P</i> -value	I^2 (%)	Closest gene (RefSeq)
rs34963268 *	1p36.12	22,710,877	G/C	0.84	0.77	1.07 (1.05-1.09)	6.28E-16	31	<i>ZBTB40</i>
rs5028523	1q24.3	172,864,224	A/G	0.53	0.05	1.04 (1.03-1.06)	1.44E-08	0	<i>TNFSF18</i>
rs12137232	1q32.1	201,885,446	G/T	0.52	0.19	1.04 (1.03-1.05)	7.71E-09	15	<i>LMOD1</i>
rs12078075	1q32.1	205,163,798	G/A	0.09	0	1.07 (1.05-1.10)	1.94E-08	0	<i>DSTYK</i>
rs2078095	1q43	240,408,346	G/A	0.28	0.23	1.04 (1.03-1.06)	2.08E-08	0	<i>FMN2</i>
rs4668039	2q24.3	169,025,379	G/A	0.2	0.52	1.04 (1.03-1.06)	3.32E-08	12	<i>STK39</i>
rs704417	3p14.1	64,252,424	T/C	0.51	0.89	1.05 (1.03-1.06)	4.35E-10	0	<i>PRICKLE2</i>
rs7623129 *	3p14.1	64,624,426	C/T	0.56	0.51	1.04 (1.02-1.05)	1.51E-08	5	<i>ADAMTS9</i>
rs2388976	4q26	115,502,406	A/G	0.44	0.45	1.04 (1.02-1.05)	1.75E-08	17	<i>UGT8</i>
rs10006803	4q31.3	151,501,208	C/G	0.5	0.45	1.04 (1.02-1.05)	2.58E-08	0	<i>LRBA</i>
rs1426947	4q34.1	175,420,523	T/C	0.42	0.66	1.04 (1.03-1.05)	7.48E-10	0	<i>HPGD</i>
rs3930345	5q14.3	82,881,255	C/T	0.8	0.75	1.05 (1.03-1.06)	6.82E-09	10	<i>VCAN</i>

rs472959	5q35.1	172,324,558	A/G	0.46	0.46	1.04 (1.03-1.05)	4.71E-09	24	<i>ERGIC1</i>
rs1294437	6p25.1	6,749,789	C/T	0.65	0.23	1.04 (1.03-1.06)	1.21E-08	0	<i>LY86</i>
rs9379084 *	6p24.3	7,231,843	G/A	0.88	0.8	1.07 (1.05-1.09)	1.79E-12	9	<i>RREB1</i>
rs209142 *	6p22.1	28,862,617	C/G	0.39	0.52	1.04 (1.02-1.05)	3.66E-08	20	<i>TRIM27</i>
rs57939401	6p21.1	45,572,071	A/G	0.1	0.13	1.07 (1.04-1.09)	3.51E-10	0	<i>RUNX2</i>
rs6912214 *	6p12.1	55,721,302	T/C	0.55	0.83	1.04 (1.03-1.05)	1.55E-08	20	<i>BMP5</i>
rs145997965 *	6q21	106,482,613	C/T	0.02	0	1.21 (1.13-1.29)	1.26E-08	0	<i>PRDM1</i>
rs6911915	6q22.1	117,809,031	C/T	0.44	0.43	1.05 (1.03-1.06)	3.99E-12	3	<i>DCBLD1</i>
rs151127921	6q23.2	133,993,925	T/C	0.02	0	1.17 (1.11-1.24)	3.19E-08	24	<i>EYA4</i>
rs1182197	7p22.2	2,863,289	A/C	0.63	0.7	1.04 (1.03-1.05)	5.32E-09	0	<i>GNA12</i>
rs12539962	7q11.23	73,167,259	C/T	0.72	0.63	1.04 (1.03-1.05)	2.96E-08	27	<i>ABHD11</i>
rs2527927	7q22.1	99,477,426	G/A	0.55	0.71	1.04 (1.03-1.06)	3.31E-10	2	<i>OR2AE1</i>
rs60911071	8p21.2	23,664,632	G/C	0.95	0.64	1.06 (1.04-1.09)	2.24E-08	0	<i>STC1</i>
rs826732	8q12.1	59,742,639	C/G	0.5	0.59	1.04 (1.03-1.06)	6.26E-10	7	<i>TOX</i>
rs11557154	9p13.3	34,107,505	T/C	0.14	0.59	1.05 (1.04-1.07)	6.02E-10	14	<i>DCAF12</i>
rs10978941	9q31.2	110,373,819	C/T	0.83	0.87	1.06 (1.04-1.08)	2.29E-12	0	<i>KLF4</i>
rs7038489 *	9q34.2	136,682,468	C/T	0.89	0.99	1.08 (1.05-1.1)	1.1E-08	48	<i>VAV2</i>
rs11789898	9q34.2	136,925,663	T/G	0.18	0.08	1.05 (1.04-1.07)	6.28E-09	36	<i>BRD3</i>
rs1775910 *	10p12.1	29,096,942	G/C	0.25	0.32	1.04 (1.03-1.06)	3.11E-08	17	<i>LOC100507605</i>
rs1773860	10p12.1	29,291,556	T/C	0.49	0.35	1.04 (1.03-1.05)	3.49E-09	6	<i>LOC100507605</i>

rs10751097	11q13.3	69,938,433	A/G	0.4	0.31	1.05 (1.03-1.06)	2.14E-12	0	<i>ANO1</i>
rs497916	11q23.3	118,758,089	T/C	0.28	0.17	1.04 (1.03-1.06)	3.37E-08	0	<i>CXCR5</i>
rs7297628	12q14.2	64,404,555	T/C	0.54	0.75	1.04 (1.03-1.05)	1.39E-08	30	<i>SRGAP1</i>
rs11178634	12q21.1	71,518,329	G/T	0.62	0.7	1.05 (1.03-1.06)	1.36E-11	34	<i>TSPAN8</i>
rs7299936 *	12q24.21	115,934,000	A/G	0.56	0.18	1.04 (1.02-1.05)	3.73E-08	0	<i>MED13L</i>
rs116964464	13q12.13	27,543,193	T/C	0.03	0.04	1.11 (1.07-1.15)	4.83E-09	3	<i>USP12</i>
rs1078563 *	13q34	110,352,851	G/C	0.33	0.28	1.04 (1.03-1.05)	1.53E-08	0	<i>IRS2</i>
rs1497077	14q22.1	52,491,655	C/T	0.66	0.76	1.04 (1.03-1.06)	3.64E-08	0	<i>NID2</i>
rs8031386	15q23	72,508,799	A/C	0.26	0.54	1.04 (1.03-1.06)	4.50E-09	12	<i>PKM2</i>
rs11247566 *	17p13.3	835,371	G/A	0.55	0.52	1.04 (1.02-1.05)	2.92E-08	35	<i>NXN</i>
rs1791373	18p11.31	3,616,779	T/A	0.43	0.14	1.04 (1.03-1.06)	1.13E-08	0	<i>DLGAP1</i>
rs10409772	19p13.3	5,840,926	A/C	0.09	0.29	1.07 (1.05-1.09)	1.33E-10	6	<i>FUT6</i>
rs9983528	21q22.3	47,772,439	A/G	0.13	0.24	1.07 (1.05-1.09)	5.10E-13	0	<i>PCNT</i>
rs4616575	22q12.1	29,406,076	T/G	0.52	0.56	1.04 (1.03-1.05)	1.49E-10	0	<i>ZNRF3</i>
rs130651	22q13.1	39,644,273	G/A	0.33	0.08	1.05 (1.03-1.07)	2.92E-10	46	<i>PDGFB</i>
rs5751474	22q13.2	43,689,542	A/G	0.79	0	1.05 (1.03-1.07)	1.80E-08	52	<i>SCUBE1</i>
rs34256596 *	22q13.2	43,778,431	A/G	0.26	0.4	1.05 (1.03-1.06)	5.86E-09	0	<i>MPPED1</i>
rs9330814 *	22q13.31	46,364,191	T/C	0.33	0.68	1.05 (1.03-1.07)	1.28E-09	33	<i>WNT7B</i>

574 **Table 2. Colorectal cancer risk associations identified by a colorectal mucosa-specific transcriptome-wide association study.**
575 SMultiXcan uses a two-sided F-test to quantify the significance of the joint fit of the linear regression of the phenotype on predicted
576 expression from multiple tissue models jointly. All associations shown were transcriptome-wide significant after Bonferroni
577 correction for 12,017 genes with an S-MultiXcan model (*i.e.* $P = 0.05/12,017 = 4.16 \times 10^{-6}$ for the $P_{S\text{-MultiXcan}}$). Genes with boundaries
578 less than 1Mb apart were considered to be in the same cluster. This resulted in 13 CRC associations, for which all TWAS-significant
579 genes were > 1 Mb away from and independent of any GWAS-significant SNP ($P_{GWAS} < 5 \times 10^{-8}$) As expected SNPs close to genome-
580 wide significance were found in all cases. Two further gene associations (*) were < 1Mb from a GWAS-significant SNP, but in analysis
581 conditional on the SNP showed a minimally changed association (**Supplementary Table 6**) and remained significant at $P = 4.16 \times 10^{-6}$.
582 # indicates the number of novel TWAS loci. z score and effect size are calculated as the mean across S-PrediXcan models from the
583 TWAS reference data sets. n models shows the number of reference data sets for which the S-PrediXcan elastic nets produced
584 genetically-predicted expression models, with the n indep showing the number of those models that were statistically independent.
585 The SNP with the lowest CRC GWAS P -value within 1Mb of the gene is also shown.
586

#	ENSEMBL identifier	Gene	Chr	Start (bp, GRCh37)	End (bp, GRCh37)	$P_{S\text{-MultiXcan}}$	Mean z score	Effect size	n models	n indep	Top GWAS SNP at <1Mb	SNP position	P_{GWAS}
1	ENSG00000171621	<i>SPSB1</i>	1	9,352,939	9,429,591	2.96E-06	4.569	0.077	3	1	rs2075971	9,407,104	1.96E-07
2	ENSG00000142632	<i>ARHGEF19</i>	1	16,524,712	16,539,104	2.32E-06	-4.610	-0.046	7	1	rs2132851	16,537,752	7.20E-07
	ENSG00000237276	<i>ANO7P1</i>	1	16,542,404	16,554,522	1.27E-06	-4.801	-0.054	3	1	rs2132851	16,537,752	7.20E-07
3*	ENSG00000237190	<i>CDKN2AIPNL</i>	5	133,737,778	133,747,589	1.37E-09	1.665	0.045	3	3	rs647161	134,499,092	8.53E-18
4	ENSG00000260653	<i>RP11-114G11.5</i>	7	57,404,172	57,419,535	1.37E-06	-4.829	-0.494	1	1	rs4242307	57,477,102	2.28E-03
5	ENSG00000204175	<i>GPRIN2</i>	10	46,994,087	47,005,643	3.38E-14	-7.582	-1.709	1	1	rs10906949	47,698,776	1.58E-04
6	ENSG00000180210	<i>F2</i>	11	46,740,730	46,761,056	2.80E-07	5.136	0.257	1	1	rs7109707	46,818,814	5.30E-07

	ENSG00000123444	<i>KBTBD4</i>	11	47,595,014	47,600,561	5.48E-07	5.008	0.053	1	1	rs7109707	46,818,814	5.30E-07
7	ENSG00000213445	<i>SIPA1</i>	11	65,405,568	65,418,401	2.81E-06	-3.033	-0.046	2	2	rs570760	65,833,631	2.88E-07
8	ENSG00000166106	<i>ADAMTS15</i>	11	130,318,869	130,346,532	3.86E-06	4.515	0.125	2	2	rs7936386	130,462,505	9.18E-08
9	ENSG00000174106	<i>LEMD3</i>	12	65,563,351	65,642,107	2.15E-06	3.040	0.076	3	3	rs59829994	65,560,831	1.39E-07
10*	ENSG00000234608	<i>MAPKAPK5-AS1</i>	12	112,277,588	112,280,706	6.15E-14	3.544	0.050	6	6	rs653178	112,007,756	2.51E-24
11	ENSG00000167173	<i>C15orf39</i>	15	75,487,984	75,504,510	2.14E-07	4.036	0.100	3	2	rs17338413	75,474,936	2.15E-07
	ENSG00000260274	<i>RP11-817O13.8</i>	15	75,660,496	75,661,925	2.93E-06	3.090	0.096	2	2	rs17338413	75,474,936	2.15E-07
12	ENSG00000166822	<i>TMEM170A</i>	16	75,476,952	75,499,395	1.05E-06	-3.464	-0.041	7	4	rs4888408	75,432,824	9.14E-07
13	ENSG00000131748	<i>STARD3</i>	17	37,793,318	37,819,737	8.11E-07	4.933	0.143	1	1	rs2313171	37,833,842	2.77E-07
	ENSG00000161395	<i>PGAP3</i>	17	37,827,375	37,853,050	9.59E-07	4.777	0.043	7	1	rs2313171	37,833,842	2.77E-07
	ENSG00000141736	<i>ERBB2</i>	17	37,844,361	37,886,606	2.96E-06	2.679	0.032	3	3	rs2313171	37,833,842	2.77E-07
14	ENSG00000152217	<i>SETBP1</i>	18	42,260,138	42,648,475	3.11E-07	4.339	0.093	2	2	rs12958322	42,309,786	2.60E-07
15	ENSG00000267100	<i>ILF3-AS1</i>	19	10,762,538	10,764,520	2.70E-07	4.689	0.079	2	2	rs10408721	10,758,319	5.71E-08

588 **Table 3. Colorectal cancer risk associations identified by a colorectal mucosa-specific transcript isoform-wide association study**
589 **(TisWAS).** As per Table 2, S-MultiXcan uses a two-sided F-test to quantify the significance of the joint fit of the linear regression of the
590 phenotype on predicted expression from multiple tissue models jointly. All associations shown were transcriptome-wide significant
591 after Bonferroni correction for 27,941 transcripts with an S-MultiXcan model (*i.e.* $P = 0.05/27,941 = 1.79 \times 10^{-6}$ for the $P_{S\text{-MultiXcan}}$). Novel
592 associations were called when >1Mb from both a GWAS-significant SNP and a TWAS locus. As expected, all these loci showed evidence
593 of a risk association in the full TWAS ($FDR < 0.05$, $P < 2.86 \times 10^{-3}$). Transcripts with boundaries < 1 Mb apart were considered to be in
594 the same cluster. This resulted in seven CRC associations. One further association (*) was identified based on conditional TisWAS
595 analysis (**Supplementary Table 8**). Other annotations are as per **Table 2**.
596

#	ENSEMBL identifier	Gene	Chr	Start (bp, GRCh37)	End (bp, GRCh37)	$P_{S\text{-MultiXcan}}$	Mean z score	Effect size	n models	n indep	Top GWAS SNP at <1Mb	SNP location	P_{GWAS}
1	ENST00000609196	ACP6	1	147,101,453	147,131,116	6.43E-11	-1.264	-0.048	4	3	rs1541187	147,051,493	1.44E-04
	ENST00000493129	ACP6	1	147,127,341	147,142,574	1.65E-23	-5.781	-0.482	2	2	rs1541187	147,051,493	1.44E-04
2	ENST00000273153	CSRNP1	3	39,183,346	39,195,066	9.99E-07	4.891	0.099	1	1	rs4676609	39,214,256	4.63E-06
3	ENST00000274695	CDKAL1	6	20,534,688	21,232,635	1.29E-06	-4.841	-0.046	1	1	rs9295474	20,652,717	7.61E-08
4	ENST00000481601	CCDC183	9	139,694,767	139,702,192	9.60E-07	-4.490	-0.048	2	2	rs2811736	139,651,954	3.12E-05
	ENST00000464157	ABCA2	9	139,902,688	139,903,240	7.39E-07	-4.951	-0.235	1	1	rs2811736	139,651,954	3.12E-05
5 *	ENST00000543000	PLEKHG6	12	6,426,733	6,427,529	3.30E-09	6.003	0.076	3	2	rs10849433	6,406,904	6.73E-17
6	ENST00000448790	TOX4	14	21,945,335	21,967,315	1.22E-07	5.290	0.498	1	1	rs3811252	22,855,779	2.11E-05
7	ENST00000478981	BNIP2	15	59,955,092	59,961,148	9.91E-07	-4.893	-0.326	1	1	rs7182962	59,945,783	6.04E-08

8	ENST00000310144	<i>PSMC5</i>	17	61,904,543	61,909,379	4.18E-10	6.247	0.553	1	1	rs12449782	61,576,249	2.18E-05
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599 **Table 4. Colorectal cancer risk associations identified by cross-tissue transcriptome-wide association study.** SMultiXcan uses a two-
600 sided F-test to quantify the significance of the joint fit of the linear regression of the phenotype on predicted expression from multiple
601 tissue models jointly. TWAS tests were performed separately for the following tissue categories: “*Colon_sigmoid*”: GTEx (n=318
602 samples; $P_{\text{Bonferroni}} = 8.12 \times 10^{-6}$ for the $P_{\text{S-PrediXcan}}$); “*Immune*”: DGN + GTEx Cells_EBV-transformed_lymphocytes + GTEx Whole_Blood
603 + GTEx_Spleen (n=1,966 samples; $P_{\text{Bonferroni}} = 3.34 \times 10^{-6}$ for the $P_{\text{S-MultiXcan}}$); “*Mesenchymal*”: GTEx Adipose_Subcutaneous + GTEx
604 Adipose_Visceral_Omentum + GTEx Cells_Cultured_fibroblasts (n=1,533 samples; $P_{\text{Bonferroni}} = 3.96 \times 10^{-6}$ for the $P_{\text{S-MultiXcan}}$);
605 “*Gastrointestinal*”: the 6 in-house colorectal mucosa datasets + GTEx Pancreas + GTEx Liver + GTEx Stomach + GTEx Terminal_Ileum +
606 GTEx Oesophageal_Mucosa + GTEx Colon_Transverse (n=2,615 samples; $P_{\text{Bonferroni}} = 3.34 \times 10^{-6}$ for the $P_{\text{S-MultiXcan}}$); “*All*”: the 6 in-house
607 colorectal mucosa datasets + all GTEx 49 tissues + DGN (n=16,832 samples; $P_{\text{Bonferroni}} = 2.31 \times 10^{-6}$ for the $P_{\text{S-MultiXcan}}$). Other annotations
608 are as per **Table 2**.

609

#	Gene	Ch r	Start (bp, GRCh37)	End (bp, GRCh37)	$P_{\text{S-MultiXcan}}$	Tissue	Mean z score	Effect size	n models	n indep	Top GWAS SNP at <1Mb	SNP location	P_{GWAS}
1	<i>RPL5</i>	1	93,297,540	93,307,481	2.27E-07	All	-1.160	-0.167	2	2	rs7530780	93,130,268	4.18E-05
2	<i>LINGO4</i>	1	151,772,740	151,778,546	2.73E-08	All	1.666	0.034	27	6	rs9826	151,778,899	3.81E-06
3	<i>FAM98A</i>	2	33,808,725	33,824,429	2.98E-06	Immune	4.672	0.166	1	1	rs1448561	33,854,344	5.92E-07
4	<i>FBLN7</i>	2	112,895,962	112,945,793	1.28E-06	All	-0.711	-0.023	28	10	rs7580507	112,879,209	2.71E-07
5	<i>ARHGEF4</i>	2	131,671,559	131,804,836	2.33E-08	All	-0.243	-0.026	14	8	rs73960398	131,795,345	4.86E-06
6	<i>GBE1</i>	3	81,538,850	81,811,312	1.95E-12	All	-0.557	-0.032	8	7	rs554330436	81,039,172	1.69E-04
7	<i>DIRC2</i>	3	122,513,642	122,599,986	1.25E-06	All	0.812	0.003	16	13	rs6774610	122,521,477	6.85E-07
8	<i>GAB1</i>	4	144,258,304	144,395,721	1.11E-07	All	1.756	0.040	10	6	rs72726477	143,517,452	2.91E-05

9	<i>FBXO38</i>	5	147,763,498	147,822,399	2.11E-06	Mesenchymal	4.677	0.287	2	2	rs35548425	147,816,153	1,80E-07
10	<i>EPB41L2</i>	6	131,160,487	131,384,462	2.70E-11	Gastrointestinal	-1.720	-0.018	8	6	rs12662663	131,398,523	6.71E-08
	<i>EPB41L2</i>	6	131,160,487	131,384,462	2.96E-09	All	-0.108	0.024	24	11	rs12662663	131,398,523	6.71E-08
11	<i>CDK6</i>	7	92,234,235	92,465,908	8.00E-14	All	0.281	0.037	8	6	rs143120528	92,258,733	2.49E-07
12	<i>PSMD13</i>	11	236,546	252,984	3.89E-06	Mesenchymal	1.737	0.113	3	2	rs7394572	432,436	4.88E-06
	<i>IFITM1</i>	11	313,506	314,456	6.73E-07	All	-0.090	-0.071	33	18	rs7394572	432,436	4.88E-06
13	<i>RHOG</i>	11	3,848,208	3,862,213	1.58E-06	Gastrointestinal	-1.862	-0.232	2	2	rs10835185	3,862,343	5.97E-08
	<i>RHOG</i>	11	3,848,208	3,862,213	8.27E-07	Mesenchymal	-4.929	-0.476	1	1	rs10835185	3,862,343	5.97E-08
	<i>OR51E2</i>	11	4,701,401	4,719,084	7.44E-06	Colon Sigmoid	4.480	0.336	1	1	rs10835185	3,862,343	5.97E-08
14	<i>ME3</i>	11	86,152,150	86,383,678	2.62E-06	Gastrointestinal	-0.215	-0.125	5	5	rs74402426	86,161,656	1.89E-05
15	<i>TAGLN</i>	11	117,070,037	117,075,052	5.80E-09	All	-2.118	-0.111	14	9	rs1035237	116,727,850	5.43E-08
15	<i>PCSK7</i>	11	117,075,499	117,103,241	2.67E-06	Mesenchymal	3.281	0.311	2	2	rs1035237	116,727,850	5.43E-08
16	<i>CLIP1</i>	12	122,755,979	122,907,179	7.61E-08	All	0.664	0.026	6	5	rs1716169	123,716,930	1.58E-06
17	<i>ATP2C2</i>	16	84,402,133	84,497,793	4.44E-07	Gastrointestinal	1.903	0.021	7	5	rs7187803	84,501,660	1.07E-05
	<i>ATP2C2</i>	16	84,402,133	84,497,793	2.89E-07	All	0.754	0.010	23	14	rs7187803	84,501,660	1.07E-05
18	<i>CBFA2T3</i>	16	88,941,266	89,043,612	1.11E-06	Mesenchymal	4.871	0.253	1	1	rs502258	88,968,547	9.90E-06
19	<i>LLGL1</i>	17	18,128,901	18,148,149	3.05E-06	Immune	-4.667	-0.469	1	1	rs6502570	17,183,255	2.63E-06
20	<i>PSMC3IP</i>	17	40,725,329	40,729,849	2.21E-06	All	1.575	0.108	11	9	rs12949918	40,526,273	1.39E-06
	<i>BECN1</i>	17	40,963,673	40,985,158	1.14E-06	Immune	4.824	0.547	2	2	rs12949918	40,526,273	1.39E-06
21	<i>SMAD4</i>	18	48,554,764	48,611,415	2.75E-06	Mesenchymal	4.750	0.653	2	2	rs12958467	48,481,751	4.69E-07
22	<i>ATP8B1</i>	18	55,313,658	55,470,547	2.54E-06	Immune	-4.704	-0.203	1	1	rs8097764	55,317,896	1.49E-07

23	<i>LIF</i>	22	30,636,528	30,640,922	4.96E-06	Colon Sigmoid	-4.566	-0.201	1	1	rs12484740	30,606,927	4.97E-06
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612 **Table 5. Colorectal cancer risk associations identified by methylome-wide association study.** SMultiXcan uses a two-sided F-test to
 613 quantify the significance of the joint fit of the linear regression of the phenotype on predicted expression from multiple tissue models
 614 jointly. All associations shown were methylome-wide significant after Bonferroni correction for 88,888 CpGs with an S-PrediXcan
 615 model ($P = 0.05/88,888 = 5.62 \times 10^{-7}$ for the $P_{S\text{-MultiXcan}}$). Pairs of CpGs or strings of adjacent CpGs within 1Mb of one another were
 616 considered to lie within the same cluster. Five CRC associations were found for which all CpGs were > 1 Mb away from GWAS-significant
 617 SNP ($P_{GWAS} < 5 \times 10^{-8}$), although near a SNP close to genome-wide significance. Two further associations for 4 CpGs (*) were identified
 618 based on conditional MWAS analysis (**Supplementary Table 15**). Novel CpG hits were all independent of each other and of GWAS SNPs
 619 and TWAS genes. Other annotations are as per **Table 2**.

620

#	CpG	Annotated Gene	Chr	Probe location (bp, GRCh37)	Probe annotation	$P_{S\text{-MultiXcan}}$	Mean z score	Effect size	n models	n indep	Top GWAS SNP at <1Mb	SNP location	P_{GWAS}
1	cg01716680	<i>GJA4</i>	1	35,259,750	S Shore	3.41E-07	-5.099	-0.164	1	1	rs57975061	34,890,238	2.42E-06
2	cg15917621	<i>NRBP1</i>	2	27,650,478	N Shore	1.61E-07	-3.301	-0.094	2	2	rs4665972	27,598,097	1.58E-07
3	cg02609692	<i>LMX1B</i>	9	129,389,125	Island	4.24E-07	5.058	0.112	1	1	rs4075850	130,169,301	1.76E-06
4*	cg12931523	<i>TTL13</i>	15	90,793,004	S Shore	7.74E-09	4.511	0.067	3	3	rs71407320	91,185,291	3.61E-08
	cg05239308	<i>TTL13</i>	15	90,793,057	S Shore	1.54E-07	5.364	0.114	3	2	rs71407320	91,185,291	3.61E-08
	cg27018984	<i>TTL13</i>	15	90,796,558	S Shelf	3.64E-09	-5.900	-0.089	1	1	rs71407320	91,185,291	3.61E-08
5	cg02086790	<i>AXIN1</i>	16	375,327	Island	2.75E-07	2.471	0.042	3	3	rs9921222	375,782	7.10E-07
6*	cg09894072	<i>PLA2G15</i>	16	68,279,487	Island	2.26E-07	5.176	0.096	1	1	rs9939049	68,812,301	1.95E-12

7	cg15135657	LOC100631378	19	38,346,511	S Shore	1.55E-07	-2.170	-0.032	2	2	rs55876653	39,146,780	2.10E-06
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623 **Figure 1. Summary of the study data and analytical design, and the number of previously unreported CRC risk loci discovered.** The
624 figure illustrates the information for the different analyses used: GWAS (green), TWAS (blue), MWAS (yellow) used to identify
625 additional risk loci. These are later used to select credible effector genes annotated to functions and tissues.

626

627 **Figure 2. Effector genes for CRC risk and the cellular processes in which they act.** Pie chart describing the proportion and list of
628 effector genes allocated to each process.

629

630 **Figure 3. Representation of effector genes and their putative actions in the colorectum.** Diagram representing the processes that
631 the combined GWAS, TWAS and MWAS analyses have unveiled as relevant to CRC risk. Exemplar effector genes from cellular processes
632 and pathways (in capitals) are chosen to depict each category.

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692

693 **Methods**

694 **The research presented in this study complies with all relevant ethical regulations, and has**
695 **been approved by the South Central Ethics Committee (UK) (reference number 17/SC/0079).**

696

697 **Data availability**

698 Summary level data for the full set of Asian and European GWAS are available through GWAS
699 catalog (accession number GCST90129505). For individual-level data, CCFR, CORECT, CORSA_2
700 and GECCO are deposited in dbGaP ([phs001415.v1.p1](#), [phs001315.v1.p1](#), [phs001078.v1.p1](#),
701 [phs001903.v1.p1](#), [phs001856.v1.p1](#) and [phs001045.v1.p1](#)). NSCCG and COIN are available in the
702 European Genome-phenome Archive under accession numbers [EGAS00001005412](#) (NSCCG),
703 [EGAS00001005421](#) (COIN). UK Biobank data are available through <http://www.ukbiobank.ac.uk/>
704 and Finnish data through [THL Biobank](#). Access to individual-level data for the remaining studies
705 is controlled through oversight committees. CCFR 1 and CCFR 2 data can be requested by
706 submitting an application for collaboration to the CCFR (forms, instructions and contact
707 information can be located at (www.coloncfr/collaboration.org). Applications for individual level
708 data from the QUASAR2 and SCOT clinical trials will be assessed by the Translational Research
709 Steering Committees that oversee those studies. Individual level data from the CORGI (UK1) study
710 will be made available subject to standard institutional agreements. Application forms for these
711 three studies, and for Scotland Phase 1, Scotland Phase 2, SOCCS, DACHS4 and Croatia, will be
712 provided by emailing a request to access.crc.gwas.data@outlook.com. For access to CORSA_1,
713 please contact gecco@fredhutch.org. For Generation Scotland (GS) access is through the GS
714 Access Committee (GSAC) (access@generationscotland.org). Applications for The Lothian Birth
715 Cohort data should be made through [https://www.ed.ac.uk/lothian-birth-cohorts/data-access-](https://www.ed.ac.uk/lothian-birth-cohorts/data-access-collaboration)
716 [collaboration](#). For details of the application process for Aichi1, Aichi2, BBJ, Guanzhou1, HCES,
717 HCES2, Korea and Shanghai cohorts, please go to <https://swhs-smhs.app.vumc.org/> or contact
718 Dr. Zheng at wei.zheng@vanderbilt.edu.
719 CRC-relevant epigenome data were obtained from the NCBI Gene Expression Omnibus (GEO)
720 database under accession number [GSE77737](#) and [GSE36401](#).

721 Genetically predicted models of gene expression and methylation have been deposited in the
722 Zenodo repository (<https://zenodo.org/deposit/6472285>).

723

724

725 **Code availability**

726 All bioinformatics and statistical analysis tools used in this study are open source, details of which
727 are available in the Methods section and in the Reporting Summary. No custom code was used
728 to process or analyse data. Details on URLs used can be found in the Supplementary Note.

729

730

731 **Statistics and reproducibility**

732 No statistical method was used to predetermine sample size. The experiments were not
733 randomized. Data exclusion from each analysis is explained below in the corresponding sections.
734 Informed consent was obtained for all participants in the study. A description of the different
735 datasets and cohorts used is included in the Supplementary Note.

736

737

738 ***Criteria for declaring new CRC risk associations***

739 Multi-omic studies present inherent difficulties for deciding on what constitutes a novel GWAS,
740 TWAS or MWAS association. To declare statistically significant associations, for GWAS we have
741 used the established threshold of $P = 5 \times 10^{-8}$. We applied this to both loci >1Mbp from a
742 previously known SNP and analyses conditioned on the most significant SNP within 1Mb region.
743 For TWAS or MWAS we also followed convention and used a Bonferroni correction $P = 0.05/N$,
744 where N is the number of gene models successfully derived from the reference tissue.
745 Furthermore, for TisWAS and cross-tissue TWAS, we used Bonferroni-corrected P -value
746 thresholds for significance in each of the reference tissue data sets separately, owing to the
747 overlap in between tissue groups and the fact that many eQTLs are present across tissues. A
748 further common practice, is that a new association should be located >1Mb from another
749 association (from this study or previously reported), whether a genome-wide significant GWAS
750 SNP, a TWAS gene or an MWAS CpG. However, use of the 1Mb distance convention introduces a

751 further problem in that, whilst the location of a GWAS SNP and MWAS CpG can be defined
752 precisely, the location of a gene cannot. We therefore defined a gene's boundaries by the
753 canonical transcript and novel associations must lie 1Mb from both those boundaries. Since
754 TWAS and MWAS associations can affect multiple nearby genes or CpGs (*e.g.* owing to co-
755 regulation or LD between eQTLs or mQTLs), we have conservatively assigned each TWAS and
756 MWAS association to a single locus (defined as a group of genes or CpGs that are significantly
757 associated with CRC risk and lie < 1Mb apart). Locus boundaries must be > 1Mb from another
758 association to be declared an independent risk association.

759 We have also performed conditional analyses across GWAS, TWAS and MWAS. This is standard
760 practice in GWAS (see below)²⁴, whereby nearby SNPs with no or limited correlation can be
761 independently associated with CRC risk. Conditioning TWAS, TIsWAS and MWAS on GWAS using
762 sMIST also allowed us to identify risk associations that were independent of the GWAS
763 associations within 1Mb, based on a $P_{conditional}$ that (i) remained Bonferroni-significant at the
764 unconditional analysis threshold, and (ii) was within one order of magnitude as $P_{unconditional}$. A
765 much larger number of TWAS and MWAS associations fulfilled only criterion (i) after conditioning
766 on a GWAS association within 1Mb (Supplementary Table 6, 8 and 15). Whilst we could not
767 exclude the possibility that some of these associations resulted from additional SNPs
768 independent of a nearby GWAS SNP for example, we conservatively did not declare these as
769 novel risk associations.

770

771 ***GWAS data analysis***

772 Meta-analysis: Within each of the 31 analytical units, we conducted logistic regression under a
773 log-additive model to examine the association between allelic dosage for each genetic variant
774 and the risk of CRC, adjusted for unit-specific covariates. Meta-analysis under a fixed-effects
775 inverse-variance weighted model was performed using META v1.7²⁵. Variants in the meta-
776 analysis only included those with an imputation quality score ($info/R^2$) > 0.4, MAF > 0.005, and
777 seen in at least 15 analytical units. The I^2 statistic was calculated to quantify between study
778 heterogeneity and variants with I^2 > 65% were excluded. A total of 8,782,440 variants were taken
779 forward in the meta-analysis. Meta-analysis of risk estimates was conducted under an inverse

780 variance weighted, fixed-effects model³. None of the analytical units showed strong evidence of
781 genomic inflation (λ ranged from 0.95 to 1.28), and the λ value for the meta-analysis was 1.30
782 ($\lambda_{1000} = 1.01$) **Supplementary figure 3**). To account for any -ancestral differences between
783 analytical units, we implemented MR-MEGA v0.1.5²⁶ , including 10 principal components (PCs)
784 in the analysis. To measure the probability of associations being false positives, the Bayesian
785 False-Discovery Probability (BFDP)³ was calculated based on a plausible odds ratio (OR) of 1.2
786 (based on the 95th percentile of the meta-analysis OR values) and a prior probability of
787 association of 10^{-5} .

788

789 Definition of known and novel GWAS SNP risk associations: We identified all previously reported
790 CRC associations at $P < 5 \times 10^{-8}$ by referencing the NHGRI-EBI Catalog of human GWAS and by
791 searching PubMed (performed June 2021)³. Additional articles were ascertained through
792 references cited in primary publications (Supplementary Table 4). Where multiple studies
793 reported associations in the same region ($r^2 > 0.1$ and within 500kb-1Mb of the index SNP), we
794 considered all variants with genome-wide significant associations. Given the improved power and
795 coverage of our study over previous works, we identified the most strongly associated variant at
796 each known signal and used lead variants for further analyses, rather than the previously
797 reported index variants (**Supplementary Table 3**). A genome-wide significant risk variant was
798 considered novel if >1Mb from a known risk variant.

799 GWAS conditional analysis: To identify independent association signals at the discovered CRC risk
800 associations, we performed conditional analyses using GCTA-COJO²⁴ on the meta-analysis
801 summary statistics. Analyses were performed separately for European and East Asian ancestry
802 populations, to account for LD structure differences. The conditioned data were meta-analyzed
803 together as described above, and associations with $P_{\text{conditional}} < 5 \times 10^{-8}$ were considered novel
804 secondary associations. As reference for LD estimation, we made use of genotyping data from
805 6,684 unrelated samples of East Asian ancestry, and 4,284 samples from combined UK10K and
806 European samples in 1000 Genomes.

807

808 **Heritability analysis**

809 We used the LDSC regression package with default parameters as implemented in LD Hub²⁷ to
 810 estimate the SNP heritability from the GWAS meta-analysis summary statistics data³. SNPs were
 811 filtered to HapMap3 SNPs with 1000 Genomes EUR MAF above 5%. SNPs with imputation info
 812 score < 0.9, MAF < 0.01 and within the major histocompatibility complex (MHC) region (i.e. SNPs
 813 between 26Mb and 34Mb on chromosome six were excluded. Precalculated LD scores files
 814 computed using 1000 Genome European data were used.

815 The contribution of risk SNPs to the familial risk of CRC was calculated as $\sum_k \frac{\log \lambda_k}{\log \lambda_0}$, where λ_0 is
 816 the familial risk to first-degree relatives of CRC cases, assumed to be 2.2²⁸, and λ_k is the familial

817 relative risk associated with SNP k , calculated as $\lambda_k = \frac{p_k r_k^2 + q_k}{(p_k r_k + q_k)^2}$, where p_k is the risk allele
 818 frequency for SNP k , $q_k = 1 - p_k$, and r_k is the estimated per-allele OR from the meta-analysis^{3,29}.

819

820

821 ***Pleiotropy analysis***

822 We explored cross-trait pleiotropic effects using the LDSC regression package with default
 823 parameters³⁰ as implemented in LD Hub. The summary statistics for 252 phenotypes were
 824 extracted from LD Hub. For comparability of results across the traits we limited our analysis to
 825 the CRC GWAS of European ancestry. After excluding GWAS performed on non-European
 826 cohorts, traits where the LD Hub output came with the following warning messages: “Caution:
 827 using this data may yield results outside bounds due to relative low Z score of the SNP heritability
 828 of the trait” and “Caution: using this data may yield less robust results due to minor departure of
 829 the LD structure”, as well as highly correlated traits, 171 phenotypes were included in the
 830 analysis. The departure of the LD structure means departure from the assumption of equal LD
 831 structure between two datasets, e.g due to differences in population structure between the
 832 study populations. SNPs from the MHC (chr6 26M~34M) region were removed for all traits prior
 833 to analysis.

834

835 ***Sample size prediction***

836 To estimate the sample size required to detect a given proportion of the GWAS heritability, we
837 made use of GENESIS software (GENetic Effect-Size distribution Inference from Summary-level
838 data)³¹, which implements a likelihood-based approach to model the effect-size distribution in
839 conjunction with LD information, using the three-component model (mixture of two normal
840 distributions). The percentage of GWAS heritability explained for a projected sample size was
841 based on power calculations for the discovery of genome-wide significant SNPs³. The genetic
842 variance explained was calculated as the proportion of total GWAS heritability explained by SNPs
843 reaching genome-wide significance at a given sample size.

844

845 ***TWAS analysis***

846 Gene expression models for the six in-house expression datasets were generated using the
847 PredictDB v7 pipeline for a total of 1,077 participants^{9,10}. Elastic net model building with 10-fold
848 cross-validation was performed independently for each dataset. The elastic net models for GTEx
849 v8 Colon Transverse were obtained from the PredictDB data repository (<http://predictdb.org/>)
850 and had been generated using the same pipeline. Models were computed using HapMap2 SNPs
851 ± 1 Mb from each gene, together with covariate factors estimated using PEER³², clinical covariates
852 when appropriate (age, sex and, where appropriate, case-control status, type of polyp and
853 anatomic location in the colorectum), and three PCs from the individual dataset's SNP genotype
854 data. Transcriptome-wide association tests were then performed for each dataset with the S-
855 PrediXcan feature using summary statistics from the GWAS meta-analysis. We used individual
856 level GWAS data from GECCO (n=8,725) to derive the LD reference covariance matrix. S-
857 MultiXcan analysis was then undertaken across datasets. Significant associations were declared
858 using Bonferroni correction (0.05/number of gene models from S-MultiXcan). As
859 recommended³³, an additional filter of a TWAS association statistic, $P_{S\text{-PrediXcan}} \leq 10^{-4}$, in at least
860 one individual reference data set was implemented to minimize potential errors due to LD
861 mismatches. Genes localizing to the HLA/MHC region (chr6:28,477,797-33,448,354bp) were
862 excluded.

863 Transcript-based TWAS analyses (TIsWAS) were likewise performed by using transcript-level data
864 from the SOCCS, BarcUVa-Seq and GTEx Colon Transverse datasets.

865 Additional TWAS analyses were similarly performed using the non-colonic mucosa tissue data
866 available from GTEx. These correspond to S-PrediXCan elastic net models from 48 additional GTEx
867 tissues with eQTL data and the DGN whole blood cohort. Five tissue groupings were tested:
868 “*Sigmoid colon*”, corresponding to muscle and other sub-epithelial tissues; “*Immune*”,
869 comprising DGN + GTEx Cells_EBV-transformed_lymphocytes + GTEx Whole_Blood +
870 GTEx_Spleen (n=1,966 samples); “*Mesenchymal*”, comprising GTEx Adipose_Subcutaneous +
871 GTEx Adipose_Visceral_Omentum + GTEx Cells_Cultured_fibroblasts (n=1,533 samples);
872 “*Gastrointestinal*”, comprising six in-house datasets + GTEx Pancreas + GTEx Liver + GTEx
873 Stomach + GTEx Terminal_Ileum + GTEx Oesophageal_Mucosa + GTEx Colon_Transverse;
874 n=2,615 samples); and “*All*”, comprising the six in-house datasets + all 49 GTEx tissues + DGN
875 (n=16,832 samples).

876 The predictive performance of the models for TWAS and TisWAS across the datasets was similar.
877 For the TWAS models the number of genes successfully predicted with $R^2 > 0.01$ (equivalent of
878 $R > 0.1$) varied between 3308 for the BarcUVa data set and 5092 for SOCCS rectum, while GTEx
879 Colon Transverse models were available for 6295 genes. The mean CV-based prediction R^2 for all
880 genes varied between 0.09 (25-75th percentile 0.04-0.12) for BarcUVa to 0.19 for INTERMPHEN
881 (0.07-0.24), compared with 0.12 (0.04-0.16) for GTEx Colon Transverse model. The numbers were
882 slightly higher when comparing the overlapping 736 genes only. The in-house TisWAS models
883 were constructed for a lesser number of transcripts (n=4632 for BarcUVa dataset and n=11262
884 for SOCCS rectum dataset) compared to GTEx Colon Transverse (n=15500), owing to greater read
885 depth and larger sample size for GTEx. The mean R^2 for all genes varied from 0.07 (0.03-0.09) for
886 BarcUVa to 0.16 for SOCCS colon (0.07-0.21). GTEx Colon Transverse had mean R^2 0.10 (0.03-
887 0.12).

888

889

890 ***MWAS analysis***

891 Methylation beta values were calculated based on the manufacturer’s standard, ranging from 0
892 to 1. Quality control and data normalization were performed in R using the ChAMP software
893 pipeline for the EPIC and 450K arrays³⁴. Briefly, we filtered out failed probes with detection $P >$

894 0.02 in >5% of samples, probes with <3 reads in >5% of samples per probe and all non-CpG
 895 probes. Samples with failed probes >0.1 were also excluded from downstream analyses. We
 896 discarded all probes with SNPs within 10bp of the interrogated CpG (from 1,000 Genomes
 897 Project, CEU population)³⁵, and probes that ambiguously mapped to multiple locations in the
 898 human genome with up to two mismatches³³. We only considered probes mapping to autosomes
 899 and those overlapping between the EPIC and the 450K arrays. Normalization was achieved using
 900 the Beta Mixture Quantile (BMIQ) method. Per probe methylation models were created using
 901 the PredictDB pipeline on the normalized methylation matrix and the genotypes as per TWAS
 902 eQTL analysis. To optimize power, we restricted our analysis to 263,341-238,443 (for the 450K
 903 array) and 377,678 (for the EPIC array) probes annotated to Islands, Shores and Shelves, and
 904 discarded “Open Sea” regions. Further analysis was performed as per the TWAS. CpGs were
 905 annotated to a known GWAS signal if within 1Mb of a genome-wide significant GWAS risk SNP
 906 and otherwise considered novel. For the MWAS models the number of CpG probes successfully
 907 predicted with $R^2 > 0.01$ (equivalent of $R > 0.1$) varied from 24325 for INTERMPHEN rectum to
 908 30385 for COLONOMICS. The mean CV-based prediction R^2 for all genes varied from 0.14 (25th-
 909 7th percentile 0.07-0.16) for INTERMPHEN proximal dataset to 0.19 for SOCCS (0.07-0.25).

910

911 ***Conditional analysis using sMiST for TWAS and MWAS findings***

912 S-MultiXcan is a powerful method for assessing predicted gene expression across multiple tissues
 913 and samples, but cannot readily undertake conditional analysis to determine independence of a
 914 TWAS or MWAS association from other GWAS, TWAS or MWAS associations. We therefore used
 915 the summary statistics-based Mixed effects Score Test (sMiST)³⁶ method to perform
 916 conditional analysis of TWAS, TIsWAS and MWAS data adjusting for GWAS risk SNPs. sMiST can
 917 assess the total effect, including both predicted molecular features (gene expression or
 918 methylation) and the residual direct effects of SNPs that are not explained by predicted molecular
 919 features, on CRC risk. To be consistent with S-MultiXcan, we only assessed the association of
 920 predicted molecular features. We first confirmed that there was a strong correlation between
 921 the sMiST and S-MultiXcan results, with minimal discordance (**Supplementary figure 4**). In view
 922 of this, we used sMiST to perform conditional TWAS and MWAS analysis for each of the

923 significantly associated genes or CpGs respectively, conditioning on the lead GWAS-significant
 924 SNP (if present) within 1Mb (**Supplementary Tables 6, 8 & 15**). We also conditioned TWAS on
 925 TWAS, TIsWAS on TIsWAS and MWAS on MWAS. We also conducted TWAS conditioned on
 926 MWAS analyses for the genes for which both significant genetically predicted expression and
 927 methylation models were produced by the PredictDB pipeline. Where multiple CpGs were
 928 annotated to the same gene, we selected the association with the lowest MWAS *P*-value. We
 929 determined the number of genes associated (at Bonferroni-corrected $P = 0.05/6,722 = 7.44 \times 10^{-6}$)
 930 with CRC risk in both TWAS and MWAS (n=43), TWAS-only (n=54), MWAS-only (n=91) or neither
 931 (n=6,534).”

932

933 ***Effector gene identification***

934 To identify the most credible target or “effector” genes at each CRC risk locus, a pragmatic
 935 approach was utilized. After excluding the MHC region, pseudogenes and transcripts of uncertain
 936 significance (generally RPNNNN or ACNNN), the following hierarchical inclusion criteria were
 937 used.

938 For significant (Bonferroni-corrected $P_{\text{TWAS}} < 0.05$) TWAS genes at a locus, the gene most strongly
 939 associated with CRC risk in any tissue, as long as its P_{TWAS} was at least an order of magnitude
 940 lower than any other gene at the locus. (N=112)

941 For loci included under (1), additional genes that remained significant (FDR < 0.05) in conditional
 942 TWAS-TWAS analysis including the lead gene. (N=9)

943 At GWAS loci not included under (1), the most significant (FDR < 0.05) TWAS gene, as long as its
 944 P_{TWAS} was at least an order of magnitude lower than any other gene at the locus. (N=17)

945 TIsWAS analysis consistent with the approach used for TWAS as described in (1-3) above. (N=16)

946 Genes harboring missense or truncating variants in LD ($r^2 > 0.9$) with sentinel GWAS SNPs. (N=1)

947 A set of 155 genes was identified, which corresponds to about two thirds of the CRC risk loci from
 948 GWAS, TWAS and MWAS (**Supplementary Table 17**).

949

950

951 ***The area under the receiver operating characteristics curve (AUC)***

952 We calculated the confounder adjusted AUC of PRS in discriminating individuals with and without
953 CRC by using the propensity score weighting to account for potentially different distribution
954 of confounders between cases and controls³⁷. We adjusted for age, sex, and four PCs as
955 confounders. We obtained the 95% confidence intervals (CI) by bootstrapping and a total of 500
956 bootstrap samples were generated. We calculated adjusted AUCs using the R package ROct.

957

958

959 Methods-only references

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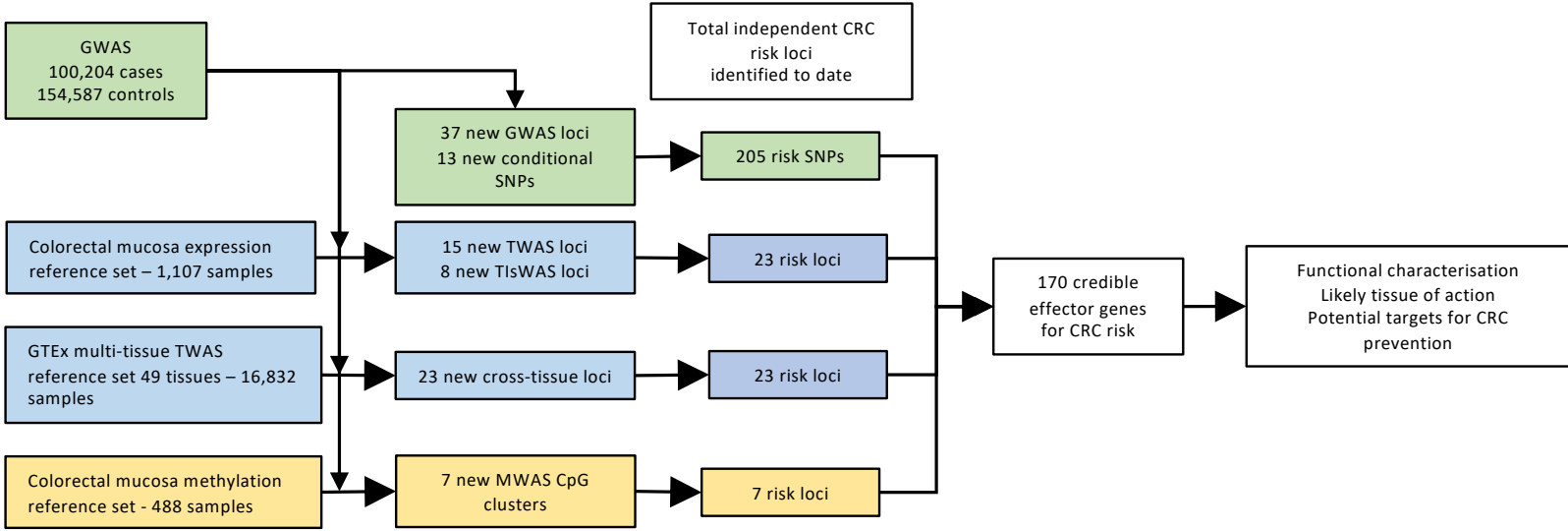
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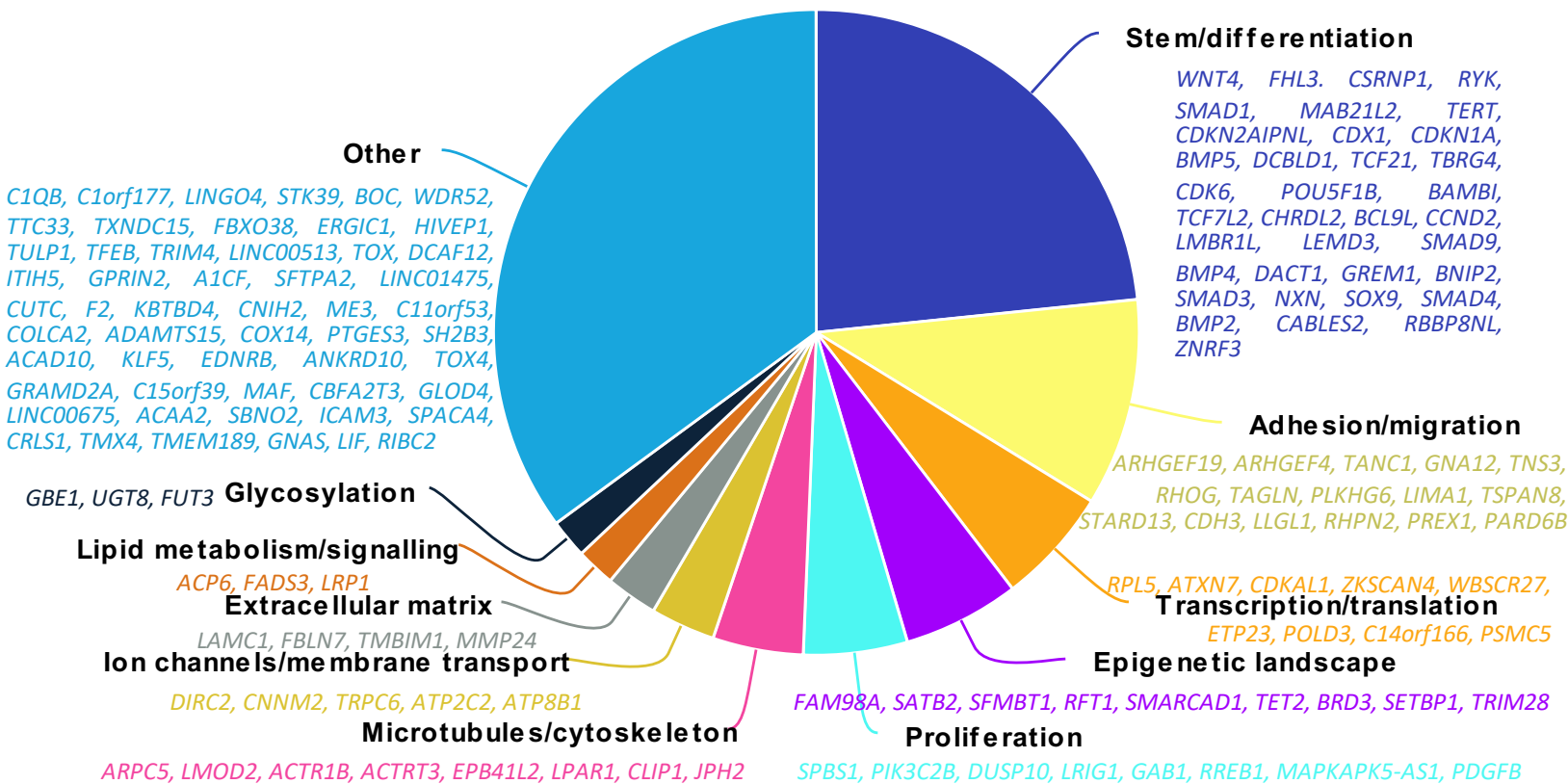
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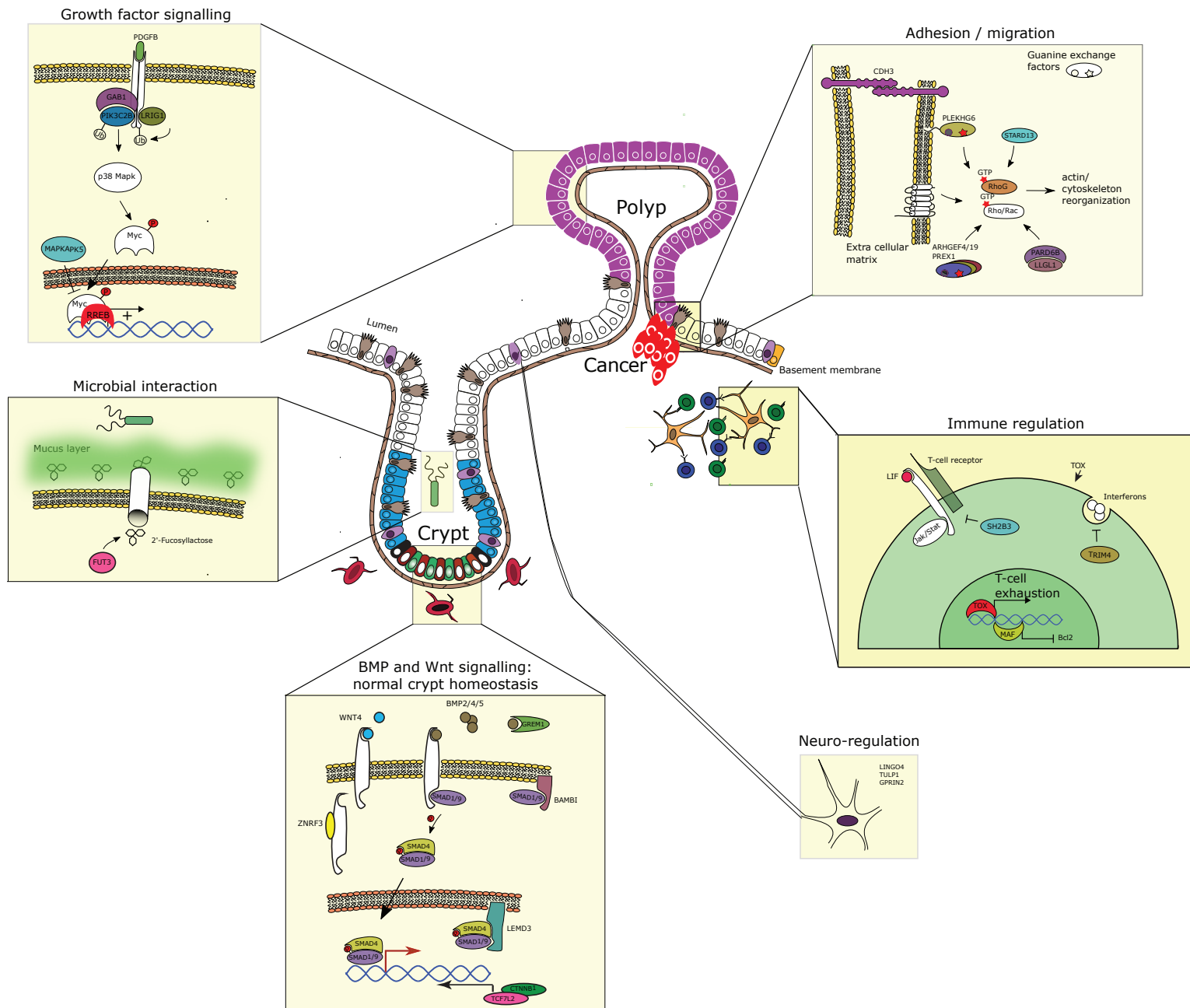
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- Cell key**
- Crypt base columnar
 - Crypt +4 stem
 - Colonocyte
 - Enteroendocrine
 - Paneth-like
 - Transit amplifying
 - Goblet
 - Myofibroblast
 - Dendritic
 - B-cell
 - T-cell
 - Bacteria
 - Adenomatous polyp
 - Cancer