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Citation for final published version:

Scurr, Martin J., Greenshields-Watson, Alexander, Campbell, Emma, Somerville, Michelle, Chen, Yuan, Hulin-Curtis, Sarah L., Burnell, Stephanie E. A., Davies, James A., Davies, Michael M., Hargest, Rachel, Phillips, Simon, Christian, Adam D., Ashelford, Kevin E., Andrews, Robert, Parker, Alan L., Stanton, Richard J., Gallimore, Awen and Godkin, Andrew 2020. Cancer antigen discovery is enabled by RNAsequencing of highly purified malignant and non-malignant cells. Clinical Cancer Research 10.1158/1078-0432.CCR-19-3087

Publishers page: http://doi.org/10.1158/1078-0432.CCR-19-3087

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Cancer antigen discovery is enabled by RNA-sequencing of highly purified malignant and non-malignant cells

Martin J. Scurr^{1^}, Alex Greenshields-Watson^{1^}, Emma Campbell¹, Michelle S.
Somerville¹, Yuan Chen¹, Sarah L. Hulin-Curtis¹, Stephanie E. A. Burnell¹, James A.
Davies², Michael M. Davies³, Rachel Hargest³, Simon Phillips³, Adam Christian⁴,
Kevin E. Ashelford², Robert Andrews¹, Alan L. Parker², Richard J. Stanton¹, Awen
Gallimore^{1*^} and Andrew Godkin^{1,5*^}

8 ¹ Division of Infection & Immunity, Henry Wellcome Building, Cardiff University, Cardiff, UK.

9 ² Division of Cancer & Genetics, Sir Geraint Evans Building, Cardiff University, Cardiff, UK.

10 ³ Dept. of Colorectal Surgery, University Hospital of Wales, Heath Park, Cardiff, UK.

⁴ Dept. of Histopathology, University Hospital of Wales, Heath Park, Cardiff, UK.

- ⁵ Dept. of Gastroenterology & Hepatology, University Hospital of Wales, Heath Park, Cardiff, UK.
- 14 [^] These authors contributed equally to this work.
- 15 * Corresponding Authors:
- 16 Division of Infection and Immunity
- 17 Henry Wellcome Building
- 18 Health Park
- 19 Cardiff
- 20 CF14 4XN
- 21 Tel: +442920687003 / +442920687012
- 22 Emails: <u>godkinaj@cardiff.ac.uk</u>23 gallimoream@cardiff.ac.uk
- 24 Running Title: Identifying novel, immunogenic tumor antigens
- 25

13

Conflict of Interest: M. Scurr, A. Gallimore and A. Godkin are co-inventors of a
 patent regarding DNAJB7 and uses thereof, co-owned by Cardiff University. All other
 authors declare no potential conflicts of interest.

29

30 **Keywords:** Tumor antigens; RNA sequencing; DNAJB7; immunotherapy; cancer 31 vaccine.

- 32
- 33 **Word count** (not including references): 4948
- 34
- 35 **Total number of figures** (not including supplementary): 6
- 36

37 Translational Relevance

In order for cancer vaccination strategies to realise their potential, they must elicit effective anti-tumor immune responses in a broad patient population. Tumor cell purification dramatically improves RNA sequencing resolution to the point where novel, highly differentially expressed, immunogenic proteins become detectable. This novel methodology of tumor antigen identification could enhance future vaccine efficacy. 44 Abstract

45 Purpose: Broadly expressed, highly differentiated tumor-associated antigens (TAA)
46 can elicit anti-tumor immunity. However, vaccines targeting TAAs have
47 demonstrated disappointing clinical results, reflecting poor antigen selection and/or
48 immunosuppressive mechanisms.

Experimental design: Here, a panel of widely expressed, novel colorectal TAAs were identified by performing RNA sequencing of highly purified colorectal tumor cells in comparison to patient-matched colonic epithelial cells; tumor cell purification was essential to reveal these genes. Candidate TAA protein expression was confirmed by immunohistochemistry, and pre-existing T cell immunogenicity towards these antigens tested.

Results: The most promising candidate for further development is DNAJB7 [DnaJ heat shock protein family (Hsp40) member B7], identified here as a novel cancertestis antigen. It is expressed in many tumors and is strongly immunogenic in patients with cancers originating from a variety of sites. DNAJB7-specific T cells were capable of killing colorectal tumor lines in vitro, and the IFN- γ^+ response was markedly magnified by control of immunosuppression with cyclophosphamide in cancer patients.

62 **Conclusion:** This study highlights how prior methods that sequence whole tumor 63 fractions (i.e. inclusive of alive/dead stromal cells) for antigen identification may have 64 limitations. Through tumor cell purification and sequencing, novel candidate TAAs 65 have been identified for future immunotherapeutic targeting.

66 Introduction

Despite understandable excitement surrounding results from cancer immunotherapy 67 studies, actual outcomes are disappointing. We recently demonstrated the principle 68 that immunological responses generated to the 5T4 oncofetal antigen through MVA-69 5T4 (TroVax) vaccination can positively influence the outcome of patients with 70 advanced colorectal cancer (CRC)(1). Whilst survival was significantly prolonged for 71 vaccinated patients mounting an anti-5T4 response, all patients had progressed 72 within 10-months. Indeed, stand-alone anti-cancer vaccines are rarely effective in the 73 74 advanced disease setting; this could be the result of inherent mechanisms of local immunosuppression, or sub-optimal antigenic targets. 75

Many upregulated tumor-associated antigen (TAA) targets are often readily detectable in healthy tissue, e.g. the autoantigen carcinoembryonic antigen (CEA)and directing immune responses against such antigens can lead to side-effects (2) and poor survival outcomes post-surgery (3). Thus, identification of TAAs that can be targeted by immunotherapy is a balance between expression on tumor and healthy tissue. The challenge is further complicated by T cell cross-reactivity which can result in off-target effects in distant tissue with potentially fatal consequences (4).

Whilst immunotherapies targeting neoepitopes hold promise, they are highly 83 focused to the individual and currently prohibitively expensive to develop. For 84 therapies relevant to the wider population such as cancer vaccines, antigens must 85 be broadly expressed in the same tumor types of multiple individuals and present at 86 minimal levels in healthy tissue. Ideal discovery pipelines would involve large scale 87 analysis of TAA candidates followed by selection based on immunogenicity and 88 tissue-specific expression. Candidates that fit these criteria could be explored further 89 90 with cancer vaccination and CAR-T cell therapy. Indeed, vaccinations targeting non-

mutated tumor antigens are capable of inducing robust T cell responses in cancer
patients (1,5).

The development of RNA-sequencing in differential expression analysis 93 provides an attractive methodology to initiate TAA discovery pipelines. However, this 94 technology is limited by the heterogeneity of the tissue in question and is not 95 extensively used in TAA discovery. For the colon, a mixture of immune cells, 96 epithelium and stroma complicates expression profiles, limiting identification of 97 significantly differential expressed genes especially when tumor immune infiltrate 98 99 varies highly between individuals and tumor location. Purification of epithelial and tumor cells prior to RNA-sequencing analysis is a novel methodology developed to 100 overcome tissue heterogeneity. In this study, we used EpCAM purification of tumor 101 and healthy colonic epithelium at two sites to improve the resolution between 102 expression profiles and thus aid identification of differentially expressed genes 103 (DEG). Gene lists were created based on expression profiles between all tissues, 104 and significance levels in a DESeg2 comparison analysis. These lists were analysed, 105 and several genes selected for further investigation. Immunogenic analysis and 106 tissue expression of the protein products of these genes in healthy tissues were 107 used to select the best candidate for cancer immunotherapy. 108

109

110 Materials and Methods

111 Excision of colonic and tumor tissue

Tumor and paired background (unaffected) colon specimens were obtained from 112 three patients undergoing anterior resection for primary rectal cancer at the 113 University Hospital of Wales, Cardiff (see Supplementary Table 1 for patient 114 characteristics). Autologous colon samples were cut from macroscopically normal 115 sections of the excised tissue, both "near" (within 2 cm) and "far" (at least 10 cm) 116 from the tumor site. All fresh tumor samples were derived from the luminal aspect of 117 118 the specimen, so as not to interfere with histopathological staging. All patients and participants gave written, informed consent personally prior to inclusion. This study 119 was conducted in accordance with the Declaration of Helsinki. The Wales Research 120 Ethics Committee granted ethical approval for this study. 121

122 Patient treatment schedule

Orally administered 50mg cyclophosphamide was taken twice-a-day on treatment days 1–7 and 15–21; no cyclophosphamide was taken on treatment days 8–14 or 22–106, or until patient relapsed. Peripheral blood samples (40ml) were taken at regular intervals during therapy.

127 Purification of tissue samples

Background colon and tumor specimens were transported and washed in extraction medium supplemented with 2% human AB serum (Welsh Blood Service), gentamicin and Fungizone (ThermoFisher). Within 30-minutes of resection from a patient, samples were minced and forced through 70µm cell strainers to collect a single cell suspension. In no instances were collagenase or DNase treatments used. Dissociated cell preparations of tumor, near and far healthy colonic tissue were stained with Live/Dead fixable Aqua (ThermoFisher) followed by surface marker

staining with CD3-APC (BioLegend) and EpCAM-PE (Miltenyi Biotec) antibodies. 135 Samples were resuspended in FACS buffer (PBS, 2% BSA) prior to sorting into 136 Live/Dead EpCAM⁺CD3⁻ populations on a FACS Aria III (BD). Tumor tissue also 137 stained with CD3 and EpCAM antibodies was additionally passed through the cell 138 139 sorter without gating, and used as an unsorted control for RNA-sequencing. All samples were sorted directly into RLT buffer (Qiagen) with β -mercaptoethanol 140 (Sigma Aldrich) and frozen at -80°C. Frozen samples were thawed and RNA isolated 141 using an RNeasy Micro kit (Qiagen). 142

143 RNA sequencing

Library preparation and RNA sequencing was carried out by VGTI-FL (Florida, USA). 144 Purified RNA was used to make libraries using an Illumina TruSeg kit. Libraries were 145 sequenced to a depth of 37-63 M read pairs on an Illumina HiSeq platform. Paired 146 end reads were processed on a Cardiff University pipeline. Reads were trimmed with 147 FastQC Trimmomatic (28) and assessed for quality using 148 (https://www.bioinformatics.babraham.ac.uk/projects/fastqc/) using the default 149 parameters. Reads were mapped to Ensembl human genome build GRCh38.89 150 Ensembl FTP downloaded from the site 151 (http://www.ensembl.org/info/data/ftp/index.html/) using STAR (29). Counts were 152 assigned to transcripts using featureCounts with the GRCh38.89 gene build GTF 153 (30). RNA-seq data have been deposited in the ArrayExpress database at EMBL-154 EBI (www.ebi.ac.uk/arrayexpress) under accession number E-MTAB-8803. 155

156 Differential expression analysis

Aligned reads were normalised using DESeq2 in R (31). Differentially expressed genes were identified between purified tumor samples, purified near, and purified far epithelium. Differential expression analysis was carried out using DESeq2 between

sample types for all donors in a paired analysis. Comparisons of tumor and near or
far normal epithelial tissue were carried out. For the three-donor expression analysis,
genes with a Log 2-fold change greater than 3.5, FPKM values in healthy tissue less
than 3.5, and FPKM values in tumor greater than 4.0 in any two of three donors and
p-adjusted < 0.05 (Benjamini and Hochberg (8), in three donors) were taken forward
for further analysis.

The analysis was expanded to genes which were significantly differentially expressed in separate comparisons for data in two of the three donors. Higher expression cut off values were used with FPKM greater than 5.0 in both donor's tumor tissue, and less than 1.0 in healthy tissues, with a log 2-fold change greater than 6 and p-adjusted < 0.05.

171 Analysis of TCGA RNA-seq data

Level 3 (raw counts, htseq.counts.gz) RNA-seq data, and sample meta data (GDC sample sheet and clinical cart files) were downloaded from the TCGA GDC portal (<u>https://portal.gdc.cancer.gov/</u>) on 15-May-2018 for colon and rectal adenocarcinoma patients (TCGA-COAD, TCGA-READ). All datasets were normalized as one matrix using DESeq2 in R and output normalized counts were used for all analyses, with data visualized using the pheatmap R package.

178 Antigens

20mer peptides overlapping by 10 amino acids, covering the entire protein sequence of each identified TAA, were synthesized by Fmoc chemistry to >95% purity (GL Biochem), and divided into pools, as shown (Supplementary Tables 2-8). Individual 9mer peptides (to measure HLA-A*02-restricted DNAJB7-specific CD8⁺ T cell responses) were synthesised by Fmoc chemistry to >90% purity (Peptide Synthetics). The recall antigens tuberculin purified protein derivative (PPD; Statens

Serum Institut) and hemagglutinin (HA; gift from Dr. John Skehel, National Institute
of Medical Research) and the T cell mitogen phytohemagglutinin (PHA; Sigma) were
used as positive controls. All antigens were used at a final concentration of 5µg/ml.

188 Peripheral blood mononuclear cell (PBMC) culture

189 Peripheral blood samples were obtained from pre-operative colorectal adenocarcinoma patients (n=17), hepatocellular carcinoma patients 190 (n=2), cholangiocarcinoma patient (n=1), head and neck carcinoma patients (n=2) and age-191 matched non-tumor-bearing donors (n=10). Blood samples were collected in 10ml 192 heparin tubes (BD) no more than 7-days prior to surgery. PBMCs were isolated by 193 centrifugation of heparinized blood over Lymphoprep (Axis-Shield). Cells were 194 washed and re-suspended in CTL Test Plus media (CTL Europe), L-glutamine and 195 penicillin / streptomycin. PBMC were plated in 96-well plates (Nunc) and cultured in 196 duplicate wells with specific antigens for 14-days, supplemented with fresh media 197 containing 20 IU/ml IL-2 on days 3, 7 and 10. 198

Generation of DNAJB7 9mer epitope-specific CD8⁺ T cell lines

PBMCs from two healthy donors and one CRC patient, who were known HLA-A*02 200 positive, were used to generate CD8⁺ T cell lines. The HLA class I epitope prediction 201 algorithms NetMHC 4.0 (32) and SYFPEITHI (33) were used to identify HLA-A*02-202 restricted DNAJB7 9mers predicted to bind with the highest affinity. PBMCs were 203 stimulated with the top five scoring DNAJB7-derived 9mer epitopes in the presence 204 of 40 IU/ml IL-2, 2 ng/ml IL-7, and 5 ng/ml IL-15, in CTL Test Plus culture media. 205 Cells were incubated in 37°C, 5% CO₂ for 9-12 days before testing their specificity 206 by IFN- γ ELISPOT or intracellular cytokine staining. CD8⁺ T cells from positive lines 207 were sorted by MojoSort CD8 T cell isolation kits (Biolegend, UK), and further 208 expanded using irradiated T2 cells, loaded with relevant peptides, and irradiated 209

autologous PBMCs as feeders. Cells were rested for a minimum of three days in
 media containing no cytokines before use in downstream assays.

212 Adenovirus vectors and cell lines

The replication deficient (Δ E1/ Δ E3) adenovirus vectors Ad5-DNAJB7 (expressing the entire DNAJB7 ORF (open reading frame)) and Ad5-EMPTY (lacking an inserted ORF) were generated in the AdZ vector system, using homologous recombineering methods as previously described (34,35). The DNAJB7 ORF was gene synthesized by GeneArt (ThermoFisher Scientific, UK), prior to being inserted into the AdZ vector. Viruses were produced in T-REx-293 and purified as previously described (36).

The colorectal tumor cell lines SW480 (European Collection of Authenticated 220 Cell Cultures (ECACC) 87092801) and Caco-2 (ECACC 86010202) were used in 221 this study, given their confirmed expression of HLA-A*02 and the coxsackie and 222 adenovirus receptor (data not shown). Cell lines were transduced using 2500 virus 223 particles (vp) per cell (SW480) or 10,000 vp/cell (Caco-2) of Ad5 vector expressing 224 DNAJB7 under the control of a CMV promoter, or control empty Ad5 vector for 3 225 hours at 37°C 5% CO₂ in media containing FBS. After the incubation period, cells 226 were washed with fresh media and cultivated according to culture collection 227 protocols (ECACC), before use in downstream assays. Ad5-GFP titration and 228 subsequent flow cytometric analyses had been performed prior to Ad5-DNAJB7 229 transduction to determine the optimal vp/cell dose for each cell line. Vp/ml of each 230 adenovirus was quantified using a Pierce Micro BCA Protein Assay Kit. 231

232 Western blot

Protein was extracted from SW480 and Caco-2 cells using 4x NUPAGE LDS sample
buffer (ThermoFisher Scientific). Protein (6-9µg) was resolved using 4-12% sodium

dodecyl sulphate polyacrylamide electrophoresis gels at 120V for 90 minutes and 235 transferred to PVDF membrane at 30V for 80 minutes at room temperature. 236 Membranes were blocked using 5% milk in PBS plus 0.05% Tween 20 (PBST) and 237 incubated with DNAJB7 antibody (1:1000, rabbit IgG, polyclonal, Bio-Techne) or β-238 actin antibody (1:2000, rabbit IgG, monoclonal, Sigma-Aldrich) overnight at 4°C. 239 Antibodies were prepared in 0.05% milk in PBST. Secondary antibody (1:3000, Anti-240 rabbit IgG, Bio-Rad) was applied for 1-hour at room temperature. Images of the 241 bands were visualized using SuperSignal Pico PLUS chemiluminescent substrate 242 (ThermoFisher Scientific). 243

244 **Real-time cytotoxicity assay (xCelligence)**

Target SW480 or Caco-2 cells were harvested and plated at 20,000 and 12,000 cells 245 per well, respectively, in a 96-well xCelligence E-plate (ACEA Biosciences). 246 Transduction was performed 48 hours prior to plating where appropriate. Suitable 247 cell densities were determined by previous titration experiments. Cell attachment 248 was monitored using the xCelligence Real-Time Cell Analysis (RTCA) instrument 249 until the plateau phase was reached. DNAJB7-specific CD8⁺ T cell lines were added 250 at an effector to target ratio of 5:1 and impedance measurements performed every 251 10 minutes for up to 72 hours. All experiments were performed in duplicate. Changes 252 in electrical impedance were expressed as a dimensionless cell index value, 253 normalized to impedance values immediately preceding the addition of effector T 254 cells. 255

256 ELISpot / FluoroSpot Assays

IFN- γ ELISpot and IFN- γ /Granzyme B FluoroSpot assays were performed as previously described (13). Briefly, PVDF 96-well filtration plates were coated with 50µl antibody (Mabtech). Cells were washed, plated, and stimulated with 5µg/ml

antigen in duplicate wells. Plates were incubated at 37° C, 5% CO₂ for 24-hours before developing spots. Spot-forming cells (SFC), i.e. cytokine-producing T cells, were enumerated using SmartCount settings on an automated plate reader (ImmunoSpot S6 Ultra; CTL Europe). Positive responses were identified as having at least 20 SFC/10⁵ cultured PBMCs, and at least double that of the negative (no antigen) control. Wells with spot counts >1000 were deemed too numerous to count and capped at this level.

267 Flow cytometry

268 To perform T cell counts, 15µl of human TBNK 6-colour cocktail (BioLegend) was added to 50µl of whole heparinized blood using a reverse pipetting technique. Red 269 blood cells were lyzed and samples run on a NovoCyte 3000 (ACEA Biosciences) to 270 271 obtain absolute cell counts. To calculate the proportion of proliferating CD4⁺ regulatory T cells, fresh PBMCs were stained with Live/Dead-Agua (ThermoFisher 272 Scientific), surface stained with CD3-FITC, CD4-BV605 and CD25-BV421 273 (BioLegend), followed by fixation / permeabilization and intracellular staining with 274 Foxp3-APC (ThermoFisher Scientific) and Ki67-PE (BD Biosciences). 275

276 Immunohistochemistry

The identified TAAs from this study were evaluated for protein expression 277 characteristics on healthy tissue and a range of tumor samples by utilising the 278 Human Protein Atlas resource (12). In addition, DNAJB7 expression was assessed 279 on formalin-fixed paraffin embedded blocks of colorectal tumor and healthy colon 280 tissue (see Supplementary Table 1 for patient characteristics), and testis tissue as a 281 positive control for DNAJB7 expression. Immunohistochemistry was performed on 282 the Leica Bond RX Automated Research Stainer. Dewaxing/hydration of 5um 283 sections was performed according to manufacturer's instructions (Leica). Antigen 284

retrieval was performed using Bond Epitope Retrieval Solution 2. DNAJB7 antibody
(HPA000534, Atlas Antibodies) was used at a dilution of 1:100 and incubated for 105
minutes. Antibody detection was performed using Bond Polymer Refine Detection
Kit, followed by hematoxylin counter staining. Following this, samples were
dehydrated, mounted then scanned using Slide Scanner Axio Scan.Z1 (Zeiss);
representative images were taken using Zen Blue software.

292 **Results**

Purification of samples prior to RNA-sequencing provided enhanced resolution of differentially expressed genes

Rectal tumor and paired background (unaffected) colon specimens were obtained from three patients undergoing resection. Autologous colon samples were cut from macroscopically normal sections of the excised tissue, both "near" (within 2 cm) and "far" (at least 10 cm) from the tumor site (Figure 1A). Dissociated single cells were sorted into Live/Dead⁻EpCAM⁺CD3⁻ populations (Figure 1B and C). EpCAM was chosen as it would enable preferential isolation of epithelial populations over stromal tissue and immune populations (6,7).

302 RNA-sequencing datasets were comparable following several normalization procedures. Differential expression comparisons were run using DESeg2 of healthy 303 tissues ("near" and "far") against purified tumor tissue in all three patients, and then 304 separate analyses for each combination of two patients. An additional comparison of 305 non-purified tumor tissue against healthy tissues was run to investigate the impact of 306 EpCAM sorting. To find relevant genes that could be targeted by immunotherapy, we 307 applied criteria that specified very low levels of expression in healthy tissue 308 combined with high expression in tumor tissue (based on FPKM and log 2-fold 309 change). Only genes assigned a p-adjusted < 0.05 (Benjamini and Hochberg (8)) 310 were taken forward for further analysis. 311

Initial gene lists gave 83 significant genes showing differential expression between tumor and far colon tissue, while 92 genes between tumor and near colon tissue. Cross referencing of these gene lists resulted in five genes that satisfied significant criteria in both comparisons (including four of those taken forward; ARSJ, CENPQ, ZC3H12B and CEACAM3). To expand our analysis, we looked at DEGs which were significantly expressed in tumor tissue of two of three patients to a higher

level (increased expression cut-offs and lower threshold of healthy tissue 318 expression). These gene lists were combined with three donor lists, and then near 319 and far tissue cross referenced (Figure 2A). This gave an initial set of 54 genes 320 which were cut to 23 based on levels of expression in healthy tissue of all three 321 donors (Supplementary Figure 1 and Supplementary Table 9). Of these 23 genes, 322 18 were protein coding. We inspected these 18 genes and selected those which 323 were most suitable for further analysis, eliminating those involved in the central 324 nervous system, or which exhibited an inconsistent expression or read mapping 325 profile in three donors gauged by visual curation of mapped reads in IGV (Integrative 326 Genomics Viewer, Broad Institute) (9). 327

The final genes selected were DNAJB7, CENPQ, ZC3H12B, ZSWIM1, CEACAM3, ARSJ and CYP2B6, based on their ideal expression profile for therapeutic exploitation (Figure 2B). Inspection of expression profiles in non-purified tumor tissue (Figure 2B and 2D) exemplified the difficulty in detecting these genes in the absence of purification, with all expression levels lower than purified tissue.

To further assess the impact of tissue purification, we looked at the DEGs between bulk and EpCAM sorted tumor samples, focusing on genes which were expressed in at least two of the three samples (Figure 2C). This emphasised the advantage of purification resulting in enhancement of the most relevant gene expression patterns and demonstrates how our novel antigens could not have been identified from bulk tumor sequencing alone (Figure 2D).

339 Comparison of Common Cancer Antigen Expression

We next wanted to assess the expression patterns of the seven novel antigens identified in the context of other antigens commonly classified in the literature as TAAs (10,11). We compiled a representative list which included antigens such as GP100, 5T4, LAGE3 and MART and looked at how their expression levels

compared within the patient samples used in this study (Figure 3A) and across both colon and rectal tumor data from the TCGA (Figure 3B). This analysis showed a clear distinction in antigen expression level, with CYP2B6, ZSWIM1 and 5T4 expression comparable to the highly expressed LAGE3. However, beyond these four genes the expression levels of other antigens were heterogeneous across tumor samples, and some appeared to be relatively low in the TCGA data, in particular CEACAM3 and DNAJB7.

A critical criterion for the analysis in Figure 2 was differential expression of 351 genes between healthy tissue and tumor tissue. The TCGA data also has several 352 paired datasets of colon and rectal tumors and corresponding heathy tissue. We 353 visually inspected the differences of the TAA panel list across these datasets in order 354 to confirm that our antigen still accorded with this criterion in a large publicly 355 available dataset (Figure 3C). Indeed, when healthy and tumor tissue expression of 356 each were compared, the differentially expressed nature of several antigens was 357 emphasised. For DNAJB7, ZSWIM1 and CENPQ, the contrast between healthy and 358 tumor datasets was skewed towards tumor expression and suggested these would 359 be better targets than ARSJ or CYP2B6 which did not show visual distinction 360 between the two tissue types. Furthermore, other cancer antigens could be classified 361 as having highly favorable (WT1, LAGE3, MART1, AFP) or hypothetically dangerous 362 (ACRBP, SPA17, KLK3) expression patterns between healthy and tumor tissues, 363 and as such their assessment as bona fide TAAs should be reconsidered. Such 364 trends were also present in our data (Figure 3D). 365

Analysis of protein expression across multiple healthy tissues highlights DNAJB7 as a cancer-testis antigen and a suitable target for immunotherapy

368 The protein expression level of each candidate TAA was evaluated using 369 publicly available immunohistochemistry data (12). Whilst each candidate exhibited

significant upregulation on tumor tissue over healthy tissue (with the possible
exception of ARSJ), DNAJB7, a protein belonging to the evolutionarily conserved
DNAJ heat shock family, was unexpectedly identified as a novel cancer-testis
antigen given its complete lack of expression on any healthy tissue bar the testis, an
immune-privileged site (Supplementary Figure 2).

We sought to corroborate this pattern of staining on paraffin-fixed samples acquired in-house, using the same anti-DNAJB7 antibody (HPA000534). In preliminary experiments, expression was higher in certain tumor samples, although antibody staining was observed in background colon tissue (Supplementary Figure 3). Given this finding and the failure to detect DNAJB7 mRNA in normal colon, we conclude that where DNAJB7 is detected, it is preferentially expressed in cancer tissue.

Furthermore, DNAJB7 was expressed on a very wide range of solid tumors, in particular on tumors of the gastrointestinal tract and accessory organs of digestion, including colorectal cancer and pancreatic ductal adenocarcinoma (Supplementary Figures 2B and 2C).

386 **DNAJB7 is a superior cancer-testis antigen**

The expression profile of DNAJB7 was compared to six other well-defined cancer-testis antigens, including NY-ESO-1, MAGE-A1 and SSX2. High protein expression of all these antigens was confirmed to be confined to the testis (Supplementary Figure 4A). In comparison to the other cancer-testis antigens, DNAJB7 was expressed on the greatest range of tumor types, with more than 67% of all patients tested exhibiting positive (low, medium or high) protein expression on their tumor, except for lymphoma (Supplementary Figure 4B).

394 Analysis of candidate TAA T_H1 responses reveal DNAJB7 to be immunogenic

Following identification of relevant genes and confirmed protein expression, 395 we assessed their immunogenicity using overlapping peptide pools and culture with 396 PBMC of CRC patients and healthy donors. Analysis of cultured PBMC by IFN-y 397 ELISpot determined three of the seven proteins to demonstrate immunogenicity in 398 399 most donors tested (Figure 4A-B). As the size of peptide libraries was highly variable for each protein, we standardized the immunogenicity relevant to the number of 400 peptides in each pool (Figure 4A). This analysis revealed CYP2B6, DNAJB7 and 401 CEACAM3 to be comparably immunogenic across multiple individuals without 402 stratification by HLA-type, and similarly immunogenic to the oncofetal antigen 5T4, a 403 tumor antigen that has successfully been targeted in CRC previously (1,13). 404 Conversely, CENPQ and ARSJ were poorly immunogenic in most donors tested. 405

406 Furthermore, our peptide pool design allowed us to interrogate immunogenicity based on a matrix format to determine the peptides responsible for 407 the positive T cell responses (example for DNAJB7, Supplementary Figure 5). This 408 type of analysis may be important for isolation of T_H1 stimulating regions of TAA 409 which can be incorporated in vaccines based on immunogenic components of 410 multiple antigens important in CRC, as well as being regions that can be targeted by 411 epitope-based modifications and strategies for enhancement of the immune 412 response (14). An example of one CRC patient revealed positive IFN- γ and 413 granzyme B responses to DNAJB7 peptide pools 3, 6 and 10, indicative of T cell 414 responses to epitopes contained within peptides 3 and 23 (Supplementary Figure 5B 415 and C). Indeed, peptide 23 was the most immunogenic region of the DNAJB7 416 protein, with responses discovered in 39% of CRC patient and healthy control 417 418 donors tested (Supplementary Figure 5D). DNAJB7 was also found to be immunogenic in patients with other tumor types, including hepatocellular carcinoma, 419

cholangiocarcinoma and the non-gastrointestinal head and neck squamous cellcancer (Figure 4C).

422

CD8⁺ T cell recognition of DNAJB7-expressing colorectal tumor cell lines

 T_{H1} responses described above were dominated by IFN- γ -secreting CD4⁺ 423 424 effector T cells, favoured by the longer 20mer peptides used to stimulate the PBMCs. However, analysis of specific IFN- γ and granzyme B production using 425 FluoroSpot assays indicates that granzyme B is abundantly produced in response to 426 the DNAJB7 peptides (9/10 donors tested, Figure 5A and Supplementary Figure 5) 427 suggesting cytotoxic T cell responses are present. In order to ascertain whether 428 these responses were indicative of CD8⁺ T cells capable of killing DNAJB7-429 expressing tumor cells, HLA-A*02-restricted CD8⁺ T cell epitopes derived from 430 DNAJB7 were identified by computer-based epitope prediction algorithms (Figure 431 5B). The top five scoring peptides stimulated cognate CD8⁺ T cells derived from 432 HLA-A*02⁺ healthy donors and a CRC patient. DNAJB7-specific CD8⁺ T cell lines 433 were successfully enriched in all donors, an example of responses to peptides 434 LTFFLVNSV and GMDNYISVT is shown (Figure 5C). 435

The HLA-A*02-expressing SW480 and Caco-2 colorectal tumor cell lines were 436 used as targets in a cytotoxicity assay, however there was minimal expression of 437 DNAJB7 in both lines (Figure 5D and Supplementary Figure 6). Cell lines were 438 successfully transduced with an Ad5-DNAJB7 viral vector and found to stably 439 increase DNAJB7 expression over a 5-day period before reducing on day 6 (Figure 440 5D and Supplementary Figure 6). Upon addition of effector DNAJB7-specific CD8⁺ T 441 cells to target colorectal tumor cells, real-time impedance traces show a highly 442 significant (P<0.0001) reduction in the number, size/shape, and/or attachment 443 quality of DNAJB7-expressing Caco-2 cells (Figure 5E), and a reduction in the 444

growth of DNAJB7-expressing SW480 cells (Figure 5F) over Ad5-EMPTY
transduced or non-transduced, untreated (UT) tumor cell lines. Non-specific
(DNAJB7-negative) T cell lines did not cause additional killing to DNAJB7-expressing
targets, compared to Ad5-EMPTY or UT cell lines (data not shown). Hence,
DNAJB7-expressing tumor cell lines present peptides on the cell surface by MHC
class I and are selectively eliminated by DNAJB7-specific CD8⁺ T cells.

451 Anti-DNAJB7 T_H1 responses are induced during cyclophosphamide treatment

We have previously demonstrated that anti-tumor T_H1 effector responses are 452 controlled by regulatory T cells (Tregs) (15), and that targeting these Tregs either by 453 depletion in vitro, or inhibition/depletion in vivo with low dose cyclophosphamide, 454 increases the anti-tumor (5T4) immune response (1,13). We sought to assess 455 whether T cell responses were induced to the novel tumor antigens in a CRC patient 456 (Figure 6) and an HCC patient (Supplementary Figure 7) receiving short-term 457 metronomic cyclophosphamide. Anti-5T4 T_H1 responses increased by >4-fold in both 458 patients, an effect previously identified as associating with improved survival 459 outcomes (13); intriguingly, anti-DNAJB7 T_H1 responses also mirrored this treatment 460 response profile in both instances, whereas no responses were induced to ARSJ, 461 CENPQ, ZSWIM1 and CYP2B6 (Figure 6B and C, and Supplementary Figure 7A 462 and B). This could suggest that responses to DNAJB7 and CEACAM3 are 463 suppressed in CRC and HCC, given that responses were unmasked by efficient 464 regulatory T cell depletion (Figure 6D and Supplementary Figure 7C). 465

466

467 **Discussion**

The pursuit of new cancer vaccines that can be administered regardless of patient HLA-type or neoantigen load relies on the investigation and discovery of novel TAAs. Paired RNA-sequencing of tumor and healthy tissue facilitates TAA identification but is limited by the diversity of cellular input in each sequencing sample. Here we purified EpCAM⁺ cellular populations from healthy colon and primary colorectal tumors; RNA-sequencing data from purified samples revealed multiple genes that showed significant differential expression across three donors.

475 In a comparison with non-purified tumor tissue, no genes were classified as significant according to the same criteria, demonstrating the power of using purified 476 tissues in antigen discovery. Further analyses of differentially expressed gene lists in 477 tissues near and far from the tumor helped identify 18 genes which showed 478 differential expression patterns suitable for therapeutics. Four protein products of the 479 identified genes exhibited significant immunogenicity in healthy donors and cancer 480 patients; of these DNAJB7 also demonstrated the most favorable expression profile 481 based on immunohistochemistry data of healthy and cancerous tissue. DNAJB7 482 belongs to the evolutionarily conserved DNAJ/Heat Shock Protein (HSP)40 family of 483 proteins and is a molecular chaperone to HSP40. It is likely that its upregulation in 484 tumors is in response to increased expression of many heat shock proteins, aiding 485 tumor cell proliferation in hostile environments (16). Indeed, other HSP40 family 486 members DNAJB6 and DNAJB8 have both been previously shown to be upregulated 487 in cancer, contributing to cancer-initiating cell maintenance (17-19). Therapies 488 targeting heat shock proteins and their molecular chaperones are already showing 489 promise in cancer treatment (20). Immunotherapeutic targeting of these proteins may 490 also yield further anti-cancer benefits, as implicated by this study. 491

Robust T_H1 responses to DNAJB7 and several other novel tumor antigens 492 were found in healthy controls, in keeping with previous findings for tumor-493 associated antigens from our laboratory (21,22) and others (23,24). It is possible 494 indicative of these responses are а normal functioning process of 495 496 immunosurveillance to remove aberrant epithelial cells. The presence or absence of such responses are now beginning to be exploited for cancer diagnostics and 497 prognostication ((24) and NCT02840058). How and why these T cells exist and are 498 maintained at such a frequency in the memory pool remains unknown: possibilities 499 range from transient upregulation of TAAs during periods of inflammation, e.g. of the 500 colon (22), incomplete thymic selection or antigenic cross-reactivity / mimicry to 501 502 microbial proteins (25).

503 There are limitations to our study, including the selection of luminal tumor sites for cell enrichment as opposed to the invasive margin (required for 504 histopathological assessment of the tumor), the use of purification procedures, i.e. 505 fluorescence activated cell sorting, that may influence mRNA expression prior to 506 RNA isolation, and low initial sample size. However, despite the relatively small 507 scale, the approach described here has successfully identified novel, highly antigenic 508 proteins expressed in cancers. These antigens could be incorporated into vaccines 509 for both therapeutic and prophylactic use. One goal of cancer vaccination in the 510 511 context of CRC immunotherapy is to reduce relapse rates following surgical intervention. Curative rates following resection of primary colorectal tumors are ~60-512 70% but could be improved if relapse was prevented by safely boosting immunity to 513 the proteins with differential expression patterns as a form of prophylactic 514 immunotherapy (26). Indeed, loss of anti-tumor immune responses associates with 515 advancing tumor stage (21,22), and these patients can benefit from anti-cancer 516 vaccination strategies. At the moment, although there is some success using a 517

single TAA in CRC (1), better therapeutic strategies are necessary with superior 518 vaccine targets combined with manipulations of immune regulation. These 519 approaches necessitate discovery of more TAAs and greater investigation of the 520 negative impacts of T cell cross-reactivity and off-target immune effects. Questions 521 over the ideal differential expression pattern, specifically the extent to which some 522 expression in healthy tissue can be tolerated relative to tumor expression are highly 523 relevant. In addition, the targeting of multiple tumor antigens is more likely to 524 overcome inherent tumor immune evasion and evolution. However, cancer-testis 525 antigens allay some of these concerns and represent an ideal tumor target for 526 527 immunotherapy (27). Indeed, here we identified that most donors have the capability 528 to mount anti-DNAJB7 T cell responses, and these responses can be significantly boosted in cancer patients receiving cyclophosphamide. Enhancing this anti-tumor 529 immune response could hold significant potential in future therapeutic and 530 prophylactic treatment strategies. 531

533 Acknowledgments

534

- This study was supported by a Cancer Research Wales programme grant (to A.
- 536 Godkin and A. Gallimore), a Wellcome Trust Collaborator Award Grant
- 537 (209213/Z/17/Z) (to A. Godkin, A. Gallimore, R. Stanton), a Cancer Research UK
- 538 programme grant (C16731/A21200) (to A. Gallimore and A. Godkin) and funding
- 539 from the Wales Cancer Research Centre (to S. Burnell). We are grateful to Prof
- Richard Houlston for useful preliminary discussion on experimental design. We thank
 Dr Catherine Naseriyan for assistance with flow cytometry and cell sorting, Anna
- 541 Dr Catherine Naseriyan for assistance with flow cytometry and cell sorting, Anna 542 Fuller for provision of peptides, Dr Mat Clement for assistance with western blots and
- 543 Dr Ruban Rex Peter Durairaj & Owen Moon for assistance with xCelligence.
- 543 DI RUDAH REX FELEI DUTAITAJ & OWEN MOON TOT ASSISTANCE WITH XCEIIIGENCE.

545 Authors' Contributions

- 546
- 547 Conception and design: M. Scurr, A. Gallimore and A. Godkin
- 548 **Development of methodology:** M. Scurr, A. Greenshields-Watson, K. Ashelford, R. 549 Andrews, A. Parker, R. Stanton and A. Godkin
- 550 **Acquisition of data:** M. Scurr, A. Greenshields-Watson, E. Campbell, M. Somerville 551 and Y. Chen.
- 552 Analysis and interpretation of data: M. Scurr, A. Greenshields-Watson, E.
- 553 Campbell, M. Somerville, Y. Chen, K. Ashelford, R. Andrews, A. Gallimore and A. 554 Godkin
- 555 Writing, review, and/or revision of the manuscript: M. Scurr, A. Greenshields-
- 556 Watson, M. Somerville, Y. Chen, S. Burnell and A. Godkin
- 557 Administrative, technical, or material support: M. Somerville, S. Hulin-Curtis, S.
- Burnell, J. Davies, M. Davies, R. Hargest, S. Phillips, A. Christian, K. Ashelford, R.
- 559 Andrews, A. Parker and R. Stanton.

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678 Figure Legends

Figure 1. Isolation of epithelial and tumor cells by EpCAM-sorting prior to RNAseq. (A) Schematic of tumor and healthy tissue resection taken at two distances from the tumor site. Samples were taken from rectal tumor of three patients. (B) Sample processing and purification flow chart. (C) Flow cytometry gating for EpCAM⁺ and CD3⁻ purification, pre and post cell sorting.

Figure 2. Identification of candidates for further investigation based on differential 684 expression analysis. (A) Workflow for obtaining gene lists of differentially expressed 685 genes based on two comparisons (purified tumor versus purified healthy colon "far," and 686 purified tumor versus purified healthy colon "near," left-hand side and right-hand side, 687 respectively). Gene lists were obtained from significantly differentially expressed genes 688 across all three patients and separately in two of three patients. These were aligned and 689 cross referenced between "far" and "near" comparisons to give a smaller gene list which 690 was further reduced based on expression in healthy tissue, and finally suitability for 691 further investigation. (B) Normalized counts for each of the seven genes selected for 692 further analysis. Counts are shown for each of the four conditions as box plots 693 representing all three patients. (C) Heatmap showing 317 genes that were differentially 694 695 expressed between EpCAM purified and bulk tumor samples and were expressed in at least two of three EpCAM purified samples. Normalized counts were scaled by gene to 696 show relative expression between each sample. (D) Heatmap of novel tumor antigen 697 gene expression (in addition to 5T4), showing differences in normalized counts between 698 EpCAM purified and bulk tumor, scaled as part (C). 699

Figure 3. (A) Heatmap showing relative expression levels of known and novel TAAs
 using normalized counts scaled by sample across EpCAM purified tumor. (B)
 Corresponding analysis using TCGA data for rectal (green bar) and colon (purple bar)

tumors scaled by sample. (C) Comparison of expression levels between tumor and solid
tissue normal in available TCGA data, scaled by gene. Sample type and tissue/tumor
location (colon or rectal) are indicated in the top two bars. (D) Corresponding analysis
performed on our EpCAM purified tumor and healthy data.

707 Figure 4. Immunogenicity of candidate TAAs. T cell responses to peptide pools spanning the entire protein sequence of each candidate TAA were assessed by cultured 708 IFN-y ELISpot (see Supplementary Tables 2-8 for peptide sequences). The total number 709 of IFN- γ^{+} spot-forming cells (SFC) per 10⁵ cultured PBMC relative to the number of 710 peptides spanning the protein was assessed and ranked by mean response (grey bars) 711 amongst all donors tested (A) and then subdivided by CRC patients (blue circles TNM 712 Stage 1/2; n=6, black circles TNM Stage 3; n=8) and healthy donors ('HD', white circles; 713 n=10) (B). (C) Patients with other gastrointestinal cancers were tested for their ability to 714 715 mount anti-DNAJB7 T cell responses (CC – cholangiocarcinoma; HCC – hepatocellular carcinoma; HNC - head and neck squamous cell carcinoma). 716

Figure 5. Enriched DNAJB7-specific CD8⁺ T cells target DNAJB7-expressing 717 colorectal tumor cell lines. T cell responses to two peptide pools spanning the entire 718 DNAJB7 protein sequence were assessed by cultured Granzyme B FluoroSpot (see 719 Supplementary Table 2 for peptide sequences). The total number of Granzyme B⁺ spot-720 forming cells (SFC) per 10⁵ cultured PBMC relative to the 30 peptides spanning the 721 DNAJB7 protein was assessed in 9 CRC patients (A). (B) HLA class I epitope prediction 722 algorithms were used to identify HLA-A*02-restricted DNAJB7 9mers predicted to bind 723 with the highest affinity; top 5 across the algorithms are indicated. (C) DNAJB7-specific 724 725 $CD8^{+}$ T cells were enriched in multiple donors, a representative example of the IFN- γ response in one T cell line to DNAJB7 epitopes LTFFLVNSV and GMDNYISVT is 726 shown. (D) The SW480 CRC cell line was transduced with Ad5-DNAJB7 or an Ad5-727

EMPTY vector, with the expression of DNAJB7 protein indicated by the band at 35kDa 728 and actin control at 45kDa. UT = untreated (non-transduced) SW480 cells. DNAJB7-729 specific T cell lines from a healthy donor ('Donor 1') and a CRC patient ('Donor 2') were 730 seeded into 96-well E-plates, co-incubated with the indicated transduced / non-731 transduced Caco-2 (E) or SW480 (F) cell lines at an effector to target ratio of 5:1. 732 Changes in impedance over a 24-hour period, normalized at the timepoint immediately 733 preceding the addition of effector T cells, are given as a dimensionless normalized cell 734 index. Experiments were performed in duplicates. Statistical results of two-way ANOVA 735 are indicated (*** P<0.0001). 736

Figure 6. Regulatory T cell depletion unmasks T_H1 responses to novel TAAs. (A) A 737 post-colectomy CRC patient received low-dose, metronomic cyclophosphamide on 738 treatment days 1-8 and 15-22, with blood samples collected weekly throughout 739 treatment. T cell responses to peptide pools spanning the entire protein sequence of 740 each candidate TAA were assessed by cultured IFN- γ ELISpot at each timepoint; 741 example images of IFN- γ ELISpot wells are shown (B). (C) The total number of IFN- γ^{+} 742 spot-forming cells (SFC) per 10⁵ cultured PBMC (mean of duplicate wells) were 743 calculated for each TAA. (D) CD3⁺CD4⁺CD25^{hi}Foxp3⁺ regulatory T cell numbers and 744 %Ki67⁺ Tregs were measured by flow cytometry during cyclophosphamide treatment. 745















5T4 MUC16 Project.ID **CEACAM3** CEACAM4 CENPQ SPA17 CTAG2 MAGEA4 GAGE1 ZC3H12B DNAJB7 MART1 ARSJ **MAGEA3** KLK3 MAGEC2 MAGEA1 SSX2 WT1 ZSWIM1 LAGE3 CYP2B6 PIWIL2 GP100 AFP ACRBP PRAME CT45A1

Sample.Type

COAD

READ







