# **Complement Membrane Attack and Tumorigenesis** *A SYSTEMS BIOLOGY APPROACH\**

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Laurence D. Towner<sup>1</sup>, Richard A. Wheat, Timothy R. Hughes, and B. Paul Morgan<sup>2</sup> From the Complement Biology Group, Division of Infection and Immunity, School of Medicine, Cardiff University, Cardiff CF14 4XN, Wales, United Kingdom

Tumor development driven by inflammation is now an established phenomenon, but the role that complement plays remains uncertain. Recent evidence has suggested that various components of the complement (C) cascade may influence tumor development in disparate ways; however, little attention has been paid to that of the membrane attack complex (MAC). This is despite abundant evidence documenting the effects of this complex on cell behavior, including cell activation, protection from/induction of apoptosis, release of inflammatory cytokines, growth factors, and ECM components and regulators, and the triggering of the NLRP3 inflammasome. Here we present a novel approach to this issue by using global gene expression studies in conjunction with a systems biology analysis. Using network analysis of MAC-responsive expression changes, we demonstrate a cluster of co-regulated genes known to have impact in the extracellular space and on the supporting stroma and with well characterized tumor-promoting roles. Network analysis highlighted the central role for EGF receptor activation in mediating the observed responses to MAC exposure. Overall, the study sheds light on the mechanisms by which sublytic MAC causes tumor cell responses and exposes a gene expression signature that implicates MAC as a driver of tumor progression. These findings have implications for understanding of the roles of complement and the MAC in tumor development and progression, which in turn will inform future therapeutic strategies in cancer.

Inflammation is now well established as a crucial contributor in the development and progression of tumors; indeed, it has been included among the second generation hallmarks of cancer (1). A key player in inflammatory responses is the complement (C)<sup>3</sup> system, an innate immune effector with important roles in defense against infection. C provides recognition, early warning signals, and the initial fast response upon exposure to foreign organisms and has evolved to amplify the response to the initial signal (2). C comprises three activation pathways that converge on a common terminal pathway at the stage of C5 cleavage; release of a 74-amino acid peptide C5a, which has potent anaphylatoxic and chemotactic activities (3), leaves the large fragment, C5b, to form the nidus of a membrane-associated complex. Sequential recruitment of C6, C7, C8, and multiple C9 molecules creates a membrane-spanning pore-like cylindrical protein structure known as the membrane attack complex (MAC). The MAC can cause osmotic lysis of certain susceptible bacteria and of metabolically inert cells (4); however, lysis of self-cells is restricted by a combination of regulatory proteins, ion pumps, and MAC removal processes (5). Non-lytic MAC triggers numerous activation events in cells that likely contribute to the proinflammatory activity of C (6).

C has been strongly implicated as an effector in tumor clearance over the past 20 years, largely because of the success of monoclonal antibody (mAb)-based immunotherapies, many of which are designed to harness C as an effector to cause killing of tumor cells (7). In this context, the mAb triggers overwhelming C activation and tumor cell destruction; however, the role of C in tumor clearance in the absence of an activating mAb is much less clear. Indeed, it has been suggested that C activation has a tumor promoting role in many malignancies (8). C activation is known to occur on tumor cells both in vitro and in vivo in many malignancies, including breast (9), papillary thyroid (10, 11), colorectal (12), and ovarian (13) cancers. The best evidence implicating a C activation product as a promoter of tumor development was provided by the demonstration that locally generated C5a recruits myeloid-derived suppressor cells into the tumor where they suppress the anti-tumor activity of CD8<sup>+</sup> T-cells (14, 15). Others have implicated C5a as a factor influencing the balance between tumor promotion and tumor clearance (16), whereas both C3a and C5a have been shown to cause proliferation in tumor cells, for example in neuroblastoma (17, 18). Studies in knock-out mice lacking C3 or C4 confirm important roles for C in tumorigenesis tumor growth being restricted in both C3 and C4 knockouts (14). Despite the large and growing body of evidence supporting a tumor promoting role of C activation (18-21), the role of MAC in tumor biology has been neglected.

Most tumors express and indeed often overexpress membrane bound C regulators CD55, CD59, and CD46 (22). As a consequence, although C is activated in the tumor microenvironment, activation will be restricted, and thus terminal pathway activation and MAC deposition may be insufficient to kill



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<sup>&</sup>lt;sup>1</sup> Present address: Thomson Reuters IP & Science, London EC1N 8JS, UK.

<sup>&</sup>lt;sup>2</sup> To whom correspondence should be addressed: Systems Immunity URI, Division of Infection and Immunity, School of Medicine, Cardiff University, Henry Wellcome Bldg., Heath Park, Cardiff CF14 4XN, Wales, UK. Tel.: 44-2920687096; E-mail: MorganBP@cardiff.ac.uk.

<sup>&</sup>lt;sup>3</sup> The abbreviations used are: C, complement; MAC, membrane attack complex; extracellular matrix; ShE, sheep erythrocytes; ShEA, antibody-sensitized sheep erythrocytes; CFD, complement fixation diluent; pNHS, pooled normal human serum; AN(TF), analyze network (transcription factor); NO, network object; EGFR, EGF receptor; MMP, matrix metalloproteinase; PCA, principal components analysis; ANOVA, analysis of variance; OmCl, O. moubata C inhibitor; qPCR, quantitative PCR.



FIGURE 1. **Optimization of sublytic complement conditions.** *A*, hemolytic activity testing lysis of ShEA by serum with or without 10  $\mu$ g/ml OmCl (C5 blocker), titrated from 16% down to 0%. *B*, susceptibility of CT26 cells to C lysis; cells in a monolayer were loaded with calcein AM then exposed to serum for 1h at 37 °C. Lysis was calculated from the release of calcein into the supernatant and expressed as the percentage of the total entrapped calcein obtained by detergent lysis of the cells. Results are the means of four separate experiments  $\pm$  S.E. *C* and *D*, expression analysis of *OPN* in CT26 cells in response to exposure to sublytic C for 1, 6, and 12 h. CT26 cells were exposed for 1 (*C*) and 6 and 12 (*D*) hours to 5% serum treated with or without a MAC-blocking dose of OmCl and *OPN* gene expression analyzed by qPCR. Expression was calculated as % of untreated control. Results are means of three determinations  $\pm$  S.E. (\*, p < 0.05; \*\*, p < 0.01).

the tumor cell. Nevertheless, MAC deposition on tumor cells at a sublytic level may have a profound impact on the target, for example by causing an immediate increase in intracellular Ca<sup>2+</sup> (23) and downstream activation of signaling cascades (24). The effects of sublytic MAC on cells in vitro include: release of inflammatory mediators such as reactive oxygen or nitrogen species, leukotrienes, arachidonic acid metabolites, and prostaglandins (5); the release of cytokines such as IL-1, TNF  $\alpha$ , IL-8, IL-6, and MCP-1 (25); increased expression of adhesion molecules such as E-selectin, ICAM-1, VCAM-1, and ELAM-1 (26); release of growth factors such as basic FGF, PDGF, EGF, PIGF (placental growth factor), and RANTES (regulated on activation normal T cell expressed and secreted) (27, 28); secretion of extracellular matrix components such as collagen IV and fibronectin (29) and regulators such as MMP2 and MMP9 (30); increased cell proliferation (31); accelerated or inhibited apoptosis (32-34) and activation of the NLRP3 inflammasome (35). Given this catalogue of effects on cell function, it is likely that sublytic MAC will significantly influence tumor cell fate in vivo. Here we take a novel approach to addressing how MAC influences tumor cell fate by adopting an unbiased systems analysis of the effects of sublytic MAC on the patterns of gene expression in a tumor cell, and we identify key pathways implicated and discuss the impact that these might have on tumor survival.

## Results

Sublytic CAttack and MAC inhibition on Tumor Cell Lines— The C5-binding protein Ornithodoros moubata C inhibitor (OmCI) has been extensively characterized and shown to specifically block formation of MAC in human and rodent plasma (36, 37). The dose of OmCI required to completely block MAC formation was titrated by assessing inhibition of hemolysis of

antibody-sensitized sheep erythrocytes (ShEA) exposed to pooled normal human serum (pNHS), an assay where target hemolysis is absolutely dependent on MAC formation (Fig. 1A). At 10 µg/ml, OmCI caused complete inhibition of pNHS-induced hemolysis. This dose was used in all subsequent experiments. The sensitivity of each of the selected tumor cell lines, CT26 and B16, to pNHS-induced complement-directed cytoxicity was determined in a calcein release assay immediately before each experiment. Both CT26 and B16 cells were efficiently killed by pNHS without need for sensitization, calcein release correlating with dose of pNHS (Fig. 1B). The dose of pNHS causing <10% specific calcein release at 1 h was chosen as the maximum sublytic dose for the subsequent experiments. The 1 h time point was chosen based on our previous work showing that MAC killing of nucleated cell targets is an acute event and does not increase further with prolonged incubation (6). To identify MAC-specific effects, cells were also exposed to the same sublytic dose of pNHS but preincubated with the inhibitory dose of OmCI (10  $\mu$ g/ml) to block MAC formation.

Preliminary qPCR experiments were carried out on RNA harvested from CT26 cells used in the above experiments to validate targets for subsequent microarray expression analyses and determine optimum time points for RNA collection. Initially, RNA was harvested at 1 h and expression of osteopontin (*OPN*), a candidate gene chosen based on evidence from the literature (38), was measured by qPCR. Relative expression of *OPN* increased more in response to treatment with pNHS compared with OmCI-treated pNHS after 1 h exposure (Fig. 1*C*). To further refine the time points of exposure to sublytic MAC before RNA harvest, the experiment was repeated for 6- or 12-h time points, calculating the expression change in relation to



FIGURE 2. **Primary microarray data analysis.** *A*, PCA plot of top three principle components. Three-dimensional plot showing the top three principle components of the microarray data as calculated using PCA. Contributing principle components (*PC*) are labeled on each axis alongside the calculated % contribution to overall variation. Each sample from the experiment is represented by a colored sphere; *red* = control, *green* = pNHS, *blue* = pNHS+OmCl. A centroid *sphere* shows how these samples are grouped according to their experimental conditions; *black* = control, *pale blue* = OmCl at 1 h, *darker blue* = 0mCl at 12 h, *light green* = pNHS at 1 h, and *darker green* = pNHS at 12 h. *B*, scatter plot comparisons between samples exposed to pNHS and pNHS+OmCl at 1 (*i*) and 12 (*ii*) hours. Data are log<sub>2</sub> transformed, median baseline-adjusted. Expression is presented as distribution around a median that represents equal gene expression in the two conditions. The parallel flanking lines represent gene expression changes of ± 1.3-fold change; data points falling outside these lines are considered to be differentially expressed. Data points are colored according to their expression levels (median baseline-adjusted) upon exposure to pNHS; *green* = below median, *red* = above median.

untreated cells. *OPN* expression increased significantly at 6 h (p < 0.01) and 12 h (p < 0.05) in MAC-exposed cells compared with the OmCI control (Fig. 1*D*). For expression analysis, we chose to use 1 h as an initial time point to capture an immediate response and 12 h to capture sustained changes.

Global Gene Expression Analysis of Sublytic C-For microarray analysis cells were exposed to sublytic C attack and compared with a MAC-inhibited control as established in the optimization experiments described above: CT26 cells seeded at  $1.6 \times 10^3$ /mm<sup>2</sup> were exposed to 5% pNHS with or without the addition of OmCI, then incubated for 1 or 12 h (4 replicates of each condition in wells of a 12-well plate) before harvest of RNA. RNA was also harvested from untreated control cells (four replicates) to establish a baseline. A total of 16 samples were used for microarray analysis, 3 from each of the 4 serum conditions and all 4 untreated controls. Quality control of microarray data was performed using principal components analysis (PCA), a method used to compress a high content dataset, enabling its description with a limited number of contributors to variation. PCA allows the effect of experimental parameters on the data to be observed and identifies data quality issues. Initial PCA showed a batch effect that was eliminated using the Partek batch remover tool to reveal the most important components (Fig. 2A). Data were plotted to explore the contributions of the top three components; PCA#1 and #3 were best correlated with time point and serum exposure and the presence of OmCI, respectively. Batch removal was not retained for downstream analysis because the robustness of the ANOVA model used rendered it unnecessary.

To better visualize the impact of experimental conditions, scatter plots of median baseline-adjusted data were prepared to compare pNHS to pNHS + OmCI at 1 and 12 h (Fig. 2B, i and ii). This graphics representation revealed that, for most parameters, expression changes were more apparent 1 h after sublytic attack than at 12 h, indicating that in this system most of the changes were transient in nature. To measure significance of differential gene expression in cells exposed to pNHS with or without MAC blockade, a two-way ANOVA model was applied using the method of moments (39). Gene lists were prepared using the ANOVA-generated -fold change and p values to identify the most significantly differentially expressed (up or down) genes at each time point (Table 1). These show those genes that altered expression significantly (p < 0.05), filtered to include genes changed by a >2-fold at 1 h or a 1.5-fold at 12 h; different filters were chosen to reduce the disparity in number of differentially expressed genes at these time points. Five genes were up-regulated and 1 was down-regulated at 1 h post-exposure, whereas 2 genes were up-regulated 12 h post-exposure with none down-regulated. This difference in the number of genes differentially regulated between the time points supports the trends shown in the scatter plots, substantially more points falling outside the set confidence intervals at 1 h compared with 12 h (Fig. 2B).

Identification of Secreted Effectors Induced by Sublytic C—To provide functional insight into the data a new gene list was prepared using less stringent thresholds for inclusion by applying the following filters to the ANOVA statistics: -fold change > 1 and an unadjusted *p* value cut-off < 0.05 for both 1 h



#### TABLE 1

Gene list of the top most significant expression changes when comparing pNHS with pNHS+OmCl at 1 and 12 h

List criteria: false discovery rate adjusted *p* value <0.05. Top: pNHS 1 h relative to pNHS+OmCl 1 h, up-regulated genes (-fold change >2) ranked by *p* value. Middle: pNHS 1 h relative to pNHS+OmCl 1 h, down-regulated genes (-fold change >2). Bottom: pNHS 12 h relative to pNHS+OmCl 12 h, up-regulated genes (-fold change >1.5) ranked by *p* value.

Probeset ID	Symbol	Transcript	p value (pNHS 1 h vs. OmCI 1 h)	-Fold change (pNHS 1 h vs. OmCI 1 h)
ILMN_2623983	Egr2	ILMN_212209	$4.57  imes 10^{-11}$	2.08117
ILMN_2600744	Rgs16	ILMN_209950	$3.03 imes10^-09$	2.74759
ILMN_2995794	Itprip	ILMN_242056	$6.49 imes10^-08$	2.16089
ILMN_2662926	Egr1	ILMN_215729	$9.47 imes10^-08$	2.37517
ILMN_2757634	Fam110c	ILMN_222981	$1.89  imes 10^-07$	2.07236
ILMN_2834777	Irf1	ILMN_209850	$5.15  imes 10^-10$	-2.19138
Probeset ID	Symbol	Transcript	p value (pNHS 12 h vs. OmCI 12 h 12)	-Fold change (pNHS 12 h vs. OmCI 12 h)
ILMN_1260323	Akr1c18	ILMN_215518	$4.16  imes 10^{-}07$	1.61755
ILMN_1225816	Hbb-bh1	ILMN_216390	$1.99 imes10^-07$	1.58676

and 12 h comparisons. This latter filter selected for MAC-induced up-regulation events that were apparent at both 1 and 12 h time points, representing sustained changes. To understand the interactions between these genes the list was interrogated using MetaCore network building tools to automatically map genes to a representative component termed the network object (NO). The "shortest path" algorithm was selected, and canonical pathways were included for network building; to interpret the resulting network, nodes were arranged to identify the starting NOs and their overlapping connections. NOs not connected to the main network were removed, and those groups displaying little connectivity to the larger network were pruned. The network was then organized by cellular location from top to bottom (Fig. 3). The analysis revealed four key highly interrelated NOs, co-regulated by canonical pathways and with roles outside of the cell. These represented four genes; AREG (encoding amphiregulin), MMP3 (encoding matrix metalloproteinase-3; mmp-3), MMP13 (encoding matrix metalloproteinase-13; mmp-13), and CXCL1 (encoding chemokine (CXC motif) ligand 1; cxcl-1) (Table 2). Other highly connected NOs of note were epidermal growth factor receptor (EGFR) and the AP-1 complex.

qPCR Validation of Identified Genes-Genes identified as differentially expressed by microarray were validated using qPCR to provide support for further bioinformatic and biochemical exploration of their relevance. The four genes identified above as secreted effectors during network analysis were selected (Table 2) together with four genes identified as showing the most significant differential expression (up or down at either time point) between MAC-exposed and MAC-inhibited pNHS-exposed CT26 cells when stringent thresholds were applied (Table 1). FAM110c (encoding family with sequence similarity 110, member C) and RGS16 (encoding regulator of G-protein signaling 16) were both increased at 1 h post-attack; IRF1 (encoding interferon regulatory factor 1) was decreased at 1 h; *HBB-BH1* (encoding hemoglobin Z,  $\beta$ -like embryonic chain) was increased at 12 h. For each of these eight genes, qPCR was performed twice, first on cDNA prepared from RNA extracted for the microarray experiment and second on RNA from a fresh replication experiment. The qPCR expression data were replicated in these experiments and largely confirmed the expression patterns found in microarray for these same genes; data are presented together for comparison (Figs. 4 and 5). In a few instances, qPCR and microarray data did not completely

replicate: *CXCL1* showed the highest up-regulation at 1 h by microarray but at 12 h by qPCR; *RGS16* peak up-regulation at 1 h in microarray was confirmed by qPCR using the same original RNA, but in RNA from the second experiment further up-regulation was seen at 12 h post exposure. Despite these minor differences, the data strongly correlated, confirming the capacity of the microarray to accurately detect expression changes.

To explore whether the observed expression changes were cell type-specific, RNA extracted from MAC-exposed and control B16 melanoma cells was analyzed by qPCR for expression of the four network identified hits, *AREG*, *MMP3*, *MMP13*, and *CXCL1*. Expression of MMP3 RNA was negligible in this cell type. Expression of RNA for both AREG and CXCL1 was markedly increased in MAC-exposed cells at both 1 and 12 h, replicating the results obtained in CT26 cells (Fig. 5). MMP13 RNA expression was low in B16 cells and not significantly different between MAC exposed and control cells.

Interconnectivity of Secreted Effectors and Regulatory Genes-In an effort to identify pathways and mechanisms by which MAC effected changes in expression of the identified genes, a model was developed that analyzed the combination of those genes identified as significantly differentially expressed under stringent statistics and those identified by network analysis as downstream secreted factors. Fig. 6 shows the shortest path network generated using the gene lists shown in Tables 1 and 2 as input. The main hub of the network contained 11 of the total 12 starting NOs; FAM110C, ITPRIP, and HBB-BH1 were unconnected and, therefore, hidden. The network shows that the starting 11 NOs are well connected with a central triangle containing EGR1, EGR2, and IRF1, suggesting they are key drivers of the gene expression response to MAC. NOs added by the algorithm included AP-1 transcription factor subunit c-JUN, several NF $\kappa$ B subunits, the glucocorticoid receptor- $\alpha$ , c-Myc, and the EGFR. AP-1 and NF $\kappa$ B are the only two transcription factor NOs connected to all four secreted NOs validated by qPCR.

*Transcriptional Regulation Network*—To interrogate the data further and explore transcription regulation patterns from a greater number of data points, a gene list was generated by applying a false discovery rate-adjusted *p* value cut-off of <0.05 to the ANOVA statistics for the pNHS *versus* pNHS+OmCI comparison. This was applied separately for 1-h and 12-h time points, and the two lists were combined to identify the most significant MAC-induced expression changes at either time





Unspecified

FIGURE 3. **Network analysis of overlap gene list.** The list includes genes up-regulated by sublytic MAC at both time points. The network was generated in MetaCore using the following options: shortest path network-building algorithm with a maximum of two steps, inclusive of canonical pathway; this latter option allows sequences of interactions that occur frequently in the cell to be counted as single steps in the shortest path. The network describes the interconnected regulation of up-regulated genes and highlights four key downstream effector genes. The network is organized so that nodes are organized by the subcellular localization of their products, from extracellular to nuclear. Nodes present in the input list are in *blue circles. Thick light blue lines* highlight the various canonical pathways of signal transduction and transcription regulation. Seed nodes are *circled in navy blue, lines* represent interactions, either transcriptional regulation or protein-protein associations; *red* = inhibition and *green* = activation.

## TABLE 2

#### Gene list entities

Four key highly interrelated NOs, co-regulated by canonical pathways and with roles outside of the cell were identified; these represented four genes, *AREG* (encoding amphiregulin), *MMP3* (encoding mmp-3), *MMP13* (encoding mmp-13), and *CXCL1* (encoding cxcl-1). These genes were sorted by *p* values at 1 h and 12 h, and -fold change display (pNHS relative to pNHS+OmCI).

Probeset ID	Symbol	Transcript	<i>p</i> value (pNHS 1 h <i>vs.</i> OmCI 1 h)	-Fold change (pNHS 1 h <i>vs</i> . OmCI 1 h)	<i>p</i> value -fold change (pNHS 12 h <i>vs.</i> OmCI 12 h)	-Fold change (pNHS 12 h <i>vs.</i> OmCI 12 h)
ILMN_1238547 II MN_2737685	Areg Mmp13	ILMN_217903	$2.31 \times 10^{-06}$ 1.06 × 10^{-06}	1.43899 1.52378	0.00185142	1.18086
ILMN_2753809	Mmp13 Cxcl1	ILMN_219123 ILMN_223377	$9.89 \times 10^{-}05$ $8.74 \times 10^{-}03$	1.23906	0.00367792	1.08968

point and regardless of direction. This new list was combined with that created to generate Fig. 3, and the entire gene list was uploaded to MetaCore. The list was used as the starting list for the Analyze Network (transcription factors) (AN(TF)) algorithm applied using the default settings.

The AN(TF) algorithm identifies transcription factors for which there are enriched numbers of targets in the starting list, then uses the list to find the shortest path back to a receptor for which there are ligands in the starting list, thereby creating networks for each transcription factor, ranked by significance, based on enrichment of starting NOs via calculated *g*-scores, *z*-scores, and *p* values. *z*-score indicates the saturation of starting NOs, the *g*-score is a modified *z*-score describing the number of canonical pathways used to build the network, and *p* value assesses the probability of the number of starting NOs falling on the generated network by chance accounting for the total number of NOs in the network and in the entire database (40).

The network with the highest *g*-score and *z*-score and smallest *p* value was selected: *g*- and *z*-scores = 187.12, *p* value =  $7 \times 10^{-211}$ . To assist in interpretation, the network was organized by aligning the most connected NOs in the center and placing the remaining NOs by protein class and in context around these main hubs (Fig. 7). With c-Myc and CREB1 as the main controlling transcription factors, EGFR and TrkB are introduced as non-seed nodes to the network as putative receptor starting points with EGFR the most interacting of the two. Other important TF hubs include c-Jun, p53, ESR1, and Oct3/4. All four secreted effectors identified in microarray are present, and the NO with most direct connections with these is c-Jun. Other





FIGURE 4. **qPCR validation of statistically significant hits.** *Microarray* = original microarray data, *Primary Validation* = RNA extracted in parallel with that used in microarray, and *Secondary validation* = RNA extracted in a fresh sublytic attack experiment. RNA was reverse-transcribed and *FAM110C*, *RGS16*, *IRF1*, and *HBB-BH1* gene expression analyzed by qPCR and calculated as expression relative to housekeeping genes  $\beta$ -actin and Pol2ra using the  $\Delta\Delta$ Ct calculation then presented as % of untreated control. Results are the means of three determinations ± S.E. \*, p < 0.05; \*\*, p < 0.01; \*\*\*, p < 0.001. *MBL adj.*, median baseline-adjusted. *Con*, control.



FIGURE 5. **qPCR validation of network identified hits.** *Microarray* = original microarray data, *Primary Validation* = RNA extracted in parallel with that used in microarray, and *Secondary validation* = RNA extracted in a fresh sublytic attack experiment. *B16 validation* = RNA extracted from fresh sublytic attack experiment using the B16 mouse myeloma cell line (*MMP3* message was not significantly detected in this cell line). In all cases RNA was reverse-transcribed, and *MMP3*, *MMP13*, *CXCL1*, and *AREG* gene expression was analyzed by qPCR and calculated as expression relative to housekeeping genes  $\beta$ -actin and Pol2ra using the  $\Delta\Delta$ Ct calculation then presented as a % of untreated control. Results are the means of three determinations ± S.E. (\*, p < 0.05; \*\*, p < 0.01; \*\*\*, p < 0.001). *MBL adj.*, median baseline-adjusted; *Con*, control.

signaling molecules with high connectivity include AKT and ERK1/2. The network contains six of the eight validated genes. Overall the AN(TF) network includes 67 of the 118 candidate objects and provides evidence for a central role of EGFR activation by sublytic MAC.

## Discussion

The role of C as a tumor promoter has attracted a great deal of attention over the last few years because of evidence for significant C activation in diverse tumors. MAC is suspected to be influential given its published activating and proliferative effects on nucleated cells (5, 6, 14); however, signaling mechanisms underlying many of these effects remain ill defined. We took a novel approach to understanding the role of the MAC, taking advantage of an available terminal pathway inhibitor, global gene expression technology, and systems biology methodology. Sublytic conditions were optimized using pNHS as a C source and OmCI to block terminal pathway activation. CT26 colon carcinoma cells were selected as a model tumor cell, and MAC-specific gene expression changes were mapped by microarray, qPCR, and network analysis. These approaches revealed a gene expression pattern in tumor cells exposed to sublytic MAC, which could significantly impact cell survival and proliferation as well as reshape surrounding extracellular matrix. The key findings were replicated in an unrelated tumor cell line, B16 melanoma.





FIGURE 6. **Network analysis of collated gene list.** Network describing the interconnected regulation of the four key downstream effector genes as well as the eight statistically significant genes. The network was generated in MetaCore using the following options: shortest path network building algorithm with a maximum of two steps, excluding canonical pathways. The network is organized to show cellular localization from extracellular (*top*) to nucleus (*bottom*). From the list, *ITPRIP, FAM110C*, and *HBB-BH1* are not represented on the network due to lack of connectivity. Seed nodes are *circled in navy blue*, and *lines* represent interactions either transcriptional regulation or protein-protein associations; *red* = inhibition, and *green* = activation.

Statistical analysis of array data comparing MAC-exposed cells with controls confirmed a set of expression changes; genes involved in Ca<sup>2+</sup> and G-protein signal transduction (*ITPRIP*, *RGS16*), early response transcription factors (*EGR1*, *EGR2*), and inflammatory responses (*IRF1*) were significantly altered. Network analysis to map the interactions of genes up-regulated at the 1-h and 12-h time points highlighted 4 further expression changes in genes encoding proteins with extracellular localization, *AREG*, *CXCL1*, *MMP3*, and *MMP13* genes; these were co-regulated by putative canonical signaling cascades including PKC, PI3k/AKT, JNK, Erk1/2, and p38.

The product of the *AREG* gene is the amphiregulin protein), an EGF-like ligand capable of triggering erbB2 activation (41). CXCL1 ligand is a potent neutrophil chemoattractant, important in infection and signals via CXCR2, a G-protein-coupled receptor (42). *MMP3* and *MMP13* both code for matrix metalloproteinases (MMPs), which function in extracellular matrix regulation and remodeling; they are important during development, wound healing, proliferation, and inflammation (43).

Validation by qPCR confirmed that the expression changes highlighted in gene array and supported by statistics and network building were real and robust. Critically, expression changes for two of these genes, *AREG* and *CXCL1*, closely replicated in an unrelated tumor cell line, B16. MMP expression in this line was extremely low so changes could not be replicated.

Network generation using these and the remaining statistically significant changes found interactions between all identified genes apart from *FAM110C* and highlighted the central importance of *IRF1*, *EGR1*, and *EGR2* in mediating the changes. EGFR is noteworthy in that it is placed in between two starting NOs, amphiregulin and Rgs16, the three connected in an extracellular to nuclear direction, supportive of EGFR activation. Rgs16 is further connected to Egr1, Egr2, and Irf1 via c-Myc. Egr1 protein is known to positively regulate *EGR2* gene expression, whereas Egr2 protein negatively regulates *EGR1* (44, 45); Egr1 protein is reported to inhibit *IRF1* expression (46). AP-1 and NF $\kappa$ B were both highlighted as possible transcriptional regulators in the network. The AP-1 and NF $\kappa$ B complexes are known to regulate *MMP3*, *MMP13*, and *CXCL1* gene expression in mouse and human cells (47–52).

Ap-1 (a heterodimer of *c-fos* and *c-jun*) and NF $\kappa$ B are known to be responsive to MAC (53) and have been cited as important regulators of the response to sublytic C (6). In particular, *c-fos* is up-regulated rapidly in MAC-exposed cells and is linked to





FIGURE 7. **Network of transcriptional regulation and ligand receptor signaling.** This network uses a greater number of genes, including all those identified as significantly changed by combining significantly differentially expressed genes with those significantly up-regulated at both time points. The network was generated in MetaCore using the following options: AN(TF)-building algorithm with "add ligands and TF targets" selected. The algorithm generates a list of possible networks, with scores based on the number of seed nodes to non-seed nodes and the presence of canonical pathway threads. The network with the highest score for these two factors was selected and manually organized, first showing the four validated genes and then the most highly connected objects regardless of functional type to their *right*. The remaining objects were sorted by function so that TFs, kinases, phosphatases, generic proteins, and binding proteins were from *left* to *right*. Ligands and receptors were placed to the *far left*. Seed nodes are *circled in navy blue*, the predicted receptor trigger is highlighted in *red*, *lines* represent interactions either transcriptional regulation or protein-protein associations; *red* = inhibition and *green* = activation.

 $Ca^{2+}$  flux and MAPK (particularly ERK) involvement (54). *MMP3* and *MMP13* up-regulation has been described in MACattacked chondrocytes in human disease and an experimental model of osteoarthritis (55). Each of the canonical signaling cascades identified in the network, PKC, PI3k/AKT, JNK, Erk1/2, and p38 have been reported to be activated in cells exposed to sublytic MAC (6).

The larger network generated using genes significantly changed either at 1 or 12 h exposure alongside those up-regulated at both time points provided greater insight into mechanisms responsible for observed gene expression changes in CT26 cells. The network placed EGFR central in the response, an assignment supported by the presence of this receptor in all three generated networks (Fig. 7). EGFR is a member of the erbB receptor-tyrosine kinase family, activated by ligand binding at the cell surface triggering phosphorylation of the intracellular tyrosine kinase domain (56). Activation of the EGFR system has also been described in response to cellular stressors such as UV, osmotic, and oxidative stress (57). MAC may cause analogous stress responses; indeed, there is evidence that it can induce expression of the EGF ligand and cause EGFR signaling activation without ligand binding (58). The response may involve  $G\alpha_i$  protein activation independent of receptor, which is known to be activated by MAC (59). Indeed, our data showing an *RGS16* expression response to MAC supports this assertion; RGS16 gene expression is induced as a feedback mechanism for G-protein signaling (60). MAC can transactivate several other

receptor-tyrosine kinases, including fibroblast growth factor receptor-2 (FGFR2) and hepatocyte growth factor receptor (HGFR) (58). Potentiation of EGFR activation may come from MMP cleavage and release of EGF family ligands such as heparin-binding EGF-like growth factor (HB-EGF) at the cell surface, a pathway supported by our demonstration of *AREG* upregulation (61).

Increased expression of AREG, CXCL1, MMP3, and MMP13 is described at mRNA and protein levels in a number of human cancers such as breast, colorectal, ovarian, and pancreatic (62-69). This increased expression of the four effector genes often correlates with tumor development and aggressivity and can be predictive of patient prognosis (70-72). Their activities promote cell proliferation, activation, and motility through various mechanisms. AREG (amphiregulin) contributes to tumorigenesis via its function as a growth factor, the development of autocrine or juxtacrine loops that promote cell proliferation and survival, and increased cell motility (63, 71), and CXCL1 acts through recruitment of myeloid-derived suppressor cells to the tumor microenvironment where they support tumor growth and metastasis and suppress the local immune response (70). MMP family members contribute through their role in regulating the extracellular matrix (73), promoting angiogenesis, tumor invasion, and metastasis (72). In addition they cleave and activate molecules in the extracellular matrix that promote proliferation, motility, and induce alterations in adhesion (74). Increased expression of MMPs is associated with poor tumor

differentiation, increased invasiveness, poor prognosis, increased likelihood of metastasis, and shorter survival time (72). In particular, MMP3 promotes tumor progression by releasing/activating E-cadherin, L-selectin, HB-EGF, and TNF $\alpha$  and is described as a central mediator of mammary tumorigenesis (69). MMP3 is reported to induce a stable epithelial to mesenchymal transition, a process that is closely linked to tumor development (75). MMP13 contribution to tumor promotion is mainly through its pro-angiogenic activity, increasing vascular density in tumors (76). Together, alterations in the expression of these four genes represent a powerful influence on tumor development. Induction of expression of these genes in response to MAC may, therefore, indicate a tumor-promoting role.

A role for C as a tumor promoting system has recently gained mainstream recognition (19). The work presented here represents a novel approach to uncover this relationship using global expression data and systems biology analysis to explore both the mechanisms and the characteristics of such a response. The approach has provided evidence to suggest that MAC deposition, which does not result in cell lysis, is a potent tumor cell activator leading to significant changes in gene expression in several critical and interlinked pathways. These data fit well with published piecemeal studies in diverse cell types. The work not only sheds light on the signaling cascades and responsive transcription factor systems that respond to MAC but also reveals a downstream gene expression response to MAC that will alter tumor behavior through induction of proliferative, migratory, and survival pathways. Interestingly, a central role for the EGFR system was identified, although it was not clear whether this was activated directly by MAC or indirectly after MAC exposure. Overall, this work provides additional evidence implicating sublytic MAC in tumor cell activation and has implications not only for our understanding of the tumor promoting effects of C but also for new approaches to cancer therapy.

## **Experimental Procedures**

*Materials*—Pooled pNHS was obtained from whole blood collected from consenting volunteers. Blood was placed in 20-ml glass vials and allowed to clot. Serum was separated by centrifugation, pooled, 0.22- $\mu$ m-filtered, and stored in aliquots at -80 °C. Sheep erythrocytes (ShEs) in Alsever's solution were purchased from TCS Biosciences (Buckingham, UK). Complement fixation diluent (CFD) was from Oxoid (Basingstoke, UK). Anti-ShE antiserum (Amboceptor) was from Siemens (Forchheim, Germany). CT26 mouse colon carcinoma and B16 mouse melanoma cell lines were from American Type Culture Collection (Manassas VA). RPMI 1640 medium (RPMI), fetal bovine serum (FBS), and additives and calcein-AM were obtained from Invitrogen. Complete medium comprised RPMI 1640 with 5% heat-inactivated FBS. All other chemicals were from Sigma.

Preparation of ShEA—ShEs were washed into CFD and resuspended at 4% (v/v) in CFD (10 ml total volume) at 37 °C. Amboceptor, diluted 1:2000 in 10 ml of CFD, was mixed with the ShE suspension and incubated for 30 min at 37 °C. The resultant antibody-sensitized sheep erythrocytes were washed and diluted to 2% (v/v) in CFD.

## Complement Membrane Attack and Tumorigenesis

*Hemolytic Assay*—Hemolytic activity in pNHS was used as a measure of MAC formation, assessed by incubating (37 °C for 60 min) triplicate serum dilutions in CFD with an equal volume of ShEA in wells of a 96-well plate. No serum (CFD alone) and 100% lysis (CFD containing 0.1% Triton-X-100) controls in triplicate were included. Plates were spun, supernatant was transferred to a flat-bottomed 96-well plate, and absorbance measured at 410 nm using a FLUOstar OPTIMA plate reader (BMG Labtech, Aylesbury, UK). Percentage hemolysis was calculated using the following equation where FI indicates fluorescence intensity,

% Lysis = 
$$\left(\frac{A^{\text{complement release}} - A^{\text{spontaneous release}}}{A^{\text{detergent release}} - A^{\text{spontaneous release}}}\right) \times 100$$
(Eq. 1)

To titrate the effect of the C5 inhibitor OmCI (a gift from Dr. Miles Nunn) on MAC formation and hemolytic activity, aliquots of pNHS were preincubated with different doses of OmCI before measurement of hemolysis as above.

Complement-directed Cytoxicity Assay-We chose the welldescribed calcein release assay to measure tumor cell killing. The cell-permeant calcein AM is taken into cells and trapped by de-esterification to calcein; release of calcein from the cells then correlates with lytic cell death. CT26 cells or B16 cells were grown as monolayers in complete medium to 80% confluence in 75-cm<sup>2</sup> tissue culture flasks then washed in saline and harvested by incubation in 10 mM EDTA in PBS (30 min). Harvested cells were washed in RPMI, diluted to a density of  $5 \times 10^5$  cells per ml in complete medium, aliquoted at 100  $\mu$ l/well into flat-bottomed 96-well plates, and incubated for 16 h at 37 °C in 5% CO<sub>2</sub>. Adherent cells were washed, and 100  $\mu$ l of complete medium containing 2  $\mu$ g/ml calcein AM was dispensed into each well. Plates were incubated for 1 h at 37 °C in 5% CO2. Calceinloaded cell monolayers were washed twice in RPMI, then pNHS dilutions (0-40% in 100  $\mu$ l of RPMI) were dispensed directly into wells and incubated for a further 1 h at 37 °C in 5% CO<sub>2</sub>. Supernatants were transferred to fresh 96-well plates, and fluorescence was measured (excitation, 485 nm; emission, 520 nm) in a Fluostar Optima plate reader. The remaining cells were lysed by the addition of 100  $\mu$ l of 0.2% Triton-X-100 in RPMI per well, and released fluorescence was measured as above. Percentage lysis was calculated using the equation FI = fluorescence intensity,

% Lysis = 
$$100 \times (Fl^{complement}/(Fl^{complement} + Fl^{detergent}))$$
(Eq. 2)

Titrating Sublytic C Attack—CT26 or B16 cell monolayers were washed with saline then incubated (37 °C, 5% CO<sub>2</sub>, 1 or 12 h) with pNHS at a dilution previously titrated to give <10% lysis at 1 h, a time point when maximum lytic killing has been reached, either in the presence or absence of a dose of OmCI that completely inhibited hemolytic activity (10 µg/ml). Monolayers were washed in RPMI, and RNA was harvested using the Genelute Mammalian Total RNA Miniprep kit (Sigma).

Global Gene Expression Analysis—RNA concentration and quality were measured using the Agilent 2100 Bioanalyzer (Agi-



lent Technologies, Stockport, UK), and global gene expression analyses performed on the Illumina Microarray platform (Illumina, Saffron Walden, UK; Cardiff University Central Biotechnology Services). Amplification of material to generate cRNA and labeling was carried out according to manufacturer's instructions. Hybridization experiments were performed using the mouse ref8v2 BeadChips (2  $\times$  8 samples) and analyzed using the iScan Reader and Control Software (Illumina). GenomeStudio Expression Module software (Illumina) was used to convert signal intensity data into expression data. Data were normalized using the quantile method and log-transformed (77). PCA and primary statistical analysis were performed using PartekGenomics Suite version 6.6 (build 6.13.0213, Partek Inc., Chesterfield, MO), and graphic representations were obtained using GeneSpring 12.0 GX (Agilent Technologies). Pathway analysis was performed using Meta-Core software (Thomson Reuters, London, UK).

*qPCR Analysis of mRNA*—To validate differences in the relative expression of genes of interest implicated from gene expression analysis, extracted RNA samples (1  $\mu$ g) were reverse-transcribed using TaqMan Reverse Transcription Reagents (Applied Biosystems, Paisley, UK) following the manufacturer's instructions; resulting cDNA was stored at -80 °C. For qPCR, sufficient cDNA for triplicate reactions of each primer pair diluted  $\frac{1}{10}$  in ultrapure H<sub>2</sub>O, was mixed with 1× SYBR Green Jump Start Readymix (Sigma) according to manufacturer's instructions. Reaction mixes were aliquoted into 48-well white PCR plates, sealed with optical flat 8-cap strips (Bio-Rad), and placed in a MiniOpticon Real-Time PCR System (Bio-Rad) controlled using the Opticon Monitor 3.1 software. Thermocycling was adjusted from the manufacturer's protocol (65 °C annealing temperature and 40 cycles) to take account of relative expression, assessed using the  $\Delta\Delta C_t$  method where calculated C<sub>t</sub> was the cycle number at which fluorescence crossed a threshold level selected as the point where PCR expansion was linear in all samples. The mean C<sub>t</sub> values for the housekeeping genes  $\beta$ -actin and Polr2A were assessed,  $\Delta\Delta C_t$  was calculated for each sample, and results are expressed as a percentage of the control.

*Author Contributions*—B. P. M. and T. R. H. conceived the idea for the project. L. D. T. conducted most of the experiments and analyzed the results. R. A. W. performed confirmatory experiments with additional cell lines. B. P. M. and T. R. H. contributed to experimental design and data analysis at all stages. B. P. M., T. R. H., and L. D. T. contributed to writing of the manuscript.

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