Systems approach for the selection of micro-RNAs as therapeutic biomarkers of anti-EGFR monoclonal antibody treatment in colorectal cancer

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miRNA plays an important role in tumourgenesis by regulating expression of oncogenes and tumour suppressors. Thus affects cell proliferation and differentiation, apoptosis, invasion and angiogenesis. miRNAs are potential biomarkers for diagnosis, prognosis and therapies of different forms of cancer. However, relationship between response of cancer patients towards targeted therapy and the resulting modifications of the miRNA transcriptome in the context of pathway regulation is poorly understood. With ever-increasing pathways and miRNA-mRNA interaction databases, freely available mRNA and miRNA expression data in multiple cancer therapy have produced an unprecedented opportunity to decipher the role of miRNAs in early prediction of therapeutic efficacy in diseases. Efficient translation of -omics data and accumulated knowledge to clinical decision-making are of paramount scientific and public health interest. Well-structured translational algorithms are needed to bridge the gap from databases to decisions. Herein, we present a novel SMARTmiR algorithm to prospectively predict the role of miRNA as therapeutic biomarker for an anti-EGFR monoclonal antibody i.e. cetuximab treatment in colorectal cancer.

Colorectal cancer (CRC) is one of the most prevalent cancers, with 1.2 million new cases every year. It is the second most commonly diagnosed cancer in females and the third most common in males; the highest incidence occurs in developed countries. By the age of seventy years, one out of every two citizens in the Western world develops benign adenomas that evolve into malignant carcinomas at an estimated yearly rate of 0.1 to 0.25%. In United States alone, the annual cost of CRC treatment is forecasted to reach $17.7 billion by 2020. However, using simultaneous strategies that reduce risk factors, increasing screening and treatment could avoid 101,353 deaths resulting in $33.9 billion in savings in reduced productivity loss.

EGFR, a transmembrane receptor tyrosine kinase has been identified as one of the most promising targets for treating metastatic colorectal cancer (mCRC). Among the 20 molecules listed by the National Cancer Institute for the treatment of mCRC (http://www.cancer.gov/cancertopics/treatment/colon-and-rectal), cetuximab is one of the most successful monoclonal antibodies (http://www.croh-online.com/article/S1040-8428(13)00102-9/fulltext). However, multiple lines of evidence suggest that only 10–20% patients with mCRC benefit from cetuximab treatment. The selective efficacy, side effects and high treatment costs of cetuximab result in the need for focused research to decipher the resistance mechanisms to cetuximab. The response of the scientific community to this need is evident through the increased no. of publications suggesting that the mutational status of KRAS, BRAF and PIK3CA, differential expression of PTEN, EGFR ligand (AREG, EREG) and EGFR gene copy number variation could serve as therapeutic biomarkers for anti-EGFR monoclonal antibody treatment in CRC. However, none of these singular molecular changes could accurately predict the response of CRC patients to cetuximab therapy. Because CRC is a systems-level disorder that involves multiple molecular mechanisms to support proliferative signalling, resist cell death, induce angiogenesis and metastasis; molecules such as miRNAs that regulate signalling pathways by affecting the expression of multiple proteins might serve as more potent therapeutic biomarkers. However, little is known regarding the role of miRNAs as a therapeutic biomarker for cetuximab treatment in CRC.
miRNAs are small (18–25 nucleotide long) non-coding, highly conserved fragments of RNA. They post-transcriptionally regulate gene expression by complementary binding to the untranslated region of mRNA\textsuperscript{14,15}. As of November 2014; 1881 primary miRNA transcripts in the human genome has been annotated in the miRBase database (http://www.mirbase.org). miRNAs are transcribed from distant regions of genome previously annotated as protein coding part and have an independent transcription machinery\textsuperscript{16}. The primary miRNA transcripts i.e. pri-miRNA are processed inside nucleus by Drosha complex. Resulting precursor miRNA (pre-miRNA), a ∼60–70 nucleotide long hairpin structure is transported to cytoplasm where it further processed by Dicer. Successively mature miRNA formation and targeting of mRNA is achieved through RICS assembly\textsuperscript{4}.

miRNA expression is typically dysregulated in cancer cells and this dysregulation has a high degree of tissue specificity, miRNAs could be used as diagnostic and therapy-related biomarkers. Additionally, miRNAs have an unusually high stability in formalin-fixed tissues, from which they could be extracted with minimal degradation\textsuperscript{16,18}. Moreover, the techniques of miRNA analysis from a single cell are established, allowing for the analysis of small amounts of miRNAs with increasing sensitivity for potential biomarker assays\textsuperscript{17}. The roles of miRNAs in CRC, EGFR signalling regulation and cetuximab treatment outcome are also evident. Multiple reports regarding the miRNA dysregulation in metastatic colorectal cancer have been published. Some groups reported that major colorectal cancer biomarkers such as EGFR and RAS are regulated by mir-7 and let-7 respectively, thus profoundly affecting downstream signalling\textsuperscript{18–22}. Recently, Bissonnette et al. demonstrated the correlation between EGFR signalling and mir-143, mir-145 in murine colon cancer models\textsuperscript{23}. Another group discovered that cetuximab-mediated EGFR inhibition abrogates the age-related increase of miR21, which is related to age-dependent colorectal cancer\textsuperscript{18}.

These findings cumulatively suggest a potentially significant role of miRNAs in EGFR signalling in CRC. Therefore, these evidences compelled us to create a workflow to identify the most critical miRNAs important for cetuximab resistance in CRC that could be further used for potential biomarker assay development. In this study, we propose ranked miRNA candidates that might contribute to cetuximab resistance in CRC patients. The inference is based on the integration of multiple published and predicted miRNA findings into cellular pathways that lead to oncogenesis and metastasis. We prioritised the biomarker candidates based on a novel algorithm, i.e., SMARTmIR (Scoring-based MARking of Therapeutic MiCroRna), that combines the network parameters and literature-derived evidence. Finally, the significance of our prediction was strongly supported by recently published experimental data that are derived from cetuximab resistant CRC patients. This study provides an actionable insight into the novel mechanism of cetuximab resistance mediated by miRNAs that might lead to identification of miRNAs as biomarkers, thereby predicting optimum responses to the drug.

### Results

**Construction of molecular pathway maps crucial for cetuximab mode of action in CRC.** The efficacy of cetuximab trarement in responsive colorectal cancer is mostly depend on the state of four cellular processes namely apoptosis, proliferation and differentiation, angiogenesis, epithelial-to-mesenchymal transition/metastasis\textsuperscript{25}. First a comprehensive framework for the analysis and a structured overview of the bio molecular space for the mechanisms of action of cetuximab therapy in colorectal cancer was created by assembling four pathway maps which are representative of those cellular processes in CRC. The landscapes of the calculated network parameters for each of the four constructed pathways are presented in Figure 1. The top three nodes of each pathway (Jak2, Akt, p53 in apoptosis; SMAD, TGF-β, Shc in metastasis; VEGFR, VEGF, Src in Angiogenesis; EGFR, c-Fos, c-Raf in proliferation and differentiation) are well-known players in the corresponding processes\textsuperscript{12,25–27}. The high no. of the nodes representing receptors and kinases in all four pathway maps emphasise the crucial role of these molecules in oncogenesis and metastasis. Our maps also demonstrate that the roles of some molecules, such as transcription factors, in metastasis are better understood and more well known compared with the other three processes.

To understand the degree of the cross-talk and the overlap between the four pathway maps, we integrated all four pathway maps into a resulting \textit{integrated map}. Table 1 presents a statistical overview of the pathway maps and the percentage of literature validation of the nodes related to the corresponding cellular processes. A list of common nodes in all four pathways are provided in Supplementary Table S1.

The high percentage of literature validations (4th column of Table 1) confirms that the constructed pathways represent a good reflection of the current knowledge accumulated in the scientific literature. The metastasis pathway map is the largest in terms of the no. of nodes and edges, followed by angiogenesis, apoptosis, proliferation and differentiation. The integrated map consists of 465 nodes and 792 directed edges, demonstrating a large overlap among the molecules involved in the four processes. The literature validations of the node’s relationships to the cellular processes are provided in Supplementary Table S2.

To analyse the CRC specificity of the assembled pathway maps, we first explored the published TCGA (The Cancer Genome Atlas) data regarding the differential expression of the nodes’ mRNAs in colon and rectum adenocarcinoma\textsuperscript{28}. This analysis demonstrated that 96% of the nodes’ mRNAs are differentially expressed (more than two standard deviations from the mean) in at least one of the 244 studied tumour samples. In addition, we examined the association of the pathway maps’ nodes to colorectal cancer in the published scientific domain and determined that 45.3% of them (211/465) have been reported in the literature to have a relationship with CRC. All of those relationships are included in Supplementary Table S3. The differential expression of each mRNA in the TCGA tumour samples is provided in Supplementary Table S4.

**miRNAome screening for putative candidate biomarker.** Upon construction of comprehensive pathway maps and verification of their association with the four cellular processes and in CRC, we screened the published miRNAome for miRNAs targeting at least one node of the four pathway maps, which could serve as potential biomarkers for further analysis. In our screening, we considered experimentally validated miRNA target interactions (MTIs) and computationally predicted MTIs. To increase the chance of the predicted MTIs being biologically relevant, overlapping MTIs between three different miRNA-target prediction algorithms\textsuperscript{9,30} were used. One method that can be used to compare the quality of MTI prediction is to calculate the percentage of the prediction that has already been experimentally validated. In that direction, the percentage of experimentally validated MTIs is calculated for predictions from three individual algorithms, i.e., Pictar, miRanda, DianaMicroT, and from the intersection of their predictions. Experimentally validated MTIs are obtained from TarBase and Pathway Studio. As demonstrated in Figure 2, the percentage obtained from intersection of the three prediction algorithms is higher than that obtained using any of the three prediction algorithms used individually (Figure 2). Using the intersection of three prediction algorithms, we are able to capture 17.5% of the experimentally validated MTIs (average for four processes). However, the results for each of the prediction algorithms are less impressive: Pictar (3.9%), miRanda (3.4%), DianaMicroT (1.7%). Therefore, the assumption of using the intersection of the three prediction algorithms proved to be a better approach and was used.
for further analysis. Total number of experimentally validated MTIs for all prediction softwares and their intersect for all four pathway maps are provided in Supplementary Table S5.

The miRNAome screening revealed 335 miRNAs that target at least one node of the four assembled pathway maps. We further analysed the 335 miRNAs based on their ability to participate in all four processes, thus having a higher probability to be therapeutic biomarkers for cetuximab treatment in CRC patients. This analysis resulted in the selection of 188 miRNAs that interact with targets in all four of the pathway maps, and these selected miRNAs were used for the ranking and validation (Figure 3).

To evaluate the relationship of those 188 miRNAs to the four cellular processes and to the neoplasm, we performed a random sampling approach. We randomly selected 20 non-overlapping sets containing 5 miRNAs each (S1, S2, S3, S4, S5, S6, S7, S8, S9, S10, S11, S12, S13, S14, S15, S16, S17, S18, S19, S20 in Figure 4). Next the association between each miRNA (from each of those 20 samples) to four cellular processes (apoptosis, proliferation and differentiation, metastasis, angiogenesis) and to neoplasms were searched in the scientific literature. The number of miRNAs from each sample that are published to regulate at least one of those four cellular process and neoplasms are summarised in Figure 4. It is evident from Figure 4 that 74% of the miRNA candidates are known to regulate at least one of the four cellular processes and 51% of those miRNA are linked to different forms of neoplasms.

All 20 of the samples and the relationship between the miRNAs from each sample to cellular process and the neoplasm are provided in Supplementary Table S6, Supplementary Table S7 and Supplementary Table S8. Based on the node degree, the top 5 miRNAs from each of the four pathway maps are provided in Supplementary Fig S1.

### Prioritisation of the selected miRNAs.

Despite the potentially strong impact of the selected miRNAs on the fundamental molecular mechanisms underlying CRC, their relation to cetuximab treatment was not used in previous filtering of the 188 miRNAs,
and the resulting candidate list remains too long for experimental validation. To further prioritise miRNA candidates that have the highest likelihood of acting as therapeutic biomarkers for cetuximab treatment, we ranked the miRNA candidates by applying a newly designed SMARTmiR algorithm, as described in the Materials and Methods section. The details of the 10 top-ranked miRNAs, including their scores and evidence of their interaction with cetuximab, and the role of the targets are summarised in Table 2. According to our predictions, those ten miRNAs might serve as the best candidates for therapeutic biomarkers for cetuximab treatment in CRC patients. All 188 miRNA with its SMARTmiR score and Higo gene id of the targets are provided in Supplementary Fig S2.

Validation of the prediction based on published experimental results. The optimal method of validating any systems biology prediction is through experimental results. In our case, the analysis of differentially expressed miRNAs in cetuximab-resistant CRC tumour samples would provide such validation. Recently, a group at the University of Helsinki studied differential miRNA expression patterns of 33 cetuximab-treated patients with metastatic colorectal cancer31. That group tested the association of each miRNA with the overall survival (OS) by applying a Cox proportional hazards regression model, and published a list of the 60 most differentially expressed miRNAs in patients with an extremely poor prognosis (resistant patients). According to our analysis, 85% (51 of 60 miRNAs) of the resistant patient-derived differentially expressed miRNAs are present in our list of the selected 188 miRNAs. Moreover, five of the ten top-ranked predicted miRNAs were found to be highly differentially expressed in resistant patients, exhibiting 4.6- to 9.7-fold changes (Table 2). More studies with greater no. of patients are needed to further validate our prediction; however, we consider this initial evidence to be very encouraging and to prove the applicability of our methodology.

Discussion

In colorectal cancer, the effects of cetuximab are mediated through various molecular pathways, including the Ras-Raf-MAPK, PI3K-AKT, protein kinase C, STAT and SRC pathways25,32. Though the efficacy of cetuximab in responsive CRC patients is mainly manifested through reduction of cell proliferation and differentiation, inhibition of angiogenesis, prevention of epithelial to mesenchymal transition (metastasis) and induction of apoptosis33. However deciphering the cetuximab resistance mechanisms a number of alternative molecular pathways were recently reviewed34. The reactivation of pro-angiogenic factors (pMAPK, VEGF) leading to increased angiogenesis in CRC is one such resistant mechanism34. In agreement with those results, our top-ranked miRNA, i.e., miR-21, is a well-known angiogenesis regulator in both in vitro and in vivo models35. Moreover, cetuximab treatment affects the expression of miR-21 in
Another group suggested that the cetuximab resistance mechanism is a phenomenon caused by an increased rate of EGFR degradation and internalisation; switching towards alternate pathways for growth and survival of CRC resistant tumour cells. In resistant cells, EGFR is localised in the sub-cellular compartments such as endosome, mitochondria and nucleus. The overexpression of nuclear EGFR is linked to SFKs (SRC-family kinase) expression, modulating the up regulation of the PI3K/AKT pathway in cetuximab resistance. Mutations of KRAS are also connected to the increased activation of SFKs, affecting the MAPK, beta-catenin, STAT and PI3K/AKT pathways in CRC resistant tumours. However, the mechanism of cetuximab resistance in CRC remain poorly understood.

To address the issue, we attempted to create a comprehensive CRC specific molecular network snapshot leading to the four cellular processes of apoptosis, proliferation and differentiation, metastasis and angiogenesis, which are crucial for mode of action of Cetuximab therapy in CRC. Pathways from Metacore were integrated. The Metacore pathway knowledge base was selected because of the high experimental validation of molecular interactions in its pathways. Of the assembled Integrated map, 96% are differentially expressed in published CRC specific RNASeq data. Based on the well-accepted node removal algorithm, that high percentage makes the Integrated map CRC specific. miRNA-target interactions have been screened from prediction algorithms (miRanda, Pictar and DianaMicroT) and literature based knowledgebase (Tarbase, Pathway Studio). TargetScanS, Pictar and miRanda used individually, or in combination, provide a good balance between precision and recall. However, Maragkakis et al. demonstrated that Pictar predictions overlap more than 75% of the predictions obtained from the other tools.

Table 2 | Top 10 miRNAs along with their scores, expression values, MTI, expression of MTI and miRNA in cetuximab sensitive to resistant CRC patients

| miRNA   | Score | No of Targets | Hugo Gene Id of differentially expressed miRNA Targets | No of TF, KNS | miRNA Fold Change |
|---------|-------|---------------|-----------------------------------------------------|---------------|------------------|
| hsa-miR-21 | 3.57  | 33            | NOTCH1 (0.004868), CD47 (0.031247), BRCA1 (0.013018), APAF1 (0.012285) | 4, 3          | 8.1              |
| hsa-miR-34a | 2.21  | 28            | YY1 (0.029778), NOTCH1 (0.004868), CD47 (0.031247), BRCA1 (0.013018), MYC (0.002497) | 4, 2          | 4.6              |
| hsa-miR-145 | 2.15  | 33            | MKL2 (0.004586), CD47 (0.031247), BRCA1 (0.013018), MYC (0.002497) | 7, 2          | N/A              |
| hsa-miR-27a | 2.01  | 48            | MAPK14 (0.009214), APAF1 (0.012285) | 4, 9          | 5.8              |
| hsa-miR-17  | 1.85  | 35            | MAPK14 (0.009214), BRCA1 (0.013018), E2F1 (0.002186), EREG (3.41e-06) | 3, 8          | 9.7              |
| hsa-miR-155 | 1.73  | 29            | CD47 (0.031247), FOXO3 (0.012217) | 6, 4          | N/A              |
| hsa-miR-182 | 1.65  | 39            | MKL2 (0.004586), CD47 (0.031247), FOXO3 (0.012217) | 5, 2          | N/A              |
| hsa-miR-15a | 1.59  | 25            | BRCA1 (0.013018), PRKAR2A (0.021049) | 1, 4          | 6.2              |
| hsa-miR-96  | 1.53  | 38            | PLCB4 (0.015726), FOXO3 (0.012217) | 4, 5          | N/A              |
| hsa-miR-10a | 1.48  | 26            | NOTCH1 (0.004868), BRCA1 (0.013018), E2F1 (0.002186) | 2, 6          | N/A              |

1st column: top 10 miRNA candidates; 2nd column: calculated score based on developed scoring function; 3rd column: total no of miRNA targets in the "Integrated map"; 4th column: Hugo gene ID of miRNA targets those were differentially expressed in cetuximab sensitive compared with resistant CRC patients with significant p-values; 5th column: no of transcription factors (TFs) and kinases (KNSs) among the targets; 6th column: expression value of the miRNA in cetuximab sensitive compared with resistant CRC patients (KRAS, BRAF wild type) published by Mosakhani et al.
Our methodology has successfully predicted possible relationships between a miRNA and cetuximab; in the case of a top-ranked miRNA, i.e., miR-21, the relationship has previously been experimentally validated. This result again demonstrates the novelty and applicability of our methodology.

The accurate prediction of miRNAs that might serve as potential therapeutic biomarkers would be of great importance for patients and for the pharmaceutical industry. Herein, we developed a novel algorithm that facilitates the prediction of potential miRNA biomarkers based on the knowledge accumulated in the public domain. To our knowledge, there is only one published alternative methodology that uses literature-based evidence for drug-associated miRNA predictions. Recently, Rukov et al. launched PharmacomiR, a miRNA Pharmacogenomics database that uses the triplet sets consisting of a miRNA, a target gene and a drug associated with the gene to predict miRNAs that could serve as potential therapeutic biomarkers. We compared the performance of PharmacomiR to that of our methodology in predicting the pharmacogenomics role of miRNAs in cetuximab treatment. PharmacomiR predicted 1102 unique miRNAs (6975 redundant miRNAs as the initial output). Next, we calculated the overlap between the no. of predicted miRNAs with the published differentially upregulated miRNAs by Mosakhani et al. in cetuximab-sensitive and cetuximab-resistant CRC patients. Clearly, our methodology has a higher prediction accuracy, with 27.1% (51 of 188) compared with 4.44% (49 of 1102) for PharmacomiR, in identifying experimentally validated potential miRNAs as therapeutic biomarkers for cetuximab therapy in CRC patients. Unlike PharmacomiR, SMARTmiR could also rank each candidate miRNA based on a novel disease specific score, making our methodology more advanced in prioritising candidate miRNAs for further experimental validation. A methodological comparison SMARTmiR and PharmacomiR in Supplementary Table S9.

Conclusion

This novel knowledge-based modelling and mining algorithm, i.e., SMARTmiR, has provided a pathway-centric methodology that facilitates the prediction of pharmacogenomic role of miRNA. However, this algorithm has limitations. The inherent issue of pathway map building is that its completeness is limited to valid and available data. Therefore, the sensitivity and specificity of the translational methodology is a function of completeness of the presented interactome. Additionally, the developed pathway maps could not integrate the dynamic nature of miRNA regulations. Annotations and encoding of the transcripts of the genes in the pathway maps are absent. However, these are outstanding challenges in representing dynamic biological systems, and the scientific community must act to solve these issues. The method is generic and can be applied to model the role of miRNA as a therapeutic biomarker for targeted therapy in other disease indications. The authors are optimistic that the application of an optimised and fully automated version of the algorithm has the potential to be used as clinical decision support tool. Moreover, this research will also provide a valuable background for colorectal cancer investigators to identify critical miRNA regulatory elements for better understanding regulatory mechanisms or designing future experimental studies in colorectal cancer.

Methods

Herein, we propose a predictive algorithm, i.e., SMARTmiR, that combines knowledge and data-driven approaches to identify miRNAs contributing to the therapeutic effects of cetuximab in CRC patients. The algorithm consists of the following four steps (Figure 5):

Step 1: Construction of pathway maps leading to oncogenesis and metastasis in CRC.
Step 2: Identification of miRNA candidate biomarkers via miRNAome screening.
Step 3: Ranking of miRNAs based on accumulated evidence and the effects on the cellular process of CRC patients treated with cetuximab.
Step 4: Validation of the prediction based on experimental data.

Construction of molecular pathway maps leading to CRC oncogenesis and metastasis. Following the first step in the workflow as illustrated in Figure 5, we created a comprehensive bio molecular space for the mechanisms of action of cetuximab therapy in colorectal cancer. In doing so we assembled four pathway maps linked to four fundamental cellular processes in oncogenesis and metastasis; namely apoptosis, proliferation and differentiation, angiogenesis, and metastasis. These pathway maps integrate known pathways from Metacore (Thomson Reuters, New York, USA) and IntAct27,28. The four cellular processes and protein-protein interactions from IntAct, BioGRID and HPRD (Human Protein Reference Database) Databases29,30. The pathways that were integrated to build the four pathway maps leading to the cellular processes are listed in Table 3. All of the molecules in the maps (nodes) are annotated with the Entrez gene ID, HGNC gene symbols and corresponding UniProt IDs. The distinct features of the pathways are as follows: (A) Node shapes and colours correspond to their functional category (e.g., receptor, ligand, transcription factor, kinase); (B) All of the edges have directionality; (C) The edges are differentiated by shapes and colour corresponding to the type of interaction (such as binding, catalysis, phosphorylation, transcription regulation, transformation etc.); (D) Reactions (edges) are tagged to PubMed IDs as evidence. Four pathway maps corresponding to proliferation and differentiation, apoptosis, angiogenesis, and metastasis are available in standardised SBML (Systems Biology Markup Language)31 for file exchange between different tools. The four pathways are provided as four Supplementary pathway maps (i.e., Apoptosis.xml, Proliferation_Differentiation.xml, Angiogenesis.xml and Metastasis.xml). The PubMed IDs supporting an edge can be found by selecting right click on the edge in the pathway map specific xml files after opening those in Cytoscape. TCGA RNA Seq data in colon and rectal adenocarcinoma (2012) has been analysed to demonstrate the CRC specificity of the constructed maps32.

Identification of miRNAs candidate biomarkers via miRNAome screening. In the second step, we identified miRNA-target interactions (MTIs) for all of the nodes in the four constructed pathway maps. The exhaustive search for these miRNAs included both experimentally validated miRNAs and predicted miRNAs.

Experimentally validated and literature reported miRNAs. Through screening public and commercial sources for experimentally validated MTIs, we identified two major resources that are widely used because of their extensive coverage and the quality of scientific evidence, i.e., TarBase and Pathway Studio (Elsevier, Amsterdam, Netherlands)33. TarBase hosts manually curated MTIs that are experimentally validated34. It also incorporates entries from other well-known databases, such as miRecords35, miRTarBase36 and miR2Disease37. All of the literature-derived MTIs that were extracted for our study from the two sources described above are included in Supplementary Table S10.

Predicted miRNAs. Several computer-aided algorithms are available for the identification of MTIs. We used Diana-Micro T 3.0, Pictar and TargetScan 6.2 to identify predicted MTIs for the four developed pathways, and to test what impact their overlapping predictions. These identified miRNAs have not been encoded into the pathway maps and are used to further filter and rank potential candidate miRNAs (described below).

Ranking of miRNAs based on accumulated evidence and their effect on the system. A novel raking formula was developed to rank the miRNAs for their potential to serve as therapeutic biomarkers for cetuximab treatment in CRC patients (Equation 1). To enhance the accuracy of the ranking function, the four assembled pathways were integrated in one Integrated map, representing a comprehensive
knowledge space of functional molecular networks that lead to the cellular process those are mostly dysregulated in colorectal cancer. Each miRNA is ranked based on its topological properties, network properties of its targets and based on literature-derived evidence of miRNA regulating signalling pathways that are important for colorectal carcinogenesis, miRNA target’s (i.e., nodes in the Integrated map) relations to cetuximab. All of the literature-derived evidence for the candidate miRNA’s relationships to cellular processes, CRC and the target’s relations to cetuximab are provided in Supplementary Table S11, Supplementary Table S12 and Supplementary Table S13; respectively. The network parameters (Betweeness centrality, Node degree) were calculated using the CentiScaPe plugin57.

\[ S_{mi} = \text{deg}(\text{mir}) + \sum_{i=1}^{n} \frac{R_i + T_i}{T_i} + p_{\text{Pathway}} + p_{\text{CRC}} + \sum_{i=1}^{n} p_{\text{Cetuximab}} \]  

(Equation 1)

\[ S_{mi} \] is the score of \( j \)-th miRNA. Each feature is given equal weightage due its natural importance and is normalised between 0 and 1 such that the calculated score could have a maximum value of 5. The overall scoring function was implemented in the Perl programming language.

The first feature of miRNA ranking is the node degree of the \( j \)-th miRNA, i.e., the no. of targets of the \( j \)-th miRNA in the Integrated map, as defined as the \( \text{deg}(\text{mir}) \). The node degree corresponds to the no. of nodes adjacent to a given node \( v \). The degree allows for an immediate evaluation of the regulatory relevance of the node. For example, in signalling networks, proteins with a very high degree are interacting with several other signalling proteins and are likely to be regulatory hubs. The second parameter was calculated based on the weighted node degree of all of the targets of \( j \)-th miRNA, depicted as \[ \sum_{i=1}^{n} \frac{R_i + T_i}{T_i} \text{Pathway}, \sum_{i=1}^{n} p_{\text{CRC}} \text{ and } \sum_{i=1}^{n} p_{\text{Cetuximab}} \].

**Table 3 | Detailed listing of the pathways used for assembling the proliferation and differentiation, apoptosis, angiogenesis, and metastasis processes**

| Cellular processes | Pathways assembled |
|--------------------|--------------------|
| Proliferation and Differentiation | EGFR signalling pathway, EGFR signalling via small GTPases, EGFR signalling via PI3, EPO-induced MAPK pathway, ERK5 in cell proliferation, PDGF signalling via MAPK cascades, PDGF signalling via STATs and NF-kB |
| Apoptosis | Apoptosis and survival, Anti-apoptotic TNFs/NF-kB/Bcl-2 pathway, Apoptosis and survival, Anti-apoptotic TNFs/NF-kB/IAP pathway, Apoptosis and survival, p53-dependent apoptosis, EPO-induced Jak-STAT pathway, EPO-induced PI3K, AKT pathway and Ca\(^{2+}\)-influx, Signal transduction of AKT signalling, Signal transduction of PTEN pathway |
| Angiogenesis | VEGF-family signalling and activation, Angiopoietin-Tie2 signalling, VEGF signalling via VEGFR2 - generic cascades, S1P1 signalling pathway Development Role of IL-8 in angiogenesis, Thrombospondin-1 signalling |
| Metastasis | TGF-beta receptor signalling, TGF-beta-dependent induction of EMT via MAPK, Regulation of epithelial-to-mesenchymal transition (EMT), Immune response of Oncostatin M signalling via JAK-STAT in human cells FGF2-dependent induction of EMT, TGF-beta induction of EMT via ROS, NOTCH-induced EMT, HGF-dependent inhibition of TGF-beta-induced EMT, TGF-beta-dependent induction of EMT via RhoA, PI3K and Il1K, Immune response of Oncostatin M signalling via MAPK in human cells, HGF signalling pathway |
target based on node degree, targeted by jth miRNA. Additionally, betweenness is calculated considering couples of nodes (v1, v2) and by counting the no. of shortest paths linking v1 and v2 and passing through a node n. The betweenness of a node in a biological network, such as a protein-signalling network, could indicate the relevance of a protein as functionally capable of holding together communicating proteins. Betweenness centrality was applied to identify the importance of a node in each of the four pathways (see the Results section). The additional features of SMARTmiR are literature-derived evidence reflecting the direct effect of each miRNA candidate in cancer aetiology, progression, spread and miRNA specific relationship to colorectal cancer. This third feature calculates the total amount of literature evidence (PubMed IDs) linking the jth miRNA to the four cellular processes relevant to tumour progression (proliferation and differentiation, apoptosis, metastasis, and angiogenesis) depicted as $E^\text{Pathway}_{i\rightarrow j}$. Similarly, the fourth feature calculates the total amount of literature evidence (PubMed IDs) that reflects the association of the jth miRNA to colorectal cancer, depicted as $E^\text{CRC}_j$.

The ultimate goal of SMARTmiR is to prioritise the role of a candidate miRNA as a therapeutic biomarker for cetuximab treatment in CRC. Therefore, the last feature, $\sum_{i=1}^n E^\text{Cetuximab}_{i\rightarrow j}$ calculates the total amount of literature evidence (PubMed IDs) of the jth miRNA target’s (ith) related to cetuximab. Separately we have also calculated the miRNA scoring with betweenness centrality of the nodes as second feature; maintaining first, third, fourth and fifth features of SMARTmiR identical. The resulting scoring is provided in Supplementary Table S14.

Validation of predicted miRNA biomarkers. We used a list of differentially expressed miRNAs in cetuximab sensitive and resistant CRC patients (KRAS and BRAF wild type) to validate the miRNAs predicted and ranked by our workflow as significant therapeutic biomarker candidates for cetuximab treatment. To further substantiate the expression of targeted miRNAs by the most significant miRNA biomarkers, we also analysed the mRNA expression data. The mRNA expression data analysis was performed using the appropriate R software packages. In particular, we used the MASS normalisation method and applied the SAM (Significance Analysis of Microarray) package to identify differentially expressed miRNAs. All differentially expressed mRNA with its corresponding p-value has been marked in “red” in the fourth column of Table 2.

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**Author contributions**
A.D. and S.B. wrote the manuscript. A.D. prepared figure 1, 3–5; S.B. prepared figure 2. A.D., P.S., M.H.A. and N.N. designed the research. A.D., S.B., P.S., M.H.A. and N.N. reviewed the manuscript.

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