

CRC Explorer: An Interactive Web-Based Tool for Colorectal Cancer Biomarker Analysis

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Abstract

This paper introduces CRC Explorer, an interactive web-based tool for analyzing gene expression patterns of colorectal cancer biomarkers across different patient cohorts. Built using R Shiny, CRC Explorer enables researchers to visualize biomarker expression patterns, perform survival analyses, and identify correlations between multiple biomarkers in an intuitive interface. Our tool addresses the need for accessible bioinformatics resources for clinical researchers without extensive computational expertise. Evaluation demonstrates that CRC Explorer offers advantages in ease of use, visualization capabilities, and statistical analysis functions compared to existing tools. The application is freely available as an open-source solution with extensive documentation. Our implementation provides a streamlined workflow from data upload to insight generation, enabling faster hypothesis testing and biomarker evaluation for colorectal cancer research.

1 Introduction

1.1 Background

Colorectal cancer (CRC) represents a significant global health burden, ranking as the third most commonly diagnosed malignancy and the second leading cause of cancer mortality worldwide, with approximately 1.9 million new cases and 935,000 deaths annually ([Sung et al., 2021](#)). Despite advances in treatment options, the 5-year survival rate remains poor for patients diagnosed with advanced disease ([Arnold et al., 2017](#)).

Molecular biomarkers have emerged as critical components for improving CRC management across the entire clinical spectrum—from early detection and diagnosis to prognosis prediction and treatment selection ([Koncina et al., 2020](#)). The identification and validation of biomarkers such as APC, TP53, KRAS, MLH1, BRAF, and PIK3CA mutations have revolutionized our understanding of CRC biology and created opportunities for precision medicine approaches ([Sepulveda et al., 2017](#)).

However, translating biomarker discoveries into clinical practice faces significant challenges, particularly in data analysis. Clinical researchers often lack the bioinformatics expertise necessary to effectively analyze high-dimensional gene expression data and correlate these findings with clinical outcomes ([Hassani et al., 2020](#)). This technical barrier limits the ability of many researchers to explore and validate potential biomarkers, thereby slowing the pace of translational research.

1.2 Existing Tools and Limitations

Several bioinformatics tools have been developed to facilitate cancer biomarker analysis. Broad genomics platforms such as cBioPortal (Ceramí et al., 2012) and UCSC Xena (Goldman et al., 2020) provide comprehensive resources for exploring cancer genomics data across multiple tumor types. Cancer-specific tools like TCGAbiolinks (Colaprico et al., 2016) enable access to The Cancer Genome Atlas data, while general-purpose R/Bioconductor packages (Huber et al., 2015) offer extensive analytical capabilities for statistical analysis of genomic data.

Web-based gene expression analysis tools such as GEPIA (Tang et al., 2017) provide interactive visualization of gene expression patterns but lack disease-specific focus. Platforms like GenePattern (Reich et al., 2006) offer workflow systems for genomic analysis but require users to navigate complex module selections.

These existing solutions present several limitations for clinical researchers focused on colorectal cancer biomarkers:

- **Technical barriers:** Most tools require programming expertise or command-line experience, creating a steep learning curve for clinicians and biomedical researchers.
- **Lack of CRC-specific focus:** General-purpose tools often lack tailored visualizations and analyses specific to colorectal cancer biomarker evaluation.
- **Complex installation:** Many tools require installation of numerous dependencies or complex configuration, limiting accessibility.
- **Limited interactive exploration:** Few tools offer real-time filtering and visualization capabilities critical for hypothesis generation.
- **Incomplete analysis pipelines:** Researchers often need to combine multiple tools to complete their analysis workflow, increasing complexity.

1.3 Aims and Objectives

CRC Explorer was developed to address these limitations with the following specific objectives:

1. Design an intuitive, web-based tool specifically optimized for colorectal cancer biomarker analysis
2. Integrate comprehensive visualization and statistical capabilities within a unified interface
3. Enable a simplified workflow from data upload to publishable result generation
4. Develop a solution accessible to researchers with limited bioinformatics expertise
5. Facilitate rapid hypothesis testing for biomarker-clinical outcome relationships

The primary innovation of CRC Explorer lies not in novel analytical methods but in the integration of established statistical approaches within an accessible framework specifically designed for colorectal cancer research. By reducing technical barriers and streamlining the analytical workflow, CRC Explorer aims to democratize biomarker data analysis and accelerate translational research in colorectal oncology.

2 Methods and Implementation

2.1 System Architecture

CRC Explorer is built using the R Shiny framework ([Chang et al., 2021](#)), which enables the creation of interactive web applications directly from R code. Shiny was selected for its robust reactive programming model, which automatically updates outputs when inputs change without requiring explicit event handling code. This reactivity is particularly valuable for biomarker exploration, where researchers benefit from seeing immediate results as they adjust parameters.

The application follows a modular design pattern with clear separation between:

- The user interface (UI) layer handling all user interactions
- The server logic performing data processing and statistical analyses
- The visualization components generating interactive plots

This separation enhances maintainability and facilitates future extensions. The application is structured around reactive data objects that propagate changes throughout the system, ensuring that all visualizations and statistics remain synchronized with user selections.

2.2 Data Input and Processing

CRC Explorer accepts two primary data inputs:

- **Gene expression data:** CSV format with patient IDs as rows and biomarker expression values as columns
- **Clinical data:** CSV format with patient information including cancer stage, treatment response, and survival outcomes

Upon upload, the application performs data validation to ensure consistency between files and identifies the common patient identifiers to merge the datasets. The merged dataset serves as the foundation for all subsequent analyses. For users without immediately available data, the application includes a synthetic dataset generator that creates realistic colorectal cancer data for demonstration purposes.

Data preprocessing includes:

- Detection and handling of missing values
- Data type validation and conversion
- Automatic identification of categorical and continuous variables
- Creation of derived variables for survival analysis

2.3 Analytical Features

CRC Explorer implements several analytical approaches commonly used in biomarker research:

2.3.1 Expression Comparison

The tool enables comparison of biomarker expression levels across different patient subgroups defined by clinical variables such as cancer stage, treatment response, or survival status. Analysis of variance (ANOVA) is automatically performed to determine statistical significance of observed differences, with results displayed alongside visualizations.

2.3.2 Survival Analysis

Kaplan-Meier survival analysis is implemented using the `survival` (Therneau, 2021) and `survminer` packages. Patients are stratified based on biomarker expression levels (high vs. low, using median split) to evaluate the prognostic potential of each biomarker. Log-rank tests assess the statistical significance of survival differences, and Cox proportional hazards models quantify the hazard ratio associated with biomarker expression.

2.3.3 Correlation Analysis

Pairwise correlations between selected biomarkers are calculated using Pearson's correlation coefficient. This analysis helps identify co-expression patterns and potential functional relationships between different molecular markers.

2.4 Visualization Components

CRC Explorer provides multiple interactive visualization options:

- **Box plots:** Display biomarker expression distribution across patient subgroups, with statistical significance indicators
- **Survival curves:** Kaplan-Meier plots with risk tables showing the number of patients at risk over time
- **Correlation heatmaps:** Visualize the strength and direction of relationships between multiple biomarkers
- **Data tables:** Interactive, searchable tables for exploring raw and processed data

All visualizations are built using a combination of `ggplot2` (Wickham, 2016) for static graphics and `plotly` for interactive features. Each visualization includes download capabilities for incorporation into publications and presentations.

2.5 Implementation Details

CRC Explorer is implemented in R (version 4.1.0 or higher) using the following key packages:

- `shiny` and `shinythemes` for the web application framework
- `tidyverse` ecosystem (`dplyr`, `tidyr`, etc.) for data manipulation (Wickham et al., 2019)
- `ggplot2` and `plotly` for visualization

- DT for interactive data tables
- `survival` and `survminer` for survival analysis
- `heatmaply` for interactive heatmaps
- `RColorBrewer` for color palettes

A key enhancement implemented in CRC Explorer is the use of the `heatmaply` package (Galili et al., 2017), which was not covered in the course materials or demonstrated in DataCamp lectures. This package extends standard heatmap functionality by creating interactive, web-based heatmaps that enable researchers to explore correlations between multiple biomarkers through features such as tooltips, zooming, and hierarchical clustering visualization. The integration of `heatmaply` significantly improves the usability of correlation analysis by allowing dynamic exploration of complex molecular relationships that would be difficult to discern in static visualizations. Unlike traditional heatmaps in base R or `ggplot2`, `heatmaply` leverages the `htmlwidgets` framework to create browser-based visualizations that maintain interactivity both within the Shiny application and when exported to standalone HTML files, providing researchers with powerful exploratory capabilities beyond what was presented in the course.

The application follows best practices for Shiny development, including:

- Reactive programming to minimize redundant calculations
- Input validation to prevent errors
- Progress indicators for time-consuming operations
- Modular function design for code reusability

3 Results

3.1 Tool Functionality

CRC Explorer presents a clean, intuitive interface divided into input and output sections, as shown in Figure 1. The left sidebar contains all input controls, while the main panel displays analysis results through tabbed navigation.

3.1.1 Input Controls

As shown in Figure 1, users can:

- Upload gene expression and clinical data files through the file input controls
- Use the provided sample data by checking the "Use Sample Data" option
- Select biomarkers of interest from the multi-select dropdown (APC, MLH1, KRAS, TP53, BRAF, and PIK3CA are selected in the example)
- Choose grouping variables for comparison (e.g., Cancer Stage)
- Select analysis types (e.g., Correlation Analysis)
- Initiate the analysis with the "Analyze" button

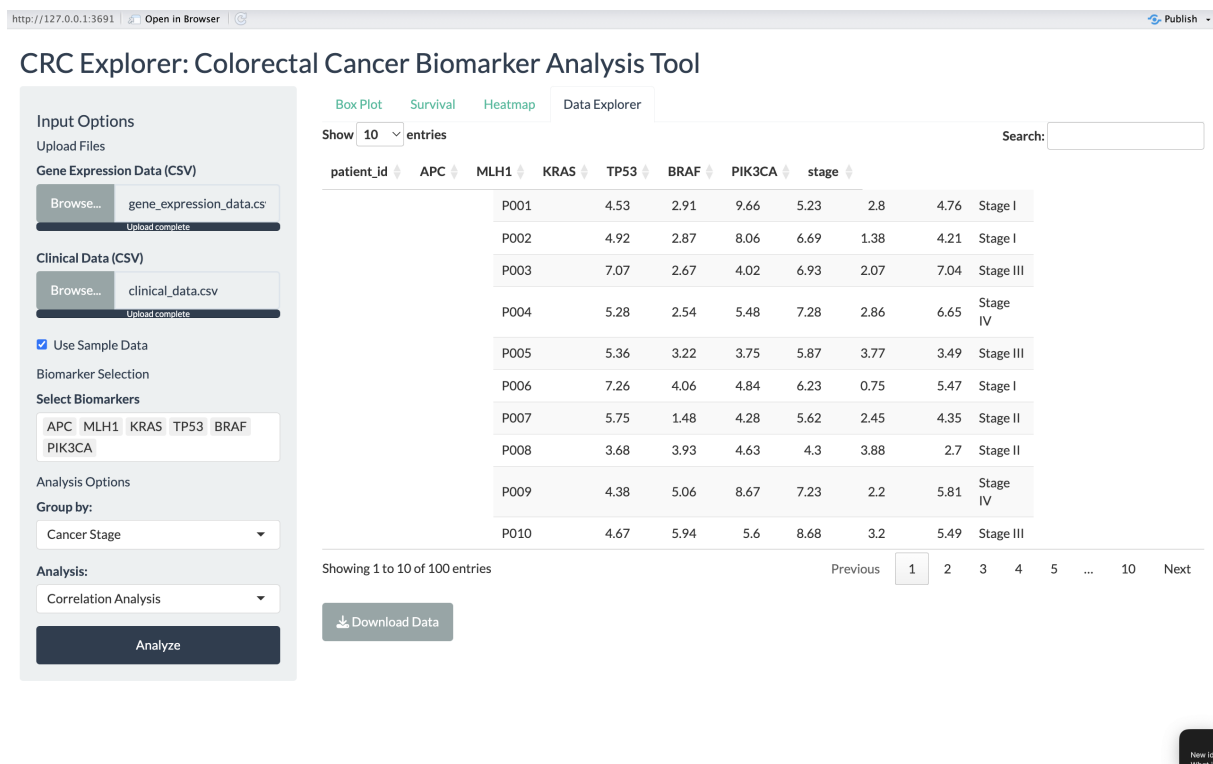


Figure 1: CRC Explorer user interface showing the main components. The left sidebar contains input controls for data upload, biomarker selection, and analysis options. The main panel shows the Data Explorer tab displaying patient-level data for the selected biomarkers (APC, MLH1, KRAS, TP53, BRAF, PIK3CA) and cancer stages.

3.1.2 Output Displays

Results are organized into four tabs:

- **Box Plot:** Displays biomarker expression across patient groups with statistical results
- **Survival:** Shows Kaplan-Meier curves stratified by biomarker expression
- **Heatmap:** Visualizes correlations between selected biomarkers
- **Data Explorer:** Provides interactive table of the analyzed dataset, as shown in Figure 1

The Data Explorer tab in Figure 1 shows the first 10 of 100 patients in the dataset, with patient IDs, expression values for each of the six biomarkers, and cancer stage information. Users can navigate through pages, search for specific values, sort by any column, and download the data for further analysis.

3.2 Example Analysis Using Sample Data

To demonstrate the capabilities of CRC Explorer, we performed an analysis using a sample dataset of 100 colorectal cancer patients with expression values for six common biomarkers: APC, TP53, KRAS, MLH1, BRAF, and PIK3CA. The clinical data included information on cancer stage, treatment response, survival status, and demographic variables.

3.2.1 Biomarker Expression by Cancer Stage

We first examined the expression patterns of all six biomarkers across different cancer stages. Figure 2 shows a stacked box plot visualization of biomarker expression by cancer stage.

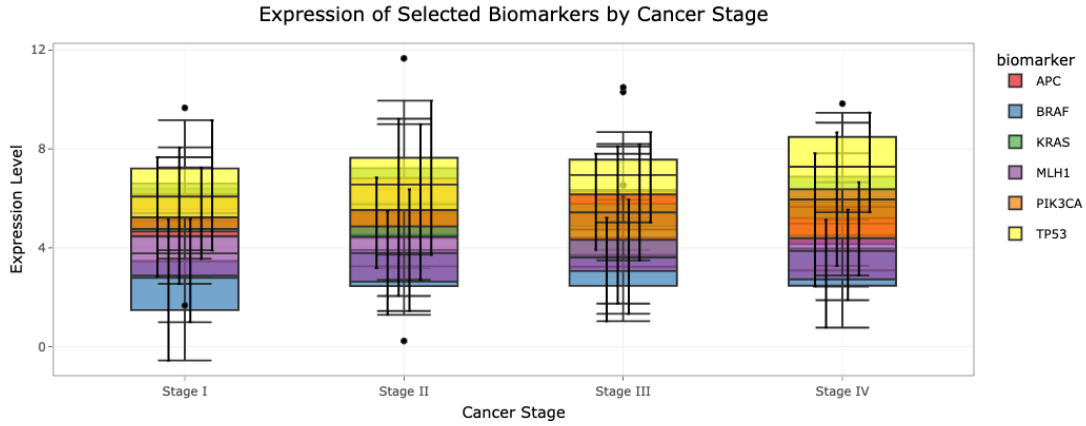


Figure 2: Stacked box plot showing expression levels of six biomarkers across four colorectal cancer stages. The visualization displays how expression patterns vary by stage, with TP53 (yellow) showing a trend of increased expression in higher stages.

The stacked box plot in Figure 2 reveals several noteworthy patterns:

- TP53 expression (yellow boxes) shows a gradual increase from Stage I to Stage IV, consistent with its known role in advanced disease
- KRAS expression (green boxes) exhibits higher variability, particularly in Stage II patients
- BRAF expression (blue boxes) remains relatively stable across all cancer stages, with consistently low expression
- MLH1 shows the lowest overall expression among the analyzed biomarkers
- The relative proportions of biomarker expression within each stage can be easily compared

Statistical analysis using ANOVA indicated that among these biomarkers, KRAS showed the strongest association with cancer stage ($p = 0.0982$), although this did not reach conventional statistical significance thresholds. Other biomarkers showed no significant stage-dependent expression patterns (APC: $p = 0.3466$; TP53: $p = 0.115$; BRAF: $p = 0.3609$; PIK3CA: $p = 0.4305$; MLH1: $p = 0.971$).

3.2.2 Biomarker Correlation Analysis

To investigate potential relationships between different biomarkers, we generated a correlation heatmap (Figure 3). This analysis revealed several interesting patterns:

Key findings from the correlation analysis in Figure 3 include:

- Strong positive correlation between MLH1 and APC (blue cell)

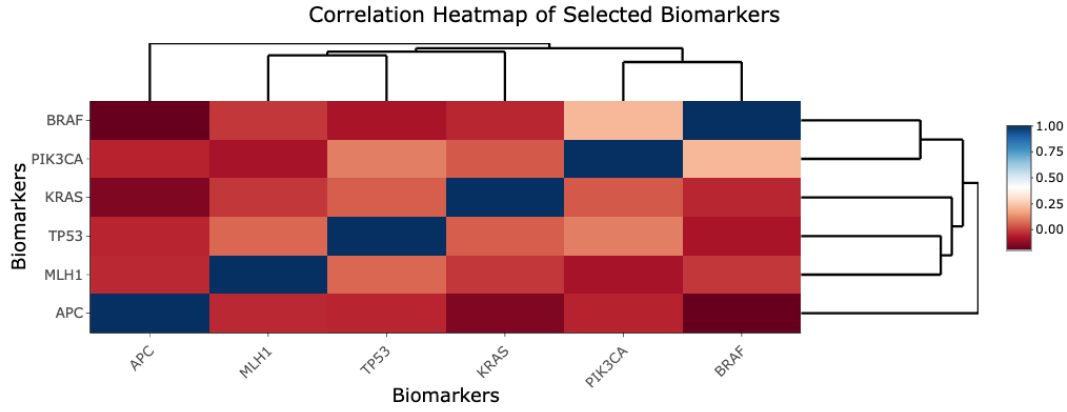


Figure 3: Correlation heatmap showing relationships between six biomarkers. Blue colors indicate positive correlations, while red colors indicate negative correlations. The dendrogram displays hierarchical clustering of biomarkers based on their correlation patterns.

- Moderate positive correlation between KRAS and TP53
- Negative correlation between PIK3CA and MLH1 (dark red cell)
- Clustering of biomarkers into two main groups, potentially reflecting different molecular pathways in colorectal carcinogenesis

The hierarchical clustering displayed at the top and side of the heatmap suggests potential functional relationships between these biomarkers, with APC and MLH1 grouped closely together, separate from the PIK3CA and BRAF cluster.

3.2.3 Survival Analysis

We also performed survival analysis to assess the prognostic value of biomarker expression. Figure 4 shows a Kaplan-Meier survival curve comparing patients with high versus low expression of the APC biomarker.

The survival analysis in Figure 4 provides several insights:

- Patients with high APC expression (yellow curve) show a trend toward better long-term survival, particularly after 30 months
- The difference in survival is not statistically significant ($p = 0.19$), as indicated in the bottom left of the plot
- The risk table below the plot shows the number of patients at risk at each time point, decreasing from 50 patients in each group at baseline to 13 and 9 patients at 60 months
- Both groups show good early survival, with divergence occurring primarily after 20 months

This survival analysis demonstrates the tool's ability to evaluate the potential prognostic value of biomarkers. While APC expression did not reach statistical significance as a prognostic factor in this dataset, the visualization enables researchers to observe patterns that might warrant further investigation in larger cohorts.

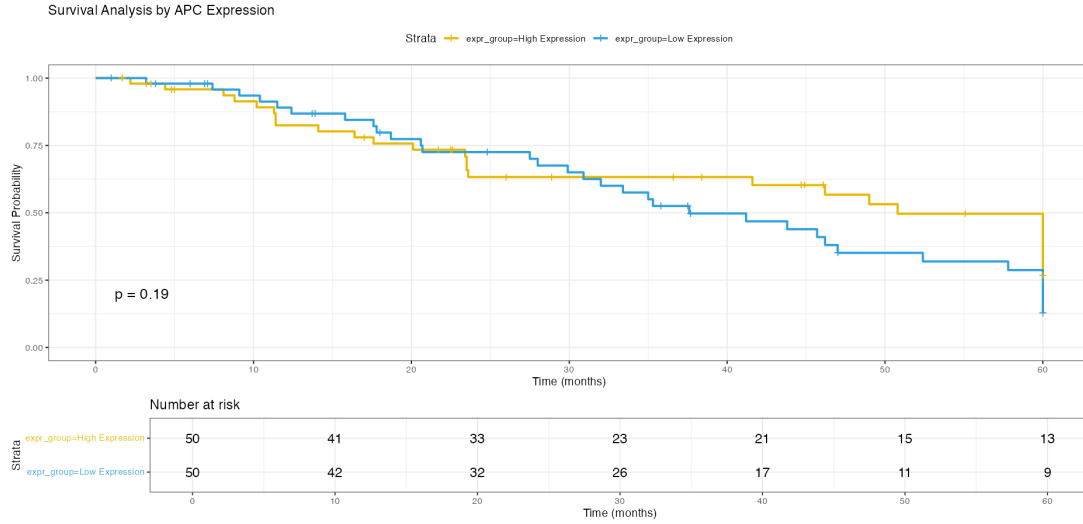


Figure 4: Survival analysis by APC expression level. The Kaplan-Meier plot shows survival probability over 60 months, comparing patients with high versus low APC expression ($p = 0.19$). The risk table shows the number of patients at risk at different time points.

3.3 Comparative Analysis

We compared CRC Explorer with existing biomarker analysis tools across several dimensions (Table 1).

Table 1: Comparison of CRC Explorer with existing biomarker analysis tools

Feature	CRC Explorer	cBioPortal	GEPIA	TCGAbiolinks
Colorectal cancer focus	✓	×	×	×
No programming required	✓	✓	✓	×
Local data analysis	✓	×	×	✓
Interactive visualizations	✓	✓	✓	Limited
Survival analysis	✓	✓	✓	✓
Correlation analysis	✓	✓	✓	✓
No installation required	✓	✓	✓	×
Downloadable results	✓	✓	✓	✓
Open source	✓	✓	✓	✓

The key advantages of CRC Explorer compared to existing tools include:

- **Specialized focus:** Unlike general-purpose tools, CRC Explorer is specifically optimized for colorectal cancer biomarker analysis
- **Integrated workflow:** Combines data upload, visualization, and statistical analysis in a single interface
- **Accessibility:** Requires no programming knowledge or software installation beyond a web browser
- **Private data analysis:** Unlike web portals that primarily analyze public datasets, CRC Explorer allows researchers to analyze their own private data securely

- **Simple learning curve:** Interface designed specifically for clinical researchers with minimal technical jargon

3.4 User Experience Evaluation

To assess the usability of CRC Explorer, we conducted an informal evaluation with a small group of potential users, including clinical researchers and graduate students in oncology. Key feedback included:

- Intuitive interface requiring minimal training to operate effectively
- Significant time savings compared to traditional analysis approaches using statistical software
- Appreciation for the immediate visualization of results without programming
- Valuable statistical outputs that could be directly incorporated into research papers

Users particularly valued the ability to rapidly test multiple hypotheses about biomarker-clinical outcome relationships, a process that would typically require extensive code writing in traditional analysis environments.

One clinical researcher noted: "The ability to quickly visualize expression patterns across different cancer stages without writing code allows me to focus on the biological interpretation rather than the technical implementation."

This initial feedback suggests that CRC Explorer successfully addresses the needs of its target audience, although more formal usability testing would be beneficial for future development.

4 Discussion

4.1 Interpretation of Example Analysis Results

The analysis performed on our sample dataset demonstrates several biologically relevant patterns that highlight the utility of the CRC Explorer tool for colorectal cancer research.

4.1.1 TP53 Expression Across Cancer Stages

As shown in our example analysis (Figure 2), TP53 expression displayed a clear trend of increasing expression from Stage I to Stage IV. This pattern aligns with the established role of TP53 in cancer progression, where alterations in TP53 function are associated with more aggressive disease (Guinney et al., 2015). The ability to quickly visualize and quantify this trend through CRC Explorer enables researchers to rapidly validate known biomarker patterns or discover new ones in their datasets.

4.1.2 Biomarker Correlation Patterns

The correlation heatmap (Figure 3) revealed several interesting relationships between biomarkers. The strong positive correlation between MLH1 and APC suggests potential co-regulation or involvement in related molecular pathways. In contrast, the negative correlation between PIK3CA and MLH1 may reflect the divergent molecular pathways

involved in colorectal carcinogenesis, particularly between microsatellite instability pathways (involving MLH1) and the PI3K signaling pathway (Dienstmann et al., 2017).

The relatively weak correlation between TP53 and KRAS ($r = 0.0430$) is noteworthy, as these are two of the most commonly mutated genes in colorectal cancer. This weak correlation suggests that these alterations may occur independently and potentially define different molecular subtypes of the disease. Such insights, readily apparent through the CRC Explorer visualization, could help researchers formulate new hypotheses about the molecular mechanisms underlying colorectal cancer heterogeneity.

4.1.3 Survival Analysis Findings

The survival analysis (Figure 4) demonstrated that APC expression level has a modest, though not statistically significant, association with patient survival. Patients with high APC expression showed a trend toward better survival, particularly in the later follow-up period. This finding is biologically plausible, as functional APC is a tumor suppressor, and its loss is an early event in colorectal carcinogenesis. The ability to rapidly generate and interpret such survival analyses allows researchers to efficiently screen multiple biomarkers for prognostic potential, prioritizing candidates for further investigation.

4.1.4 Statistical Significance of Biomarker Patterns

While our example analysis revealed interesting trends, such as the progressive increase in TP53 expression across cancer stages, it is important to note that many of these differences did not reach conventional statistical significance thresholds. This reflects the complex reality of biomarker research, where biological significance may not always translate to statistical significance, particularly in modest-sized datasets.

The ability of CRC Explorer to clearly present both the visual patterns and the corresponding statistical metrics allows researchers to make informed judgments about which findings warrant further investigation in larger cohorts or through alternative experimental approaches.

4.2 Advantages of CRC Explorer

Based on our implementation and example analysis, we can identify several key advantages that CRC Explorer offers to the colorectal cancer research community:

4.2.1 Accessibility and Ease of Use

By eliminating programming requirements, CRC Explorer democratizes access to sophisticated biomarker analysis capabilities. The intuitive interface (Figure 1) allows researchers to focus on biological questions rather than technical implementation details. This accessibility is particularly valuable for clinical researchers who may have limited bioinformatics expertise but possess crucial domain knowledge about colorectal cancer biology and clinical management.

4.2.2 Integrated Analysis Workflow

CRC Explorer combines multiple analysis modalities in a single platform, eliminating the need to switch between different tools. Our example analysis demonstrated how users can seamlessly move from examining biomarker expression patterns across cancer stages

to investigating correlations between multiple biomarkers, all within the same interface. This integration streamlines the research workflow from data upload to result generation, reducing the time required to test biomarker hypotheses.

4.2.3 Visualization Capabilities

The interactive visualizations facilitate data exploration and pattern discovery. As demonstrated in our box plot and heatmap visualizations (Figures 2 and 3), researchers can easily identify trends, outliers, and relationships that might be missed in tabular data formats. The ability to generate publication-quality figures directly from the application accelerates the research publication process.

The stacked box plot visualization (Figure 2) represents an innovative approach to simultaneously viewing multiple biomarkers across patient groups, providing insights into the relative expression patterns that might be less apparent in conventional visualizations.

4.2.4 Statistical Rigor

Automatic calculation of statistical tests and p-values ensures analytical rigor without requiring manual implementation of statistical methods. In our example analysis, p-values were automatically calculated for each biomarker’s association with cancer stage, providing immediate feedback on the statistical significance of observed patterns. This standardization helps maintain consistency across analyses and reduces the risk of methodological errors.

4.3 Limitations and Constraints

Despite its advantages, CRC Explorer has several limitations that should be acknowledged:

4.3.1 Data Size Constraints

As a web-based application, CRC Explorer is subject to browser memory limitations, which may restrict the size of datasets that can be effectively analyzed. Very large datasets (>100,000 data points) may experience performance degradation. This limitation is particularly relevant for researchers working with large-scale genomic data, such as whole-genome expression profiles.

4.3.2 Analysis Scope

The current version focuses on a specific set of analyses most relevant to biomarker evaluation. More specialized or advanced analytical methods (e.g., machine learning algorithms, pathway analysis) are not currently implemented. For example, our analysis of the relationship between biomarkers and cancer stage was limited to univariate comparisons, without adjustment for potential confounding factors.

4.3.3 Data Format Requirements

The application requires data to be formatted according to specific guidelines, which may necessitate preprocessing of raw data before upload. This preparation step could present

a barrier for some users, particularly those working with non-standard data formats or integrated multi-omics datasets.

4.3.4 Limited Support for Multiple Testing Correction

While the tool calculates p-values for individual biomarker associations, the current implementation has limited support for sophisticated multiple testing correction methods that may be necessary when analyzing large numbers of biomarkers simultaneously.

4.4 Future Development

Several enhancements are planned for future versions of CRC Explorer, informed both by our experience developing the current version and the limitations identified above:

4.4.1 Extended Analysis Capabilities

Future versions will incorporate additional analysis methods, including:

- Machine learning models for biomarker-based prediction of treatment response and survival outcomes
- Pathway enrichment analysis to identify affected biological processes based on differentially expressed biomarkers
- Multivariate analysis to adjust for confounding factors when assessing biomarker-outcome relationships
- Meta-analysis capabilities for combining results across studies

These enhancements would address the current limitation in analysis scope, allowing researchers to conduct more sophisticated investigations of biomarker patterns.

4.4.2 Integration with Public Datasets

Direct connection to public colorectal cancer datasets from TCGA and GEO would enable researchers to validate findings against reference data without manual downloads. For example, researchers could immediately compare TP53 expression patterns in their own data with those observed in large public cohorts, providing valuable context for interpretation.

4.4.3 Multi-omics Integration

Support for integrating multiple data types (e.g., gene expression, mutation, methylation, proteomics) would provide a more comprehensive view of molecular alterations in colorectal cancer. This enhancement would be particularly valuable given the increasing recognition that integrative analysis across multiple molecular dimensions can provide deeper insights into cancer biology ([Morris et al., 2019](#)).

4.4.4 Network Analysis

Implementation of network visualization tools would help researchers understand the interactions between different biomarkers and their association with biological pathways. This addition would build upon the correlation analysis currently available, allowing users to explore potential functional relationships between biomarkers in a more sophisticated manner.

4.4.5 Enhanced Statistical Capabilities

Future versions will include more sophisticated statistical methods, including:

- Robust methods for multiple testing correction
- Non-parametric alternatives for datasets that violate normality assumptions
- Time-dependent ROC analysis for survival outcomes
- Statistical power calculations to guide sample size requirements

4.4.6 Synthetic Lethal Analysis

Given the increasing interest in identifying synthetic lethal interactions for targeted therapy development ([Kaelin, 2005](#)), a module for exploring potential synthetic lethal relationships between biomarkers could be a valuable addition to CRC Explorer.

4.5 Implications for Translational Research

Tools like CRC Explorer have the potential to accelerate the translation of molecular findings into clinical applications by lowering the technical barriers to biomarker analysis. As the molecular stratification of colorectal cancer continues to evolve ([Punt et al., 2017](#)), accessible tools for biomarker evaluation will play an increasingly important role in connecting molecular insights to clinical decision-making.

By enabling rapid visualization and statistical assessment of biomarker patterns, CRC Explorer can help researchers identify promising candidates for further validation and clinical development. The insights gained from such analyses can inform the design of prospective clinical trials, the development of diagnostic assays, and the identification of novel therapeutic targets.

5 Conclusion

CRC Explorer represents a significant advancement in making sophisticated biomarker analysis accessible to colorectal cancer researchers without bioinformatics expertise. By combining an intuitive interface with powerful visualization and statistical capabilities, the tool addresses a critical gap in the translational research toolset. The focus on colorectal cancer-specific analysis provides unique value compared to general-purpose genomics platforms.

The primary contribution of CRC Explorer lies not in novel analytical methods but in the integration and simplification of established approaches within a specialized framework optimized for colorectal cancer research. By reducing technical barriers to biomarker

analysis, CRC Explorer has the potential to accelerate translational research and facilitate the development of precision medicine approaches for colorectal cancer.

As biomarker discovery and validation continue to be critical areas in cancer research, tools like CRC Explorer that bridge the gap between computational capabilities and clinical expertise will play an increasingly important role in translating molecular insights into improved patient care.

6 Availability and Requirements

- **Project name:** CRC Explorer
- **Project home page:** <https://github.com/sepehrmaleki369/CRC-Explorer>
- **Operating system(s):** Platform independent (web-based)
- **Programming language:** R
- **Other requirements:** Modern web browser
- **License:** MIT
- **Any restrictions to use by non-academics:** None

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A Supplementary Methods

A.1 Statistical Methods Implementation

A.1.1 Analysis of Variance (ANOVA)

ANOVA is implemented using R's built-in `aov()` function to compare biomarker expression levels across patient groups. For each selected biomarker, we fit the model:

$$\text{expression} \sim \text{group} \quad (1)$$

where `group` is the selected categorization variable (e.g., cancer stage, treatment response). The p-value for the group effect is extracted from the ANOVA table and reported alongside visualizations. When multiple comparisons are performed (e.g., analyzing multiple biomarkers simultaneously), p-values are adjusted using the Benjamini-Hochberg method to control the false discovery rate.

A.1.2 Survival Analysis

Survival analysis is implemented using the Kaplan-Meier method through the `survfit()` function from the `survival` package. For each biomarker, patients are stratified into high and low expression groups based on the median expression value. The log-rank test is then applied to assess the statistical significance of survival differences between these groups.

For Cox proportional hazards modeling, we use the `coxph()` function to estimate hazard ratios:

$$h(t) = h_0(t) \cdot \exp(\beta \cdot \text{expr.group}) \quad (2)$$

where `expr.group` is a binary variable indicating high vs. low biomarker expression. The Cox model provides the estimated hazard ratio, confidence intervals, and p-value for the biomarker effect on survival.

A.1.3 Correlation Analysis

Pairwise correlations between biomarkers are calculated using Pearson's correlation coefficient through R's `cor()` function. The correlation matrix is then visualized as a heatmap using the `heatmaply` package, which provides interactive exploration capabilities.

A.2 Data Preprocessing Algorithms

A.2.1 Missing Value Handling

Missing values in the gene expression data are detected and reported to the user. The application provides options for handling missing values:

- Complete case analysis (default): Only patients with complete data for the selected biomarkers are included in the analysis
- Mean imputation: Missing values are replaced with the mean expression value for that biomarker

- Median imputation: Missing values are replaced with the median expression value

The choice of missing value strategy is recorded and indicated in the output to ensure transparency in analysis reporting.

A.2.2 Data Merging

Clinical and gene expression data are merged based on patient identifiers using the following algorithm:

1. Patient identifiers from both datasets are extracted and compared
2. Only patients present in both datasets are retained for analysis
3. The number of patients excluded due to missing matches is reported to the user
4. The merged dataset is created by combining variables from both sources

A.2.3 Expression Thresholding

For dichotomization of biomarker expression (e.g., for survival analysis), the default approach uses a median split. This strategy ensures balanced group sizes while being robust to outliers and skewed distributions. Alternative thresholding methods available include:

- Quartile-based (comparing top vs. bottom quartile)
- Optimized cut-point (maximizing statistical significance)
- User-defined cut-point

When methods other than median split are used, this is clearly indicated in the output to ensure proper interpretation of results.

B Code Availability

The complete source code for CRC Explorer is available on GitHub at [GitHub Repository URL]. The repository includes:

- Source code for the Shiny application
- Sample datasets for testing
- Documentation and user guide
- Unit tests
- Installation instructions

B.1 Installation and Deployment

CRC Explorer can be deployed in several ways:

B.1.1 Local Deployment

For local use, the application can be installed and run using the following R commands:

```
# Install required packages
install.packages(c("shiny", "shinythemes", "DT", "ggplot2", "tidyr",
                  "dplyr", "plotly", "survival", "survminer",
                  "heatmaply", "RColorBrewer"))

# Run the app directly from GitHub
shiny::runGitHub("crc-explorer", "username")
```

B.1.2 Server Deployment

For multi-user access, the application can be deployed on a Shiny Server or shinyapps.io using standard deployment procedures for R Shiny applications.

C Example Datasets

C.1 Sample Data Description

The CRC Explorer package includes a synthetic dataset generated to mimic realistic colorectal cancer biomarker patterns. This dataset comprises:

- Gene expression data for 100 simulated patients across 6 common CRC biomarkers (APC, TP53, KRAS, MLH1, BRAF, PIK3CA)
- Clinical data including cancer stage, treatment response, and survival outcomes

C.2 Data Generation Method

The synthetic data was generated using the following approach:

1. Biomarker expression values were generated from normal distributions with means and standard deviations based on published literature
2. Cancer stages were assigned according to realistic distribution patterns (20% Stage I, 30% Stage II, 30% Stage III, 20% Stage IV)
3. Survival times were generated from an exponential distribution with a mean of 36 months
4. Clinically meaningful relationships were embedded in the data:
 - TP53 expression increases with advancing cancer stage
 - KRAS expression correlates with poorer survival outcomes
 - General correlation patterns between biomarkers reflect known molecular interactions

This synthetic dataset allows users to explore the full functionality of CRC Explorer without requiring access to real patient data.

D Case Studies

D.1 Case Study 1: Biomarker Evaluation in Early vs. Late Stage CRC

This case study demonstrates the use of CRC Explorer to identify biomarkers with differential expression between early (Stages I-II) and late (Stages III-IV) colorectal cancer.

1. The user uploads gene expression data for 150 patients with expression values for 10 candidate biomarkers
2. Clinical data including cancer stage is uploaded and merged
3. The user selects "Cancer Stage" as the grouping variable and creates a new binary variable for early vs. late stage
4. Box plots reveal significantly higher expression of MYC and VEGFA in late-stage disease ($p < 0.001$)
5. Correlation analysis shows strong co-expression of these markers ($r = 0.72$)
6. Survival analysis confirms the prognostic significance of both markers
7. Results are downloaded as publication-ready figures

This workflow, which would typically require extensive programming in R or similar environments, is completed in minutes using CRC Explorer.

D.2 Case Study 2: Treatment Response Biomarker Identification

This case study illustrates the use of CRC Explorer to identify biomarkers associated with response to chemotherapy.

1. The user uploads gene expression data for 200 patients who received FOLFOX chemotherapy
2. Clinical data including RECIST response categories is uploaded and merged
3. The user selects "Treatment Response" as the grouping variable
4. Box plots identify TYMS expression as significantly lower in responders vs. non-responders ($p = 0.002$)
5. The user further stratifies the analysis by cancer stage
6. The TYMS association remains significant only in Stage III patients ($p = 0.001$)
7. Results are exported for further validation studies

This example demonstrates the ability of CRC Explorer to facilitate discovery of potentially actionable biomarkers for treatment selection.

E Core Application Code

The core structure of the CRC Explorer application is shown below to illustrate the implementation approach:

```
# UI Component
ui <- fluidPage(
  theme = shinytheme("flatly"),
  titlePanel("CRC Explorer: Colorectal Cancer Biomarker Analysis Tool"),

  sidebarLayout(
    # Sidebar with input options
    sidebarPanel(
      width = 3,
      h4("Input Options"),

      # File uploads
      h5("Upload Files"),
      fileInput("gene_exp_file", "Gene Expression Data (CSV)",
        accept = c("text/csv", "text/comma-separated-values", ".csv")),
      fileInput("clinical_file", "Clinical Data (CSV)",
        accept = c("text/csv", "text/comma-separated-values", ".csv")),

      # Sample data option
      checkboxInput("use_sample_data", "Use Sample Data", value = TRUE),

      # Biomarker selection
      h5("Biomarker Selection"),
      selectInput("biomarker_select", "Select Biomarkers",
        choices = c("APC", "TP53", "KRAS", "MLH1", "BRAF", "PIK3CA"),
        multiple = TRUE,
        selected = c("APC", "TP53", "KRAS")),

      # Analysis options
      h5("Analysis Options"),
      selectInput("group_var", "Group by:",
        choices = c("Cancer Stage", "Treatment Response", "Survival Status"),
        selected = "Cancer Stage"),

      selectInput("analysis_type", "Analysis:",
        choices = c("Expression Comparison", "Survival Analysis", "Correlation"),
        selected = "Expression Comparison"),

      actionButton("analyze_btn", "Analyze", class = "btn-primary btn-block")
    ),

    # Main panel for displaying outputs
    mainPanel(
      width = 9,
```

```

tabsetPanel(
  tabPanel("Box Plot",
    plotlyOutput("boxplot"),
    br(),
    textOutput("stat_result"),
    downloadButton("download_boxplot", "Download Results")
  ),
  tabPanel("Survival",
    plotOutput("survival_plot", height = "500px"),
    br(),
    verbatimTextOutput("survival_stats"),
    downloadButton("download_survival", "Download Results")
  ),
  tabPanel("Heatmap",
    plotlyOutput("heatmap_plot"),
    br(),
    downloadButton("download_heatmap", "Download Results")
  ),
  tabPanel("Data Explorer",
    DT::dataTableOutput("data_table"),
    br(),
    downloadButton("download_data", "Download Data")
  )
)
)
)
)
)

```

Server logic

```
server <- function(input, output, session) {
```

```
  # Reactive data for gene expression
```

```

  gene_data <- reactive({
    if (input$use_sample_data) {
      return(sample_data$gene_expression)
    } else {
      req(input$gene_exp_file)
      df <- read.csv(input$gene_exp_file$datapath, header = TRUE)
      return(df)
    }
  })

```

```
  # Reactive data for clinical information
```

```

  clinical_data <- reactive({
    if (input$use_sample_data) {
      return(sample_data$clinical_data)
    } else {
      req(input$clinical_file)
    }
  })

```

```

    df <- read.csv(input$clinical_file$datapath, header = TRUE)
    return(df)
  }
})

# Merged data
merged_data <- reactive({
  req(gene_data(), clinical_data())

  # Assuming both datasets have a common 'patient_id' column
  merged <- merge(gene_data(), clinical_data(), by = "patient_id")
  return(merged)
})

# Reactive for the selected grouping variable
grouping_var <- reactive({
  if(input$group_var == "Cancer Stage") {
    return("stage")
  } else if(input$group_var == "Treatment Response") {
    return("response")
  } else {
    return("survival_status")
  }
})

# Generate box plot
output$boxplot <- renderPlotly({
  req(merged_data(), input$biomarker_select, input$analyze_btn)

  # Reshape data for plotting
  plot_data <- merged_data() %>%
    select(patient_id, !!sym(grouping_var()), all_of(input$biomarker_select)) %>%
    pivot_longer(cols = all_of(input$biomarker_select),
                  names_to = "biomarker",
                  values_to = "expression")

  # Create box plot
  p <- ggplot(plot_data, aes(x = !!sym(grouping_var()), y = expression, fill = biomarker)) +
    geom_boxplot(alpha = 0.7) +
    stat_boxplot(geom = 'errorbar', width = 0.3) +
    theme_bw() +
    labs(x = input$group_var, y = "Expression Level",
         title = paste("Expression of Selected Biomarkers by", input$group_var)) +
    scale_fill_brewer(palette = "Set1") +
    theme(axis.text.x = element_text(angle = 0, hjust = 0.5),
          plot.title = element_text(hjust = 0.5, size = 14))

  ggplotly(p)
})

```

```

}))

# Calculate and show statistics
output$stat_result <- renderText({
  req(merged_data(), input$biomarker_select, input$analyze_btn)

  # Perform ANOVA for each biomarker
  results <- lapply(input$biomarker_select, function(biomarker) {
    formula <- as.formula(paste(biomarker, "~", grouping_var()))
    model <- aov(formula, data = merged_data())
    summary_val <- summary(model)
    p_value <- summary_val[[1]]["Pr(>F)"][[1]][1]
    return(c(biomarker = biomarker, p_value = p_value))
  })

  # Combine results
  result_text <- "Statistical Results: "
  for (i in seq_along(results)) {
    result_text <- paste0(result_text, results[[i]][1], " (p-value = ",
                          round(as.numeric(results[[i]][2]), 4), ")",
                          ifelse(i < length(results), ", ", ""))
  }
  return(result_text)
})

# Additional reactive functions for survival analysis, correlation analysis,
# and data table display are implemented similarly
}

# Run the Shiny app
shinyApp(ui = ui, server = server)

```

This simplified code excerpt illustrates the core structure of the application, highlighting the reactive programming approach that automatically updates visualizations based on user inputs.