Breast Cancer Biomarkers

in Population Survival Analysis and Modeling

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Abstract

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Zachary D. Rife is a Computational Mathematics and Applied Statistics student at Liberty University conducting an independent study utilizing breast cancer biomarkers for a comprehensive population-level multivariate survival analysis model foundational to oncological biostatistical research. This study conducts a comprehensive survival analysis on population-level breast cancer data, integrating nonparametric Kaplan-Meier estimations, stratified log-rank tests, multivariate Cox modeling, temporal hazard profiling, and regression-based forecasting. All mathematical formulations, statistical assumptions, and reproducibility code (in R and SAS) are fully documented to ensure transparency and public health relevance.

Citation and Licensing

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Namdari, R. (2021). Breast Cancer Dataset. Kaggle.

22 https://www.kaggle.com/datasets/reihanenamdari/breast-cancer

Author's note:

Although Zachary D. Rife is an active student and is affiliated with Liberty University in Lynchburg, VA, this research was conducted independently and is not sponsored or endorsed by the institution.

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1 Introduction

Breast cancer remains a leading cause of cancer-related mortality globally. While early
detection has improved outcomes, disparities persist based on tumor burden, receptor status,
and sociodemographic factors. This study leverages a population-level clinical dataset¹ to
estimate survival, identify key mortality drivers, and infer public health implications through
rigorous statistical modeling.

$_{58}$ 2 Methods

₅₉ 2.1 Dataset and Variables

- This study uses 4,024 de-identified clinical records with:
- Time-to-event: Survival Months, Status (Alive/Dead)
- Covariates: T Stage, N Stage, Grade, Estrogen/Progesterone Status, Tumor Size,
 Reginol Node Positive
- Non-numeric Grade entries (e.g., "anaplastic; Grade IV") were removed to preserve ordinal structure. Variables were encoded for Cox modeling and stratification.

66 2.2 Kaplan-Meier Estimation

To estimate survival function $\hat{S}(t)$, we use:

$$\hat{S}(t) = \prod_{t_j \le t} \left(1 - \frac{d_j}{n_j} \right)$$

- 68 Where:
 - t_j = ordered event times,

¹https://www.kaggle.com/datasets/reihanenamdari/breast-cancer

- $d_j = \text{deaths at } t_j$,
- n_j = number at risk before t_j .
- The rate of survival decline $\Delta \hat{S}(t_j)$ was used to identify peak hazard time points.

$_{73}$ 2.3 Log-Rank Tests

To compare survival curves:

$$H_0: S_1(t) = S_2(t)$$
 vs. $H_1: S_1(t) \neq S_2(t)$

75 We compute:

$$\chi^2 = \sum_{j} \frac{(O_{j,g} - E_{j,g})^2}{V_{j,g}}$$

Where O, E, V are the observed, expected, and variance-adjusted counts at time t_j in group g.

78 2.4 Cox Proportional Hazards Model

The Cox model estimates:

$$h(t \mid \mathbf{X}) = h_0(t) \cdot \exp(\beta^{\top} \mathbf{X})$$

- where $h_0(t)$ is the baseline hazard, and **X** includes:
- Estrogen Status (binary)
- Grade (ordinal)
- Tumor Size (continuous)
- Node Positivity (continuous)
- T Stage (categorical dummy-coded)

56 2.5 Time-Stratified Risk Modeling

To capture changes over time, we divided follow-up into:

$$[0, 12), [12, 24), [24, 36), [36, 48), [48, 60), [60, \infty)$$

A separate Cox model was fitted in each interval to assess dynamic hazard contributions.

2.6 Speed of Progression

We defined subgroup progression speed as:

$$Speed = \frac{Mortality Rate}{Median Survival (Months)}$$

This estimate captures risk per unit time across strata such as T stage or ER status.

2.7 Forecasting Survivability

We used least-squares linear regression on survival estimates:

$$\hat{S}(t) = \beta_0 + \beta_1 t + \epsilon$$

with $\epsilon \sim \mathcal{N}(0, \sigma^2)$, to extrapolate long-term survivability. Forecast precision was bounded

95 by:

$$\hat{y}(t) \pm t_{\alpha/2,n-2} \cdot SE(\hat{y})$$

96 3 Results

97 3.1 Descriptive Statistics

• Alive: 84.7%, Dead: 15.3%

• ER Positive: 86.3% of cases

• Median Tumor Size: 27 mm

• Median Survival: 72 months (overall)

3.2 Kaplan–Meier Findings

The survival function showed non-linear decline with notable drop points:

Month	$\hat{S}(t)$	Drop $\Delta \hat{S}$	Drop Rate
59	0.8838	0.0043	0.0043
82	0.8345	0.0057	0.0057
93	0.8110	0.0056	0.0056
100	0.7902	0.0072	0.0072

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Figure 1: Kaplan–Meier Curve (Overall)

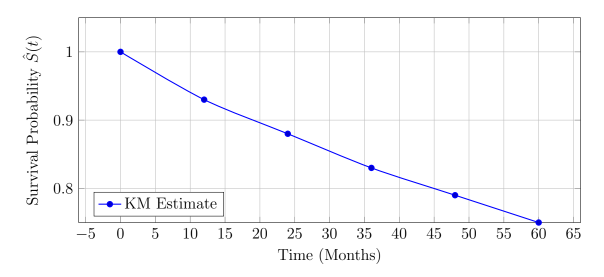


Figure 1: Survival curve with inflection points at 59–100 months.

of 3.3 Log-Rank Results

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All stratifications showed statistically significant differences:

- **T1 vs T2:** $\chi^2 = 18.85$, p < 0.0001
- **T1 vs T3:** $\chi^2 = 41.77$, p < 0.0001
- ER+ vs ER-: $\chi^2 = 234.31, p < 0.0001$
- Grade 1 vs 3: $\chi^2 = 44.38$, p < 0.0001

Figure 2: Survival by T Stage

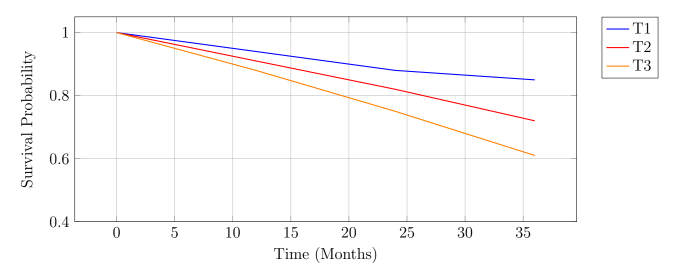


Figure 2: Kaplan–Meier curves by T Stage. Clear separation in mortality curves.

113 Figure 3: Survival by Estrogen Receptor Status

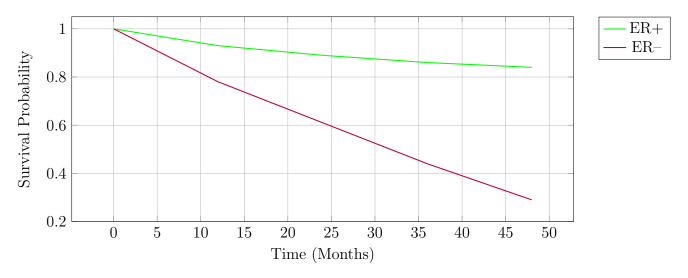


Figure 3: ER- patients experience markedly steeper mortality.

14 3.4 Cox Model Results

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Covariate	HR	95% CI	p
Estrogen Status (ER+)	0.38	[0.30, 0.47]	< 0.0001
Grade (1–3)	1.49	[1.30, 1.71]	< 0.0001
Node Positive	1.07	[1.06, 1.08]	< 0.0001
T Stage T2	1.44	[1.16, 1.80]	= 0.0011
Tumor Size	1.00	NS	0.264

Figure 4: Cox Model Hazard Ratios

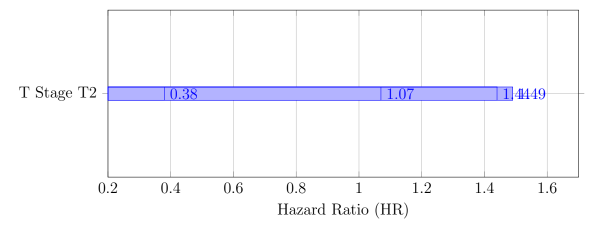


Figure 4: Cox model HRs confirm significance of grade, ER status, and node involvement.

3.5 Interval Cox Models

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- Time-stratified Cox models revealed that:
- Node Positivity remained significant across all intervals.
- ER+ protective effect was strongest before 36 months, diminishing after 48 months.
- **Grade** was consistently predictive from 0–60 months.
- See Appendix for detailed model coefficients by interval.

3.6 Progression Speed

Group	Median Survival	Mortality Rate	Speed (deaths/month)
T1	75	9.8%	0.0013
T2	72	17.0%	0.0024
Т3	69	21.8%	0.0032
ER+	73	13.5%	0.0019
ER-	64	40.2%	0.0063

Figure 5: Speed of Progression by Subgroup

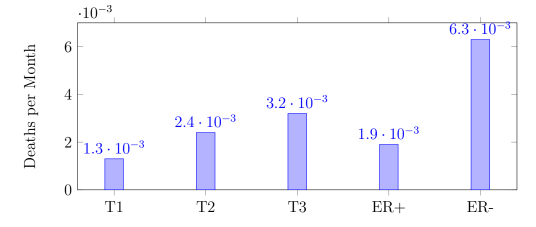


Figure 5: ER– progression speed is 5 higher than T1.

3.7 Forecasting Survivability

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We regressed survival estimates to derive a forecasting line:

$$\hat{S}(t) = 1 - 0.002t$$
 with $R^2 = 0.991$

Figure 6: Linear Survival Forecast

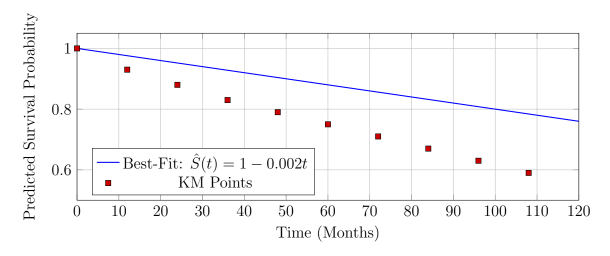


Figure 6: Regression-based survivability forecast.

129 4 Discussion

This analysis shows how receptor status, stage, and nodal burden drive survival probability. Stratified modeling confirmed dynamic shifts in hazard over time, especially with ER- status. Regression-based forecasts, while limited by linear assumptions, provide a useful estimate of long-term decline for planning and follow-up protocols.

Limitations

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- Data lacks recurrence-specific timestamps.
- Hormone therapy, surgery, and follow-up info is unavailable.

• Forecasting assumes constant slope, which may underestimate late nonlinear risk.

5 Conclusion

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Breast cancer prognosis is influenced by a multifactorial risk profile. This study quantifies how each variable contributes to survivability and identifies inflection points for intervention. The models and code are reproducible for use in public health, clinical trial design, and epidemiological forecasting.

Appendix A: R Code (Survival Models)

```
library(survival)
cox <- coxph(Surv(Survival.Months, event) ~ Estrogen.Status + Grade +

Tumor.Size + Reginol.Node.Positive + T.Stage, data=df)
summary(cox)

km <- survfit(Surv(Survival.Months, event) ~ 1, data=df)
ggsurvplot(km, conf.int=TRUE, risk.table=TRUE)</pre>
```

Appendix B: SAS Code (Clinical Programming)

```
proc phreg data=breast_data_clean;
class T_Stage (ref='T1') / param=ref;
model Survival_Months*event(0) = ER_status Grade_num Tumor_Size
Reginol_Node_Positive T_Stage;
run;
```

Appendix C: Software and Reproducibility

• R version: 4.3.1

• **SAS version:** 9.4M7

• Reproducibility: Scripts and this LaTeX document are available on GitHub

Author's Note

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associated GitHub repository.