# Variance component mixture modelling for longitudinal T-cell receptor clonal dynamics

David Swanson<sup>1</sup>, Alex Sherry<sup>2</sup>, and Chad Tang<sup>2</sup>

<sup>1</sup>Department of Biostatistics, <sup>2</sup>Department of Radiation Oncology, University of Texas MD Anderson Cancer Center

October 18, 2024

#### Abstract

Analyses of T-cell receptors has shown promise in elucidating the association between immune response and human disease, from cancer and infectious disease to autoimmune conditions. Methods to identify expanding and contracting T-cell receptor clones have so far been limited to longitudinal pairwise comparisons of changes in clone frequency, then adjusted for multiplicity to control the false discovery rate. Mixture models show promise in playing a useful role in distinguishing dynamic T-cell receptor clones from static ones, allowing one to relate these features to exposures and different markers of health and prognosis. While it is common to mix on the location or scale parameter of a family of distributions, mixing on the parameterization of the model is exploited in our approach, with one mixture component allowing TCR clones to have variable mean over longitudinal followup, whereas the other component does not. Our model leverages Bayesian hierarchy, utilizing the posterior predictive distribution of Gamma-Poisson conjugacy to arrive at negative binomial and negative multinomial distributions of components, each modified according to an offset term for receptor read count normalization. An EM-algorithm is developed to estimate hyperparameters, and validity of the approach is demonstrated in simulation. The model identifies a statistically significant and clinically relevant increase in TCR clonal dynamism among metastasis-directed radiation therapy in a cohort of prostate cancer patients.

# 1 Introduction

Immune response plays a central role in human health and addressing the many foreign agents people are regularly exposed to. T-cells are an important piece of one's adaptive immune system, in part by recognizing antigens with receptors that exhibit an exceptional degree of diversity via highly variable gene recombination in portions of the receptor [15, 8]. Study of T-cell receptor (TCR) repertoires has gained attention in recent years because immunose-quencing receptors allows one to examine TCR clonal dynamics and their association with biomarkers, disease, and other aspects of human biology [23, 19, 16]. Clones of receptors binding an antigen tend to expand, which can be measured by calculating the proportion of a certain clone within the repertoire and examining the quantity over time [7]. Other clones contract or are relatively static proportionally within the repertoire, and these dynamics have been looked at longitudinally [11]. Identifying expanding and contracting clones has taken different forms in the past, though a popular approach places the problem within a pairwise time comparison and multiple testing framework, identifying significant changes in clonal proportions over two points in time within a Beta-Binomial conjugate prior setting which effectively

moderates estimated rate changes by pulling clones with low read counts toward the null [20]. The procedure then applies false discovery proportion control to address the severe multiple testing burden involved in the analysis [3]. The approach has been successful at identifying expanding and contracting clones and relating the dynamics to a variety of diseases. Cancer has been a disease of primary interest with respect to longitudinal TCR clonal movement and trajectories and their associations with treatment and prognosis, with recent studies demonstrating clonal expansions and contractions significantly associated with radiation therapy directed at metastases [22].

Comprehensive statistical modelling approaches to understanding longitudinal T-cell receptor clonal dynamics have so far not been pursued in the literature to our knowledge, which might account for variable follow-up by observation, missingness, and error term specification so that what constitutes significant clonal frequency change is explicit. Several challenges arise when trying to develop a model for these dynamics. Because of heterogeneity in the total number of template reads within the biological unit being interrogated, one models a fraction, rather than an absolute count, since it is the clonal fraction relative to others that elucidates biological fluctuation and therefore interest. Second, because of extreme antigenic and therefore receptor diversity, most clones are not "public", or shared, across the population being modelled, unlike expression of a certain gene or many gene mutations which though varying across observations have a common interpretation in one's sample [2]. Therefore one generally cannot relate any one clone and its expansion or contraction behavior with patient phenotype, rather, one must consider dynamics of the repertoire as a whole. Lastly, while it is often feasible to identify change in some proportion from a statistical perspective, identifying biologically relevant changes, which presumably are of greater interest for prognostic models and other uses, is a different and more nuanced question. Inclusion of an error term that is monotonic increasing in the clone's mean parameter may capture biological mechanism best.

We try to address these challenges with an interesting and novel variance component mixture model, which leverages Bayesian conjugacy under two different parameterizations of a variance component model, one component corresponding to dynamic (expanding or contracting) clones, and one component corresponding to static clones [17, 21]. The model posits clonal mean frequencies being sampled from a Gamma prior which we use hierarchically to sample from a Poisson on that clone's mean parameter  $\lambda$ , with offset term corresponding to the total number of reads within the biological unit being modelled. Under the dynamic component, a different clonal mean parameter  $\lambda$  is hypothesized for each longitudinal followup, giving flexibility to the within-clone frequency variability over time. In contrast, the same  $\lambda$  parameterizes all follow-up times under the static component, constraining the entirety of that clone's frequencies for the times observed. For both parameterizations, we marginalize out  $\lambda$ , leveraging conjugacy to arrive at a marginal likelihood that is a product of negative binomial probability mass functions (pmf) for the dynamic component clones and the negative multinomial pmf for the static component clones, both of which are modified due to the offset term. We fit  $\alpha$  and  $\beta$  with empirical Bayes, and use the expectation-maximization (EM) algorithm to calculate the probability of component membership for each clone, be it dynamic or static, iterating until convergence [10].

In section 2, we derive the model and means of its fitting, grounding it in hierarchical Bayesian modelling and the broad literature on mixture models. In section 3, we examine sensitivity of the model to different sets of parameters, proportion mixing, and powerings and compare with the multiple testing framework in which such analyses are often done. In section 4 we analyze a cohort of prostate cancer patients who are randomized to metastasis-directed therapy or not and the influence of that exposure on TCR clonal dynamics. We conclude in Section 5 with consideration of model extensions and improvement.

# 2 Methods

Finite mixture models have played a central role in the development of statistical modelling with broad use across data-generating mechanisms where there is belief that subsets of observations arise under a discrete set of different parameters [13, 6, 9]. Since parameters are generally unknown for each subset, or component, nor is component membership known among observations, these models can be difficult to fit and considerable work has been devoted to understanding how and behavior of subsequent estimates. Simulation-based methods like Markov chain monte carlo (MCMC) and other optimization routines have both proved valuable tools in model fitting [robertMCMC, 18]. It may be most common to see these mixture models mixing on location and scale parameters in exponential families of distributions for clustering data, though more sophisticated mixtures of regressions or mixtures of experts have also been proposed and applied to different modelling settings [5, 14].

It is less common to mix on the parameterization of a model, which is our proposition here. Our mixture consists of two different Gamma-Poisson hierarchies which upon marginalization yields a negative multinomial distribution governed by two hyperparameters  $\alpha$  and  $\beta$ , whereas the other component is a product of negative binomial distributions governed by the same parameters, where the number of terms is the follow-up times within clonotype. Both of these variance components are a function of the number of clonotype TCR reads and the total number of reads across clones per person-time. While variance components are typically used in mixed models to account for within-observation correlation and augment fixed effects, here they are useful to partition dynamic clones from static ones, that is those that exhibit significant heterogeneity in clonal proportion across time versus those that do not [21, 24, 1].

We fit the model by first randomly assigning  $E[D_{ij}]$ , optimizing with respect to  $\alpha$  and  $\beta$ , then iterating until convergence, defined as the sum of changes in component probabilities being less than some  $\epsilon$ .

#### 2.1 Model framework

Consider the  $i^{th}$  clone template count for person j at time k,  $C_{ijk}$ , and the total number of template counts for person j at time k,  $O_{jk} = \sum_{i=1}^{T_j} C_{ijk}$ , where  $T_j$  is the number of follow up times of observations j. Since there is significant variability in the total number of template reads and it is of primary biological interest how proportions of clones change over time, we model the proportion  $C_{ijk}/O_{jk}$  over time and patient.

We seek to distinguish the "dynamic" clones (denoted  $D_{ij} = 1$  for clone i person j), who exhibit significant contraction and/or expansion over longitudinal follow-up, from the static ones ( $D_{ij} = 0$ ). The hierarchical Bayesian formulation envisions that among the static component, clonal mean parameter  $\lambda_{ij} \sim \text{Gamma}(\alpha, \beta)$  is invariant over time k and sampled from a Gamma distribution on hyperparameters  $\alpha$  and  $\beta$ . Template count reads are assumed Poisson-distributed  $C_{ijk} \sim \text{Pois}(\lambda_{ij}O_{jk})$ , where the  $\lambda_{ij}$  is scaled by the total number of reads at a person-time  $O_{jk}$ , effectively treating its logged value as an offset. In contrast, under the dynamic component, separate  $\lambda_{ijk}$  are assumed sampled for each time point, in which case subsequently  $C_{ijk} \sim \text{Pois}(\lambda_{ijk}O_{jk})$ . Integration of  $\lambda_{ij}$  and  $\lambda_{ijk}$  yields the respective posterior predictive distributions, which due to conjugacy are here the negative multinomial and negative binomial distributions accordingly. These distributions are appropriate for modelling clonal template counts because of the right-skewed and significant dynamic ranges of this outcome. These distributions are modified according to the  $O_{jk}$  via their use as an offset, implicitly modelling clonal proportion dynamics.

#### 2.2 Static model

We have the static component clonal model as

$$C_{ijk} \mid (\lambda_{ij}, O_{jk}, D_{ij} = 0) \sim \text{Pois}(\lambda_{ij} O_{jk})$$

with

$$\lambda_{ij}|(D_{ij}=0)\sim \mathrm{Gamma}(\alpha,\beta)$$

The likelihood for  $C_{ij} = (C_{ij1} \dots C_{ijT_j})$  given  $\lambda_{ij}$  and  $O_j = (C_{j1} \dots C_{jT_j})$ ,  $l(C_{ij} | \lambda_{ij}, O_j) = \prod_k l(C_{ijk} | \lambda_{ij}, O_{jk})$ , with  $T_j$  the number of time points observed for clone i on person j due to an unbalanced design (ie, missingness of follow up for a subset of patients), is

$$l(C_{ij}, | \lambda_{ij}, O_{j}, O_{j}) = \prod_{k=1}^{T_j} \frac{\exp(-\lambda_{ij}O_{jk}) (\lambda_{ij}O_{jk})^{C_{ijk}}}{C_{ijk}!} = \frac{\exp(-\lambda_{ij}\sum_k O_{jk}) \lambda_{ij}^{\sum_k C_{ijk}} \prod_{k=1}^{T_j} O_{jk}^{C_{ijk}}}{\prod_{k=1}^{T_j} C_{ijk}!}$$

Since our prior on  $\lambda_{ij}$ ,  $p(\lambda_{ij}|D_{ij}=0,\alpha,\beta)$ , is Gamma $(\alpha,\beta)$ , the conditional joint distribution of  $(C_{ij},\lambda_{ij})|O_{j},D_{ij}=0$  is

$$p(C_{ij}, \lambda_{ij} | O_j, D_{ij} = 0, \alpha, \beta) = l(C_{ij}, |\lambda_{ij}, O_j, p(\lambda_{ij} | D_{ij} = 0, \alpha, \beta))$$

$$= \frac{\exp(-\lambda_{ij} \sum_k O_{jk}) \lambda_{ij}^{\sum_k C_{ijk}} \prod_{k=1}^{T_j} O_{jk}^{C_{ijk}}}{\prod_{k=1}^{T_j} C_{ijk}!} \cdot \frac{\beta^{\alpha}}{\Gamma(\alpha)} \lambda_{ij}^{\alpha-1} \exp(-\beta \lambda_{ij})$$
(1)

Collecting terms of equation (1) and leveraging Gamma-Poisson conjugacy, we recognize the kernel of the Gamma on updated parameters  $\alpha' = \sum_k C_{ijk} + \alpha$  and  $\beta' = \sum_k O_{jk} + \beta$ . Then, integrating out  $\lambda_{ij}$ , we obtain the inverse normalizing constant of the Gamma( $\alpha'$ ,  $\beta'$ )

$$p(C_{ij} \cdot | O_j \cdot, D_{ij} = 0, \alpha, \beta) = \int p(C_{ij} \cdot, \lambda_{ij} | O_j \cdot, D_{ij} = 0, \alpha, \beta) \, d\lambda_{ij}$$

$$= \frac{\beta^{\alpha} \prod_{k=1}^{T_j} O_{jk}^{C_{ijk}}}{\Gamma(\alpha) \prod_{k=1}^{T_j} C_{ijk}!} \int_0^{\infty} \lambda_{ij}^{\sum_k C_{ijk} + \alpha - 1} \exp\left(-\lambda_{ij} (\beta + \sum_k O_{jk})\right) d\lambda_{ij}$$

$$= \frac{\beta^{\alpha} \prod_{k=1}^{T_j} O_{jk}^{C_{ijk}}}{\Gamma(\alpha) \prod_{k=1}^{T_j} C_{ijk}!} \cdot \frac{\Gamma(\sum_k C_{ijk} + \alpha)}{(\beta + \sum_k O_{jk})^{\sum_k C_{ijk} + \alpha}}$$

$$= \frac{\Gamma(\sum_k C_{ijk} + \alpha)}{\Gamma(\alpha) \prod_{k=1}^{T_j} C_{ijk}!} \left(\frac{\beta}{\beta + \sum_k O_{jk}}\right)^{\alpha} \prod_{k=1}^{T_j} \left(\frac{O_{jk}}{\beta + \sum_k O_{jk}}\right)^{C_{ijk}}$$
(2)

which for discrete  $\alpha$  we recognize Equation (2) as a negative multinomial probability mass function on size parameter  $\alpha$ , success probability  $\beta/(\sum_k O_{jk} + \beta)$ , and "offset-normalized" near equal-valued failure probabilities  $O_{jk}/(\beta + \sum_k O_{jk})$ ,  $k = 1, \ldots, T_j$ , and evaluated at vector  $(C_{ij1}, \ldots, C_{ijT_j}, \alpha - 1)$ , for clone read counts  $C_{ij}$ , noting  $\Gamma(n) = (n-1)!$  for integers n.

#### 2.3 Dynamic Model

For the dynamic model component we parameterize clonal mean  $\lambda_{ijk}$  to vary across time k. While  $\lambda_{ij}$  in the static model can be considered a random effect and has an expectation which is identifiable,  $\lambda_{ijk}$  is non-identifiable and and comparison of the two components is only possible under marginalization and the induced posterior predictive distributions. Integration of  $\lambda_{ijk}$  yields a product of posterior predictive distributions which are negative binomial distributed

$$p(C_{ij} \mid O_{j}, D_{ij} = 1, \alpha, \beta) = \prod_{k=1}^{T_{j}} \int p(C_{ijk}, \lambda_{ijk} \mid O_{j}, D_{ij} = 1, \alpha, \beta) \ d\lambda_{ijk}$$

$$= \prod_{k=1}^{T_{j}} \int l(C_{ijk} \mid \lambda_{ijk}, O_{j}) p(\lambda_{ijk} \mid D_{ij} = 1, \alpha, \beta) \ d\lambda_{ijk}$$

$$= \prod_{k=1}^{T_{j}} \int \frac{\exp(-\lambda_{ijk} O_{jk}) \lambda_{ijk}^{C_{ijk}} O_{jk}^{C_{ijk}}}{C_{ijk}!} \cdot \frac{\beta^{\alpha}}{\Gamma(\alpha)} \lambda_{ijk}^{\alpha-1} \exp(-\beta \lambda_{ijk}) \ d\lambda_{ijk}$$

$$= \prod_{k=1}^{T_{j}} \frac{\Gamma(C_{ijk} + \alpha)}{\Gamma(\alpha) C_{ijk}!} \left(\frac{\beta}{\beta + O_{jk}}\right)^{\alpha} \left(\frac{O_{jk}}{\beta + O_{jk}}\right)^{C_{ijk}}$$
(3)

where we write the distribution in terms of the  $\Gamma(\cdot)$  function, and which in  $C_{ijk}$  we recognize as the product of  $T_j$  negative binomial mass functions on size parameter  $\alpha$  and success probabilities  $\beta/(\beta + O_{jk})$  for  $k = 1, ..., T_j$ . We develop our mixture model using these mass functions in equations (2) and (3) for the static and dynamic components, respectively.

## 2.4 Mixing and Model fitting

Having derived the component expressions we can write the likelihood in terms of them and the mixing distribution described with binary  $D_{ij}$  and  $P(D_{ij} = 1|\pi) = \pi$ . We write

$$\begin{split} p(C_{ij}. \mid O_{j}.\alpha, \beta, \pi) &= p(C_{ij}., D_{ij} = 1 \mid O_{j}., \alpha, \beta, \pi) + p(C_{ij}., D_{ij} = 0 \mid O_{j}., \alpha, \beta, \pi) \\ &= p(C_{ij}. \mid O_{j}., D_{ij} = 1, \alpha, \beta) P(D_{ij} = 1 \mid \pi) + p(C_{ij}. \mid O_{j}., D_{ij} = 0, \alpha, \beta) P(D_{ij} = 0 \mid \pi) \\ &= \pi \cdot p(C_{ij}. \mid O_{j}., D_{ij} = 1, \alpha, \beta) + (1 - \pi) \cdot p(C_{ij}. \mid O_{j}., D_{ij} = 0, \alpha, \beta) \end{split}$$

since  $P(D_{ij})$  is not a function of  $\alpha$  nor  $\beta$ . Hyperparameters  $\alpha$  and  $\beta$  are estimated using empirical Bayes and the model is fit using the EM algorithm [10]. We calculate the expectation of the complete data log-likelihood

$$\sum_{ijk} \log p(C_{ijk}|O_{jk}, \alpha, \beta, \pi) = \sum_{ijk} \log \left( p(C_{ijk}, D_{ij} = 1|O_{jk}, \alpha, \beta, \pi) + p(C_{ijk}, D_{ij} = 0|O_{jk}, \alpha, \beta, \pi) \right)$$

$$= \sum_{ijk} \log \left( \pi \cdot p(C_{ijk}|O_{jk}, D_{ij} = 1, \alpha, \beta) + (1 - \pi) \cdot p(C_{ijk}|O_{jk}, D_{ij} = 0, \alpha, \beta) \right)$$

for the  $(m+1)^{st}$  iteration under  $p(D_{ij}|C_{\cdot j}, O_{j}, \alpha^{(m)}, \beta^{(m)}, \pi^{(m)})$  given current estimates of  $(\alpha^{(m)}, \beta^{(m)}, \pi^{(m)})$ , then maximize the expression with respect to  $\alpha, \beta, \pi$  to find  $\alpha^{(m+1)}, \beta^{(m+1)}$ , and  $\pi^{(m+1)}$ . That is, we find

$$Q(\alpha, \beta, \pi | \alpha^{(m)}, \beta^{(m)}, \pi^{(m)}) = E_{p_D}[\log p(C_{ijk}, D_{ij} | O_{jk}, \alpha, \beta, \pi)]$$

where for brevity of notation we use  $p_D = p(D_{ij}|C_{ijk}, O_{jk}, \alpha^{(m)}, \beta^{(m)}, \pi^{(m)})$ , and subsequently calculate

$$(\hat{\alpha},\; \hat{\beta},\; \hat{\pi}) = \arg\max_{\alpha,\beta,\pi} Q(\alpha,\beta,\pi | \alpha^{(m)},\beta^{(m)},\pi^{(m)})$$

Since  $\alpha$  and  $\beta$  are present in both components, maximization is difficult. While an additional, nested EM is possible to estimate  $(\alpha^{(m+1)}, \beta^{(m+1)})$ , here we use BFGS for the maximization step [4, 12].

Because our mixture consists of two components so that  $D_{ij}$  can be considered Bernoulli for all i, j, calculation of  $p(D_{ij}|C_{ijk}, O_{jk}, \alpha^{(m)}, \beta^{(m)}, \pi^{(m)})$  is reduced to finding its expectation

which is

$$E[D_{ij}|C_{ij\cdot},O_{j\cdot},\alpha^{(m)},\beta^{(m)},\pi^{(m)}] = p(D_{ij} = 1|C_{ij\cdot},O_{j\cdot},\alpha^{(m)},\beta^{(m)},\pi^{(m)})$$

$$= \frac{p(D_{ij} = 1|C_{ij\cdot},O_{j\cdot},\alpha^{(m)},\beta^{(m)},\pi^{(m)})}{p(D_{ij} = 1|C_{ij\cdot},O_{j\cdot},\alpha^{(m)},\beta^{(m)},\pi^{(m)}) + p(D_{ij} = 0|C_{ij\cdot},O_{j\cdot},\alpha^{(m)},\beta^{(m)},\pi^{(m)})}$$

$$= \frac{p(D_{ij} = 1,C_{ij\cdot}|O_{j\cdot},\alpha^{(m)},\beta^{(m)},\pi^{(m)})}{p(D_{ij} = 1,C_{ij\cdot}|O_{j\cdot},\alpha^{(m)},\beta^{(m)},\pi^{(m)}) + p(D_{ij} = 0,C_{ij\cdot}|O_{j\cdot},\alpha^{(m)},\beta^{(m)},\pi^{(m)})}$$

$$= \frac{\pi^{(m)} p(C_{ij\cdot}|O_{j\cdot},D_{ij} = 1,\alpha^{(m)},\beta^{(m)})}{\pi^{(m)} p(C_{ij\cdot}|O_{j\cdot},D_{ij} = 1,\alpha^{(m)},\beta^{(m)}) + (1-\pi^{(m)}) p(C_{ij\cdot}|O_{j\cdot},D_{ij} = 0,\alpha^{(m)},\beta^{(m)})}$$
(5)

where equation (4) uses  $p(D_{ij}|C_{ij}, O_j, \alpha^{(m)}, \beta^{(m)}, \pi^{(m)}) \propto \pi^{(m)D_{ij}}(1-\pi^{(m)})^{1-D_{ij}} p(C_{ij}, |O_j, D_{ij}, \alpha^{(m)}, \beta^{(m)})$ , and also  $p(D_{ij} = 1|C_{ij}, O_j, \alpha, \beta, \pi) = 1 - p(D_{ij} = 0|C_{ij}, O_j, \alpha, \beta, \pi)$ , and (5) is calculated with component expressions (2) and (3). We iterate EM steps until the convergence criterion is reached, the mean of changes  $\sum_{ij} (D_{ij}^{(m)} - D_{ij}^{(m+1)})/n$  for iteration m to m+1 less than some  $\epsilon$ , chosen here  $10^{-8}$ .

# 3 Simulation

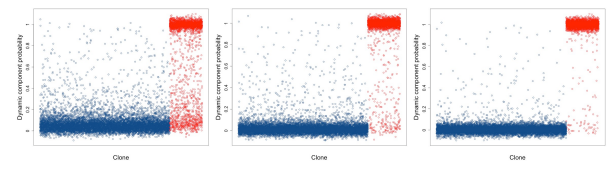
### 3.1 Generating process

We generated 60,000 clonotypes under a combination of  $\alpha$  and  $\beta$  values as shown in Table 1 and with total observation-time offsets sampled from an exponential distribution that approximated the empirically observed total observation-time reads. Clonotype mean proportions were sampled from a Gamma( $\alpha$ ,  $\beta$ ) for the different parameter combinations, and subsequently multiplied by the offset  $O_{jk}$  to be used as the mean parameter in sampling from a Poisson distribution. This mean parameter was either varying or constant across the assumed four followup times as a function of whether the clone was considered dynamic or static, respectively. Depending on the simulation scenario, we fit the variance component mixture model to two, three, or four of the timepoints generated to examine the influence of followup on clonotype classification sensitivity and specificity and generated 10% of clones as dynamic and 90% static. We iterated 100 times for each parameter combination to assess variability. Parameter estimates for  $\hat{\alpha}$ ,  $\hat{\beta}$ , and  $\hat{\pi}$  were calculated with four followup periods.

#### 3.2 Simulation Results

Simulations indicate model fits reflect the parameters under which data are generated. Parameters estimate demonstrate a lack of bias and low variance as shown in Table 1. Convergence criteria under the parameter combinations described were achieved in fewer than 20 EM iterations.

Table 2 shows that under the parameters considered, specificity was nearly perfect and relatively invariant to the component membership threshold range of 0.75 and 0.95. Sensitivity to identifying dynamic clones decreased as a function of the threshold used, whereas it increased significantly in the number of followups, each holding the other variable constant and as expected. The results indicate, as expected, that the more information one has within clone longitudinally, the greater the ability to distinguish between dynamic and static behavior. This behavior is demonstrated visually in Figure 1, as one can see more dense dynamic and static clones near the top and bottom, respectively, of the figures as the number of modelled followups increases.



- (a) Clonotype membership probability with two observed follow-ups.
- (b) Clonotype membership probability with three observed follow-ups.
- (c) Clonotype membership probability with four observed follow-ups.

Figure 1: Model estimated clonotype membership probability for those generated under the static (blue points) versus dynamic models (red points) with two, three, and four within-clone observed follow-ups.

Scenario:					
$\alpha$	$\beta$	$\pi$	$\hat{\alpha} (95\%  \text{CI})$	$\hat{\beta} (95\% \text{ CI})$	$\hat{\pi} (95\% \text{ CI})$
1	100	0.2	1.01 (1.01,1.02)	101.25 (100.32,102.41)	0.2 (0.2,0.2)
1	200	0.2	$1.02 \ (1.02, 1.03)$	204.04 (202.11,206.48)	$0.21\ (0.2,0.21)$
2	100	0.2	$2.01\ (1.99, 2.02)$	100.34 (99.28,101.33)	$0.2\ (0.2,0.2)$
2	200	0.2	2.02(2,2.03)	200.84 (199.19,203.23)	$0.2\ (0.2,0.2)$

Table 1: Component and mixing parameter estimates and confidence intervals under four combinations of  $\alpha$  and  $\beta$  based on 100 iterations of each scenario.

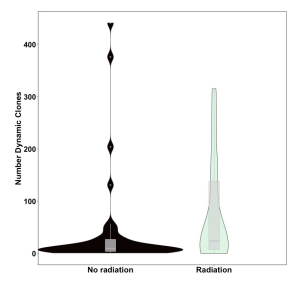
	Sensi	tivity	1-Specificity		
Membership Threshold:	0.75	0.95	0.75	0.95	
2 observed times	0.70 (0.69,0.70)	$0.64 \ (0.64, 0.65)$	0.99 (0.99,0.99)	1.00 (1.00,1.00)	
3 observed times	$0.90 \ (0.90, 0.91)$	$0.88 \ (0.87, 0.89)$	$1.00\ (1.00,1.00)$	$1.00\ (1.00,1.00)$	
4 observed times	$0.97 \ (0.97, 0.97)$	$0.96 \ (0.96, 0.96)$	$1.00 \ (1.00, 1.00)$	1.00 (1.00,1.00)	

Table 2: Sensitivity and specificity rates as a function of component membership threshold and number of observed followup period.

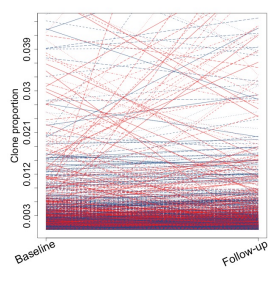
## 4 Results

We applied the model to a longitudinal cohort of 108 prostate cancer patients to whom androgen-deprivation therapy was applied and metastasis-directed radiation therapy (MDT) was randomized [22]. We performed analyses of baseline-followup cross sections, and baseline with two followups, the latter including approximately 15% missingness across the second two followup periods. The total number of productive clonotypes considered across person-times was approximately 4.9 million. After filtering clones to have at least 8 template reads in total across followup, we arrived at 57,436 clones on 91 observations for the baseline-followup analysis, and 86,381 on 104 observations for the baseline with two followups analysis. For each

person-time, the total number of template reads was calculated for use as the logged offset so that variation and therefore component membership would be interpreted on the scale of clonotype proportion, rather than the count of template reads. We fit the model to these data across MDT treatment strata, and also within treatment strata to examine heterogeneity in estimated  $\hat{\alpha}$ ,  $\hat{\beta}$ , and  $\hat{\pi}$ . Once convergence criteria was reached, we examined cluster membership and parameter estimates. Using a threshold component membership probability of > 0.75 to determine the number of unique dynamic clones, we counted the number within patient and partitioned it into those expanding and contracting based on the trajectory of the proportion change over baseline and followup. (EXCLUSIVELY FOR 2 FUS...)

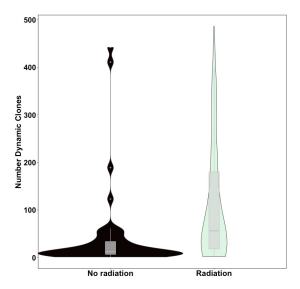


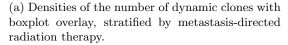
(a) Densities of the number of dynamic clones with boxplot overlay, stratified by metastasis-directed radiation therapy.

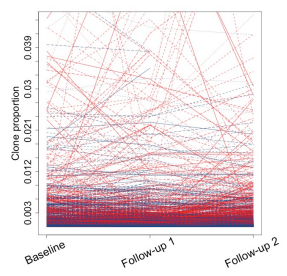


(b) Proportion clonotype over time with dynamic clones shown in red and static clones in dark blue as estimated by the model. Static clones are downsampled by a factor of 2 for visibility of dynamic clones.

Figure 2: Two followup period analysis of baseline-followup cross sections using variance component mixture model.







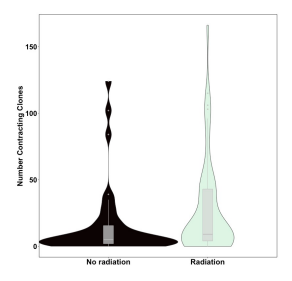
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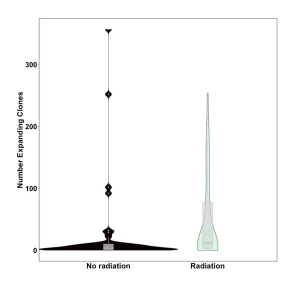
Figure 3: Three followup analysis using variance component mixture model.

Results indicate a significant increase in the number of dynamic clones under MDT for both longitudinal analyses. Figures 2a), 3a), and 4 all show qualitative marked increases in the number of dynamic, expanding, and contracting clones in the MDT treatment stratum, which are formally tested and found significant in nearly all cases by  $\chi^2$  tests and log-linear models (Table 3). The non-MDT stratum had a small number of outlier observations that exhibited a large number dynamic clones.

One observes in Figures 2b) and 3b) that dynamic clones tend to exhibit significant slopes as the clonotype proportion changes dramatically over followup, in contrast to static clones which are flatter. One also sees that the higher average clonotype proportion, the larger change in proportion is needed for a clone to be considered dynamic, incorporating expected biological variability into the model error term.

Within MDT treatment strata models reveal some heterogeneity in estimated  $\alpha$  and  $\beta$  as shown in Table 4, suggesting differences in shape of clonal dynamics over time as a function of treatment. Since  $\alpha$  and  $\beta$  govern both the dynamic and static mixture components, interpretation is not straightforward, though the implication of estimates is that among the MDT stratum, clonotypes mean values tend to be higher and have smaller variance than the non-MDT stratum across time and components. Since clonotype proportions are sensitive to the number of unique clones within observation-times because they must sum to one, interpretation might indicate a tendency toward greater "focus" in dynamism among MDT stratum clones as those useful in immune response expand and those that are not contract.





- (a) Contracting clone count densities with boxplot overlay, stratified by metastasis-directed radiation therapy.
- (b) Expanding clone count densities with boxplot overlay, stratified by metastasis-directed radiation therapy.

Figure 4: Analysis of baseline-followup cross sections using variance component mixture model for number of contractions and number of expansions.

	2 observed follow-ups (FU's)			3 FU's
Clone group:	Dynamic	Expansions	Contractions	Dynamic
$\chi^2$ test p-value	0.01	0.002	0.14	0.001
log-linear parameter (p-value)	$0.64 \ (10^{-08})$	$0.68 \ (10^{-08})$	$0.56 \ (10^{-08})$	$0.98 \ (10^{-08})$

Table 3: Hypothesis tests for dynamic clone stratification by MDT treatment stratum.  $\chi^2$  tests are based on dichotomizing at 50 total dynamic clones or 25 total expanding/contracting clones depending on the case. Log-linear model parameter indicates the log expected proportion increase in the number of dynamic, expanding, or contracting clones, depending on the case, for the MDT stratum as compared to no MDT.

	2 observed follow-ups			3 observed follow-ups		
	Combined	No MDT	MDT	Combined	No MDT	MDT
$\hat{\alpha}$	0.36	0.36	0.37	0.33	0.37	0.32
$\hat{eta}$	749.14	603.41	857.13	715.33	648.81	752.05
$\hat{\pi}$	0.24	0.22	0.25	0.20	0.19	0.20

Table 4: Model parameter estimates for two and three study follow-ups, across and within the metastasis-directed radiation therapy (MDT) and lack of MDT (No MDT) treatment strata.

# 5 Discussion

In this work we have placed T-cell receptor clonal dynamics in a mixture model framework grounded in Bayesian hierarchy and argued the approach can be a useful tool for understanding these high-dimensional data that duly acknowledge expected biological variability. Since receptors shared across subjects are rare relative to those unique to each subject, traditional statistical high-dimensional regression tools are not as useful and one must try to relate summarizations of clonal population dynamics to subject phenotype, in our case a count of model-determined dynamic clones. Additionally partitioning dynamic clones into those that expand and contract as a function of their trajectory further allows one to examine the kind of dynamism most sensitive to intervention, in our case metastasis-directed radiation therapy.

While the mixture of negative multinomial and product of negative binomial components arrived at through the posterior predictive distribution of Gamma-Poisson conjugacy was both theoretically convenient and reflective of observed data, future work may well attempt flexible alternatives for which more computationally intensive fitting procedures are necessary. Since the single parameter Poisson distribution equates the mean and variance, enforcing the relationship between clone frequency and degree of variability constituting "dynamism", a model allowing one to specify that relationship could help clinicians and subject experts place greater weight on, say, dynamism among the low-frequency group of clones or vice versa, depending on hypothesized biological mechanism. The mean variance equivalence of the Poisson does have advantages in that since coverage increases in the root of the variance, one is better powered to identify dynamism among higher frequency clones for less variability relative to that mean, consistent with a biological prior that higher frequency clones are inherently more interesting and should be easier to label as dynamic. But still, exploring sensitivity to different and more flexible mean-variance specifications could improve characterization of the dynamism associated with subject phenotype.

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