Inferring ongoing cancer evolution from single tumour biopsies using synthetic supervised learning

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Abstract

Variant allele frequencies (VAF) encode ongoing evolution and subclonal selection in growing tumours. However, existing methods that utilize VAF information for cancer evolutionary inference are compressive, slow, or incorrectly specify the underlying cancer evolutionary dynamics. Here, we provide a proof-of-principle synthetic supervised learning method, TumE, that integrates simulated models of cancer evolution with Bayesian neural networks, to infer ongoing selection in bulk-sequenced single tumour biopsies. Analyses in synthetic and patient tumours shows that TumE significantly improves both accuracy and inference time per sample when detecting positive selection, deconvoluting selected subclonal populations, and estimating subclone frequency. Importantly, we show how transfer learning can leverage stored knowledge within TumE models for related evolutionary inference tasks — substantially reducing data and computational time for further model development and providing a library of recyclable deep learning models for the cancer evolution community. This extensible framework provides a foundation and future directions for harnessing progressive computational methods for the benefit of cancer genomics and, in turn, the cancer patient. TumE is publicly available for use at https://github.com/tomouellette/TumE.
Introduction

Cancer is a disease characterized by unrelenting tissue growth and clonal evolution. During evolution, genetic and epigenetic aberrations provide the reservoir for dysfunctional cellular phenotypes that maintain a tumour’s replicative advantage, while, over time, fluctuating physiological and ecological properties within the tumour microenvironment drive the need for updated adaptations that sustain immortality. Overall, the complex interplay between mutation accumulation and microenvironmental changes leads to a high degree of both cellular and genetic heterogeneity and, by proxy, composite subclonal structure in tumours.

Naturally, the desire to better understand the evolutionary and subclonal dynamics in growing tumour populations has become a major task for cancer genomics - with goals of forecasting tumour progression, developing adaptive evolutionary therapies, and deconvoluting the genetic architecture that drives adaptation.

However, a significant hurdle in understanding cancer evolution in vivo are the clinical constraints surrounding serial sequencing, through space or time. For this reason, tumour biopsies are primarily sequenced in bulk from a single site and at a single time point. Although multi-region and single-cell data are becoming increasingly utilized, single time point, bulk sequenced biopsies still represent the major accessible data source for precision genomics guided treatment and for studying cancer genomics and evolution in patients. Given this limitation, a reasonable strategy for inferring evolution in single tumor biopsies has been to utilize theoretical population genetics to capture signatures of selection from the variant allele frequency (VAF) distribution. The premise being that fitness-altering mutations will deterministically change in frequency over time, leading to characteristic and quantifiable deviations in the VAF distribution relative to some neutral evolutionary scenario.

VAF-based methods have been employed to differentiate between positive selection and neutral evolution, to examine growth patterns, to quantify subclonal fitness and time subclonal emergence, and to build population genetics informed mixture models that account for neutral dynamics, that shape, to some extent, all tumour populations. With that said, existing VAF-based methods used to infer cancer evolution, although mechanistic and useful, have apparent limitations. For example, single statistics are maximally compressive and cannot infer complex information, approximate Bayesian computation methods suffer from the curse of dimensionality and can be prohibitively slow due to a rate-limiting simulation step required for each sample, and mixture models, used to identify subclonal populations, are only implicitly connected to an underlying model of evolution and, until recently, have been built under incorrect assumptions that have led to systematic overestimation in the number of subclonal populations in sequenced tumours.

To address these limitations, we contribute a proof-of-principle synthetic supervised deep learning approach, TumE, for quantifying and classifying the evolutionary and subclonal dynamics in bulk sequenced
tumours biopsies using purity-corrected variant allele frequency (VAF) information from diploid genomic regions. By generating synthetic VAF distributions, as a proxy for evolutionary ground truth, from plausible simulations of tumour evolution, we were able to build inference models that accurately classify and quantify evolutionary (e.g. positive selection versus neutral evolution) and subclonal dynamics (e.g. subclone frequency) in real patient tumours while capturing uncertainty in our estimates, via a form of approximate Bayesian inference called Monte Carlo dropout\textsuperscript{26,27}. Importantly, our method further highlights the power of utilizing deep learning for inference - namely exploiting stored knowledge via transfer learning. By recycling our models for new evolutionary prediction tasks, we reduce the computational burden associated with the generation of synthetic or simulated data. We validated our synthetic supervised learning approach in millions of synthetic tumours and applied TumE to 95 copy-number and purity corrected whole-genome (WGS) and whole-exome (WES) sequenced tumour biopsies.

**Results**

**Inferring cancer evolution using synthetic supervised deep learning**

Synthetic supervised, or simulation-based, deep learning has been shown to be equivalent to amortized approximate inference under a generative model\textsuperscript{28}. Therefore, by optimizing a neural network using realistic synthetic data $x$ generated from a stochastic generative process $p(x,z|\Theta)$, where $\Theta$ indicates the prior or parameters that define the simulation and $z$ indicates the latent variables generated during simulation, we can build inference models that approximate our true posterior of interest $p(\Theta,z|x)$. In our case, by optimizing a neural network using synthetic VAF distributions sampled from $p(x,z|\Theta)$, we can build inference models for evolutionary inference in sequenced tumour biopsies (Figure 1A-C; Methods).

**Figure 1.** (a) TumE integrates a generative sampling process and stochastic simulation of cancer evolution to build well-specified synthetic variant allele frequency (VAF) distributions with respect to data observed in bulk sequenced tumour biopsies. Assuming copy-neutral diploid regions of tumour genomes, the generative sampling process uses the observation that neutral VAF distributions can be described by a power-law or Pareto neutral "tail"\textsuperscript{16,29} in addition to a dispersed clonal peak. By sampling empirically valid Pareto distributions, rapid realizations of the null hypothesis of neutral evolution encoded in the VAF distribution can be created (Methods). The stochastic branching process model of tumour evolution is then used to link parameters and latent states, relevant to positive subclonal selection, back to VAF distributions (Methods). (b) Synthetic supervised learning utilizes neural networks capable of handling the complete dimensionality of the simulated VAF distributions, $x$, to solve the inverse problem of identifying the evolutionary parameters and latent states, $y$, assigned to each synthetic VAF distribution. (c) We can then quantify model uncertainty using a computationally efficient form of Bayesian deep learning called Monte Carlo dropout\textsuperscript{26,27}. Approximate posteriors are generated by performing $T$ stochastic passes through the trained neural network.
To generate synthetic data that properly captured evolutionary dynamics in patient tumours, we implemented a simulation framework, i.e. a stochastic generative process $p(x, z|\Theta)$, combining two complementary approaches to improve the speed and efficiency of synthetic data generation — one for tumours subject to positive selection and one for tumours evolving neutrally (Figure 1A). For growing tumours simulated with positive selection, we utilized a well-established framework of cancer evolution that models exponential tumour growth under a stochastic branching process and coupled this with a virtual biopsy procedure to account for sequencing noise observed in real patient tumours (adapted from ref7). In our model, we allowed for a completely stochastic arrival of driver mutations that multiplicatively increased the fitness of mutated subclones and tracked the frequency of each subclone until the time of virtual biopsy (Methods). In this study, we define a subclone as a subpopulation of cells with a fitness or growth rate advantage relative to the background population (Methods) and consider subclones detectable if they are between ~10 - 40% VAF (20 - 80% cellular fraction). For tumours that lacked selected subclones (neutrally evolving), we implemented a generative sampling process based on the observation that VAF distributions from tumours without positively selected subclones can be described by a power-law or Pareto distribution in conjunction with a dispersed clonal peak (Supplementary Figure 1). Concisely, this process involved i) sampling allele frequencies from empirically realistic Pareto distributions to generate the neutral power-law ‘tail’ in the VAF distribution, ii) adding additional diploid clonal heterozygous mutations at 50% VAF, and then iii) injecting additional sequencing noise under a beta-binomial model (Methods). In general, a complete VAF distribution indicative of positive selection, and computed from heterozygous diploid mutations, includes a neutral power-law tail, a heterozygous clonal peak centered at ~50% VAF, and additional subclonal peak(s) in the intermediate frequency ranges (~10 - 40% VAF); whereas a neutrally evolving tumour, or one with undetectable selected subpopulations, lacks the characteristic subclonal peak(s) (Figure 1A). To ensure positively selected and neutrally evolving synthetic tumours were not out of distribution with each other given the alternate data generation approaches, we simulated synthetic tumours in pairs, assigning the neutral VAF distributions with equivalent parameters and mutations with respect to the paired positive selection simulation (Methods; pseudo algorithms and examples provided alongside Supplementary Figure 2 & 3).

Using this framework, we generated approximately 40 million synthetic tumours across varying mutation rates, selection coefficients, and sequencing noise parameters. We selected broad simulation parameter ranges that were consistent with previous computational studies and empirically estimated values (Methods; Supplementary Table S1). By generating synthetic tumours using well-specified simulations (comparison of real and synthetic data outlined in Methods and Supplementary Figures 3 - 5), we were able to explicitly link each VAF distribution to the parameters and latent states that defined the underlying subclonal and evolutionary dynamics. We then used the millions of annotated synthetic VAF distributions to train hundreds of neural networks using a random hyperparameter search to make inferences on the evolutionary mode (positive selection or neutral evolution), the number of subclones (0, 1, or 2), and the
subclone frequency at borderline to optimal sequencing depths (50 - 250X) for evolutionary analysis in cancer genomics (Figure 1B; Methods). To capture model-based uncertainty in our estimates, we implemented a form of Bayesian approximation for deep learning called Monte Carlo (MC) dropout\textsuperscript{26,27} (Figure 1C; Methods). We used MC dropout to mitigate overconfident estimates in cases of high uncertainty or broad approximate posteriors. In general, we structured both neural network training and prediction to favour the more parsimonious explanation of the data (fewer subclones and neutral evolution; Methods). We show how using a classification threshold based on a lower bound of the MC dropout approximate posterior helps mitigate model overconfidence across changing subclone mutation and frequencies in Supplementary Figure 6. Following training, we selected the top scoring models, for predicting the evolutionary mode, number of subclones, and subclone frequency, for further validation (Methods).

We outline the full synthetic supervised learning pipeline in Methods. In addition, we highlight that even though we model VAF distributions in patient tumours using point mutations from diploid regions, mutations in our framework, as with previous approaches\textsuperscript{7,13,16}, are agnostic to the underlying functional alteration, e.g. missense, silent, driver or copy number driving selection in patient tumours. This is because genome-wide linkage, a by-product of zero recombination, results in hitchhiking of any additional point mutations on the genetic background of any selected clone\textsuperscript{3,18}.

**Comparison of synthetic supervised learning to existing methods**

To evaluate TumE performance on inferred estimates of selection, number of subclones, and subclone frequency, we simulated an additional ~2.8 million synthetic tumours under neutral evolution (0 subclones) and positive selection (1 or 2 detectable selected subclones) assessing the impact of variable sequencing depths (50 - 250x coverage) and read count overdispersion (0 - 0.3 rho) (Methods). We first compared TumE against frequency-based summary statistic approaches for differentiating between neutral evolution and positive selection and found that TumE significantly outperforms recently developed VAF summary statistics\textsuperscript{12} (two-sided Wilcoxon test, \( p = 2.7 \times 10^{-12} \)) as well as common population genetic summary statistics\textsuperscript{20,21} (two-sided Wilcoxon test, \( p = 1.9 \times 10^{-8} \)), based on AUROC (Figure 2A). Further, TumE outperforms each statistic individually when compared across all sequencing depth and overdispersion combinations analyzed here (ROC analysis; Supplementary Figure 7).

We next compared TumE against the only mixture model approach, MOBSTER\textsuperscript{16}, that explicitly and correctly takes into account the neutral dynamics within sequenced tumour VAF distributions to detect subclones. We found that TumE provides comparable or improved performance for predicting the number of subclones (precision-recall, Supplementary Figure 8) and for predicting subclone frequency (Figure 2B; correlation and mean absolute percentage error, Supplementary Figure 9) across all empirically relevant depth (50 - 250x coverage) and read count overdispersions (0 - 0.003 rho) combinations. However, as expected, we found that the performance of TumE and MOBSTER both degrade as sequencing depth
decreases (≤ 75x coverage) and overdispersion increases (≥ 0.01) under a beta-binomial sequencing noise model (Supplementary Figure 8 & 9). Furthermore, additional analysis of subclone frequency estimates in the 2 subclone setting revealed that as inter-subclone distance increases, i.e. overlap of subclonal peaks decrease, the mean percentage error for predicting the frequency of both the lowest and highest frequency subclone decreases towards zero (Supplementary Figures 10 - 12).

**Figure 2** (a) In a cohort of 2.8 million synthetic tumours, TumE outperformed all existing common population genetic\textsuperscript{20,21} and cancer evolution\textsuperscript{7,12} specific summary statistics when differentiating between positive selection and neutral evolution, based on AUROC (two-sided Wilcoxon test). (b) Further, for predicting the true frequency of selected subclones, TumE provides comparable or better performance relative to the current state-of-the-art mixture model MOBSTER\textsuperscript{16} that properly accounts for neutral dynamics in tumour populations. The panel shows correlation coefficient ($r$) between the true and predicted subclone frequency in 80,000 synthetic tumours sequenced at 150x mean sequencing depth. (c) In an orthogonal dataset of 150 synthetic tumours\textsuperscript{16} with either 0 or 1 detectable subclones, TumE was significantly faster at estimating the number of subclones (two-sided Wilcoxon test) than existing mixture model based methods sciClone\textsuperscript{24} and MOBSTER\textsuperscript{16} (measured in inference time per sample). In addition, only TumE and MOBSTER consistently identified the correct number of subclones, as both methods directly account for the neutral dynamics observed in tumour populations. (d) TumE estimates in a synthetic tumour sequenced at 120x mean sequencing depth and a subclone at 54% cellular fraction.

Given our simulation framework was based on certain approximating assumptions to improve computational speed and efficiency (namely small population size and no cell death; outlined in Methods), we sought to perform additional validation of evolutionary estimates in an alternative dataset of synthetic tumours\textsuperscript{16}. The orthogonal dataset, described in Caravagna et al. 2020\textsuperscript{16}, consisted of 150 synthetic tumours, 40 effectively neutral and 110 with one detectable subclone (between 10 - 45% VAF), sequenced to 120x depth and grown to a population size of >10\textsuperscript{8} cells at birth rate of 1 and death rate of 0.2. To frame our predictions relative to existing methods, we applied TumE, MOBSTER, and a variational Bayesian mixture model sciClone\textsuperscript{24} to the synthetic dataset. To make comparisons fair, we limited the maximum
number of subclonal cluster assignments to 2 for both MOBSTER and sciClone, as this was the upper bound on TumE estimates (Methods). For sciClone, this meant setting the maximum number of mixture components to 4 (neutral tail, 2 subclones, and a clonal peak) as sciClone doesn’t properly account for neutral dynamics (Pareto tail) observed in sequenced tumour populations. Both TumE and MOBSTER consistently identified the correct number of detectable subclones in the majority of cases while sciClone systematically overestimated the number of subclones, even after correcting estimates for the clonal peak and neutral tail (Figure 2C). However, relative to both sciClone and MOBSTER, TumE provided orders-of-magnitude faster estimates (two-sided Wilcoxon test, \( p < 2.2 \times 10^{-16} \), Figure 2C), reducing run times per sample from minutes to \( \sim 1 \) second. We provide individual estimates with TumE for each of the 150 synthetic tumours, and an additional 750 synthetic tumours of variable sequencing depth from ref\(^{16}\), in Supplementary Figures 13 & 14. We provide an example TumE output for a synthetic tumour with a single detectable subclone in Figure 2D.

In this study, we note that the birth and death rate were set to fixed values (birth rate = \( \log(2) \), death rate = 0, in line with ref\(^7\)) to additionally improve the computational efficiency of the stochastic simulations of positively selected tumour populations. Therefore, one additional factor that may impact the accurate detection of selection and subclones with TumE is variable birth and death rates in growing tumours. For example, an elevated cell death can lead to an increase in the number of passenger mutations that are swept to higher frequencies during subclonal selection. In regards to the VAF distribution, this elevated number of mutations ‘trailing’ the subclonal peak may obscure lower frequency subclones or, alternatively, lead to spurious identification of additional subclones due to an elevated number of neutral mutations entering the subclonal frequency range. To assess the impact of variable growth rates, we generated an additional 6 million synthetic tumours across 26 different birth and death rate combinations (simulation parameters outlined in Supplementary Table S1). Overall, we find that our estimates are robust to changes in tumour growth rates. Any errors that do occur only appear to increase the number of parsimonious explanations of the data (e.g. classifying 2 subclones as 1; Supplementary Figure 15). In addition, the prediction of subclone frequency also remained consistent across all the birth and death rate combinations evaluated here (Supplementary Figure 16).

### Analysis of whole-genome and exome sequenced tumour biopsies

To make the utility of synthetic supervised learning concrete, we first evaluated TumE in ‘gold-standard’ tumour biopsies commonly used to evaluate mixture model based approaches, namely a deep sequenced (~320x coverage, 90.7% purity) acute myeloid leukemia (AML) sample from Griffith et al.\(^{31}\) and a deep sequenced (~226x coverage, 71.2% purity) breast adenocarcinoma sample retrieved from the pan-cancer analysis of whole genomes (PCAWG)\(^{11}\) but originally from ref\(^{32}\). In both cases, we recovered the correct evolutionary mode, number of subclones, and subclone frequencies (Figure 3A & 3B). In addition, because we provide accurate subclone frequency estimates, we performed heuristic clustering of the clonal,
subclonal, and neutral tail mutations by using the expected variance under a binomial sequencing noise model (Methods). This heuristic approach facilitates subclonal clustering at almost zero additional computational cost (as observed in the total runtime per sample of ~1s).

Figure 3. TumE estimates in deep whole-genome or whole exome sequenced tumour biopsies. (a) A deep-sequenced primary acute myeloid leukemia (AML) sample from Griffith et al.\textsuperscript{31}. TumE estimated two subclones, a neutral tail, and a clonal peak. P(Selection) indicates the probability of selection. P(0, 1, 2 subclone) indicates the probability estimate for the number of subclones. Each probability estimate is provided with the 89\% equal-tailed interval generated from 50 Monte Carlo dropout samples. A sample is labeled positive selection if the lower bound of the 89\% interval is above P = 0.5, and the number of subclones is assigned to a sample if the lower bound of the 89\% interval is greater than 0.5 (Methods). Subclone frequency estimates are shown with the complete approximate posterior. (b) A deep-sequenced breast adenocarcinoma from the pan-cancer analysis of whole genomes\textsuperscript{11} (PCAWG). TumE estimated two subclones, a neutral tail, and a clonal peak. (c) We applied TumE to a single mismatch repair deficient (MMR) gastrointestinal tumour sequenced across 5 spatially distinct regions. We first identified an intermediate frequency subclone in region P with TumE. (d) Under the hypothesis that TumE could reveal the fixation process of region P subclones in other regions, we annotated each of the remaining regions with the clonal, subclonal, and neutral tail mutations identified in region P. We identified ongoing subclonal selection in 2 out of the 4 remaining regions (N and T) consistent with an increase in frequency of subclonal and neutral tail mutations from region P. In cases where neutral evolution was the most parsimonious explanation, we observed complete fixation of the subclonal region P mutations (region AE and H).

We next evaluated TumE in whole-exome sequenced (WES) mismatch repair deficient (MMR) gastrointestinal tumours biopsied across multiple spatially distinct regions (collected from von Loga et al.\textsuperscript{33}). As evolutionary inference requires high-quality genomes, we only included samples that had a mean effective coverage (mean sequencing depth \textbf{\textit{*}} purity) greater than 50x and a minimum purity of 50\%. We note that ~70x mean sequencing depth has been suggested as the minimal threshold for accurate estimates\textsuperscript{7,16}, as
we also observed (Supplementary Figures 8 & 9). Following removal of low quality biopsies, we retained biopsies from two tumours with one tumour retaining 5 spatially distinct (WES) biopsies. TumE estimates in the 5 spatially distinct biopsies from a single tumour revealed the fixation process of a positively selected subclone, from intermediate frequency to metastasis fixation (Figure 3C & 3D). In addition, the application of TumE to multi-region samples highlighted the ability of TumE to pick up signatures of selection not directly encoded in distinct subclonal peaks but in the asymmetry of the diploid heterozygous cluster (region N & region T, Figure 3D).

Finally, we evaluated TumE in 85 whole-genome sequenced (WGS) tumour biopsies, spanning 8 different cancer types, retrieved from the pan-cancer analysis of whole genomes (PCAWG). In total, 38.8% of samples showed evidence for positive, or subclonal, selection whereas the majority, 61.2%, were adequately described by neutral evolutionary dynamics (Supplementary Table S2). Alternative methods applied to large cancer cohorts, including PCAWG, have estimated that as few as 3% to upwards of 96% of samples show evidence for ongoing subclonal selection. The discrepancy is likely explained in modeling approaches. For example, low estimates are a by-product of utilizing mixture models that rely on distinct and ‘clean’ subclonal peaks whereas high estimates likely occur from not taking into account the neutral dynamics in tumour evolution. In contrast, TumE generates a non-linear encoding of the VAF distribution, extracting novel representations that increase accuracy while simultaneously accounting for the correct neutrality evolutionary dynamics observed in tumour populations. All samples analyzed in this study, including MMR gastro-esophageal and deep-sequenced AML, are outlined in Methods and Supplementary Figure 16 - 18.

**A transfer learning framework to infer additional evolutionary parameters**

One drawback of simulation-based deep learning approaches is the requirement for the repeated generation of synthetic data for training. Although this allows for fast inference at test time through amortization, altering the models assumptions or changing the parameters being inferred generally requires simulating a completely new set of data and training an entirely new set of models - a computationally expensive process. Practically, overcoming this limitation would provide substantial reductions in the amount of time and data needed to build accurate models and would make simulation-based approaches more accessible to the general user. Therefore, we hypothesized that our trained deep learning models could be used as a source of ‘stored’ knowledge for related evolutionary inference tasks that also used the VAF distribution as input.

To explore this possibility, we implemented a transfer learning pipeline, based on domain adaptation, to make inferences on additional parameters using a previously developed cancer evolution simulator, TEMULATOR, that was built under a modified set of assumptions relative to our multiplicative fitness
framework (Methods, viable parameter combinations for detectable subclones outlined in Supplementary Figure 20). In this study, we employ open set domain adaptation\textsuperscript{36} where the structure of the input space, i.e. the VAF distribution, is retained whereas the outputs, the evolutionary tasks, are modified. Briefly, this pipeline involved generating new synthetic tumour sequencing data using TEMULATOR, performing

![Architecture renovation](image)

**Figure 4.** (a) Transfer learning approach utilizing ‘renovated’ pre-trained neural networks for alternative evolutionary inference tasks in tumour cellular populations. TEMULATOR is an alternative cancer evolution simulator that generates synthetic tumour sequencing data by deterministically initiating subclones at user specified fitnesses and time points\textsuperscript{37}. (b) Pre-trained models provide significant reductions in testing loss, over non-pretrained models, when updating neural network weights on reduced dataset size of 500,000 synthetic VAF distributions (~1.25% of the total dataset size used to originally train TumE). (c) TumE transfer (TumE-T) effectively recovers evolutionary parameters from TEMULATOR simulations (75 - 200x mean sequencing depth, 100% tumor purity) with mean and median percentage errors less than 10% in all cases. A full description of performance across variable sequencing depths, mutation rates, and subclone frequencies is provided in Supplementary Figure 23. (d) We find consistency between the subclone cellular fraction estimated by TumE-T and the subclone frequency (cellular fraction / 2) estimates generated from TumE, indicating nearly identical tasks are easily transferred through pre-training. (e) Per genome per division mutation rate estimates in 95 WES and WGS samples from von Loga et al.\textsuperscript{33} (MMR-GE = mismatch deficient repair gastro-esophageal cancer), Griffith et al.\textsuperscript{31} (AML = acute myeloid leukemia), and PCAWG\textsuperscript{11}. (f) Subclone fitness (1+s) estimates (relative growth rate advantage of subclone over background population) and (g) subclone emergence time estimates in 30 tumour biopsies identified with 1 subclone in the PCAWG data. Subclone fitness and emergence time estimates were scaled to a final tumour population size of 10\textsuperscript{10} cells, similar to ref.\textsuperscript{7}. PCAWG sample identifiers are provided on the x-axis. Boxplots for subclone fitness and emergence time indicate median estimate and 1.5x interquartile range (whiskers) over 500 Monte Carlo dropout samples from TumE-T.

architecture ‘renovation’ on pre-trained TumE neural networks to replace existing task-specific branches with new ones, and re-tuning the neural network weights and hyperparameters for optimization on the new
evolutionary inference tasks (Figure 4A). The evolutionary inference tasks included predicting subclone fitness, subclone emergence time, mutation rate, and subclone cellular fraction (subclone frequency * 2).

To highlight the benefit of using pre-trained models on related evolutionary inference tasks, we opted to update network weights with only 500,000 synthetic VAF distributions, representing only a fraction (~1.25%) of the data used in the original training of TumE. Each VAF distribution was generated by simulating synthetic tumours with TEMULATOR at a birth rate of 1, death rate of 0.2, final population size of ~10^4, and with either 0 or 1 detectable subclone. The remaining parameters, such as mutation rate, were uniformly sampled from empirically plausible ranges (Supplementary Table S3).

Initially, we used the 500,000 synthetic VAF distributions to compare pre-trained vs non-pretrained models for predicting the evolutionary and subclonal parameters in the presence of 1 subclone. To ensure valid comparisons, we performed a random hyperparameter search, tuning the learning rate and number of fully connected layers in the new task specific branches, across both the pre-trained and non-pretrained model groups. Both groups shared identical neural network architectures. When initially evaluating ~300 pre-trained and non-pretrained models on an external test set of 3000 synthetic tumours, we found that pre-trained models obtained a significantly lower testing loss (mean absolute error across all tasks, two-sided Wilcoxon test, p < 2.22 x 10^{-16}, Figure 4B). Further, when evaluating the top performing pre-trained and non-pretrained models on an additional 400,000 synthetic tumours, pre-trained models obtained significantly lower mean percentage errors, relative to non-pretrained models, for predicting the mutation rate, subclone emergence time, subclone fitness, and subclone frequency (two-sided Wilcoxon test, p < 1.7 x 10^{-8} on all tasks, Supplementary Figure 21).

Next, we selected the top performing pretrained model, TumE transfer (TumE-T), for further validation. We initially found a modest yet systematic underestimation of the mutation rate (~50% mean percentage error). However, this was easily corrected with a post-hoc adjustment by re-fitting the predicted mutation rate to a set of 1000 synthetic tumours using polynomial regression (degree = 2). Evaluating the updated mutation rate estimates on a holdout set of 100,000 synthetic tumours validated the post-hoc adjustment (Supplementary Figure 22). Overall, we were able to effectively recover all evolutionary parameters in the 100,000 synthetic tumours with mean and median percentage errors lower than 10% in all cases (Figure 4C). The performance was also consistent across sequencing depths and mutation rates, however, as expected, we could only effectively assign subclonal parameters, such as fitness, at detectable subclone frequencies (~10 - 40% VAF; Supplementary Figure 23).

Applying TumE-T to the 95 WGS and WES samples described above, we found strong correlation between predicted subclone cellular fraction and subclone frequency estimated by the original TumE models, suggesting nearly identical tasks are easily transferred to new source-target distributions when using pre-trained models (Figure 4D). With respect to mutation rates, estimates were consistent with the general
trends observed empirically\textsuperscript{11,33} - with mismatch repair deficient tumours showing extremely high mutation rates (>100 per genome per division) and acute myeloid leukemia showing very few (Figure 3E). For subclone fitness and subclone emergence time estimation, we had to take into account the difference between simulated and true population sizes\textsuperscript{7,19}. In this regard, we rescaled our estimates to account for a true tumour population size of $10^{10}$, similar to ref\textsuperscript{7}. With rescaling, TumE-T subclone fitness estimates, defined as the relative growth advantage of the selected subpopulation over the background population, ranged from $\sim$1.9 to 2.6 (Figure 4F) while subclone emergence time estimates ranged $\sim$20 to 24 tumour doublings (Figure 4G) in samples with ongoing subclonal selection. We note that emergence times of $\sim$20 to 24 tumour doublings represent approximately 0.001 to 0.16\% of the final tumour volume, which is consistent with theory and empirical evidence suggesting subclones must arise early during tumour growth to reach detectable frequencies\textsuperscript{7,14,17,38}.

Discussion

In this study, we developed a synthetic supervised learning approach, TumE, for cancer evolutionary inference. Overall, the synthetic supervised learning approach, TumE, provides four major advantages. First, by generating synthetic data using models of cancer evolution, we are able to explicitly account for the neutral and non-neutral evolutionary dynamics observed in tumour VAF distributions\textsuperscript{7,12,16}, thereby avoiding systematic overestimates in the number of subclones due to misclassifying low frequency neutral ‘tails’. Second, by using neural networks that can naturally handle high-dimensional VAF distributions as input, we avoid information loss that comes with compressing data into a single statistic, or distance metric, prior to inference, improving model accuracy across all evolutionary inference tasks considered here. Third, by separating simulation and model training from prediction, via amortized inference, we significantly decrease inference time per sample, reducing time from minutes to seconds relative to existing methods. Finally, we show how we can use open set domain adaptation\textsuperscript{35,36}, a form of transfer learning, to recycle our models for alternative evolutionary inference tasks that use VAF distributions as input - drastically reducing the number of synthetic samples and computational time required for further model development. Our library of pre-trained models benefits all researchers building inference machines for cancer evolution prediction, even in a non-deep learning setting. For example, providing fast, evolutionary-informed peak initializations for mixture model based methods.

We mention some current limitations. Firstly, as a neural network requires optimization on a finite, static set of data, estimates, without transfer learning, are constrained to a pre-defined search space. In this study we focused on cancer evolution in the context of 2 detectable selected subpopulations captured from frequency information in diploid genomic regions. Although multiple studies have shown it’s rare to detect 2, or even 1, subclones\textsuperscript{7,14,16,17} in noisy one-dimensional VAF distributions, it’s possible we do not capture extreme cases of selected subclonal heterogeneity. Furthermore, focusing on diploid regions may obscure the detection of ongoing selection if mutations are concentrated in copy number aberrated segments.
However, constraining analyses to diploid regions provides a strong baseline for model development, while genome-wide linkage provides biological justification for analyzing diploid regions. Finally, our model of tumour evolution was structured to reflect the biopsy material available here, namely bulk sequenced single site and time point data. We note that tumour growth over space and time can have profound effects on the detectability of selection\textsuperscript{39,40}. In this regard, TumE estimates can still be applied in a localized setting and aggregated globally. Nevertheless, more structured ways of integrating a synthetic supervised learning approach with multi-region data are necessary for maximizing utility.

Altogether, in this study, we exhibit how coupling well-specified synthetic data with neural networks provides fast and accurate amortized estimates that go beyond the current paradigm of single statistics, mixture models, and approximate Bayesian computation for classifying and quantifying ongoing selection in tumour populations. The integration of generative and simulation-based models of cancer evolution with modern deep learning frameworks facilitates robust and efficient estimates of evolutionary and subclonal dynamics in growing tumour populations. This extensible framework provides future avenues for harnessing progressive computational methods for the benefit of cancer genomics and, as an end goal, the cancer patient.

**Methods**

**Synthetic data generation**

We generated synthetic data that encoded the evolutionary dynamics observed in the variant allele frequency (VAF) distribution (namely the neutral tail, subclones, and clonal peaks) using two complementary approaches dependent on the underlying evolutionary mode - one for tumors subject to positive selection and one for tumours evolving neutrally.

For tumours simulated under positive selection, we utilized a well-established framework of cancer evolution that models exponential tumour growth under a stochastic branching process\textsuperscript{7,12,13,15,19,30} and coupled this with a virtual biopsy procedure to account for sequencing noise observed in whole-genome/whole-exome sequenced tumours from real patient tumours. For implementation, we adapted a previous cancer evolution framework developed by Williams et al.\textsuperscript{7} Briefly, this model simulates exponentially growing tumour populations under a stochastic branching process using a rejection-kinetic Monte Carlo (MC) algorithm, where a given cell accrues mutations at some Poisson-distributed per genome per division rate $\mu$ and divides or dies with probabilities proportional to its birth or death rate. This branching process continues by randomly sampling existing cells, weighted by cellular fitness, until a final tumour population size $N$, sufficient to recapitulate the features of empirical VAF distributions, is reached. Following completion of each simulation, a virtual biopsy procedure to account for sequencing noise observed in real patient VAF distributions is implemented. In this sequencing noise model, the observed
frequency for a given mutation ($VAF_{obs}$) relative to the true underlying frequency ($VAF_{true}$) in a tumour of population size $N$ is given by

$$VAF_{obs} = \frac{R_{obs}}{D_{obs}}$$

where $D_{obs}$ indicates the total observed read depth, $R$ indicates the number of observed reads covering the mutation, $VAF_{true}$ indicates the true population frequency of the mutation, and $\rho$ indicates the overdispersion parameter for the beta-binomial.

In this study, we modify the Williams et al\textsuperscript{7} framework in two ways. Firstly, we implement a fully stochastic arrival of subclones (driver mutations) rather than deterministically injecting a subclone with a specified fitness at a given time $t$. Secondly, the fitness of a subclone or cell is dictated by the multiplicative fitness of all driver mutations. Therefore, when a driver mutation does occur, based on some probability $p_d$, it is assigned a selection coefficient $s > 0$ sampled from an exponential distribution which increases the cell’s growth rate ($b - d$) by a factor of $(1 + s)$ i.e. the fitness. In the case of multiple driver mutations, the fitness of a given cell increases multiplicatively i.e. $\Pi (1 + s)$. Although this random injection of driver mutations is more computationally intensive, it implicitly captures a wider variety of potential frequency distributions without hard coding additional parameter settings - for example, when additional subclones, beyond 1 or 2, are present at undetectable frequencies (e.g. $>40\%$ or $<10\%$). In this study, we consider up to 2 detectable subclones but allow for up to 3 selected subclones to be present at the time of biopsy (see Simulation Parameter Selection below for more details).

For tumours simulated under neutral evolution, we use a generative sampling process for producing neutral VAF distributions, rather than using the stochastic simulation framework. We implement this sampling process because we use a small $N$ population size approximation to generate VAF distributions in our stochastic simulations (using a small $N$ allows us to increase simulation speed and efficiency, which makes generating millions of synthetic VAF distributions practically feasible). Although using a small $N$ is reasonable since the VAF distribution contains no information on population size\textsuperscript{7} (a final simulated tumour population size $N$ of $10^3$ - $10^4$ has been shown to be sufficient to recapitulate the properties of empirical VAF distributions\textsuperscript{7}), neutral stochastic simulations have a higher probability of returning late-occurring spurious subclones due to chance or, in empirical terms, genetic drift. Given the quality of the synthetic data impacts deep learning model performance, we utilize the fully synthetic generative sampling scheme to avoid misspecified data relative to the expected null model of neutral evolution.
The neutral generative sampling process we implement is based on the observation that neutrally evolving asexual, non-recombining populations, such as cancer populations, have VAF distributions (excluding clonal mutations) that follow a power-law or Pareto distribution\textsuperscript{16,29}. Therefore, a VAF for any mutation $i$ in the neutral tail of a frequency distribution can be realized by sampling

$$VAF_i \sim f(x \mid \alpha, m) \text{ with } f(x \mid \alpha, m) = \alpha m^\alpha x^{-(\alpha + 1)}$$

where $\alpha$ is the shape parameter and $m$ the scale parameter for the Pareto distribution.

Given that the generative process for neutral tails is known\textsuperscript{16,29}, if we have empirically valid shape and scale parameters that define the Pareto distribution, we can generate realizations of neutral allele frequency distributions that are well-specified. Previously, Caravagna et al\textsuperscript{16} fit Pareto distributions (and beta distributions) to thousands of patient tumours extracting both shape and scale parameters. We used these Pareto distribution fits from diploid regions of patient tumours with greater than 50x sequencing coverage to build sampling distributions for the shape and scale parameters. We then used these sampling distributions to generate allele frequencies under a Pareto distribution and, in addition, randomly assigned clonal mutations to each neutral synthetic VAF. In practice, as previously noted\textsuperscript{16}, the scale parameter can be set to the minimum observed frequency as this is the maximum likelihood estimate for the Pareto distribution.

We note that we added additional noise to synthetic neutral distributions to better account for variability observed in empirical data in two ways. Firstly, for any synthetically generated neutral distribution, we randomly trimmed the low frequency neutral tail at a frequency $f(10\text{ - }30\% \text{ VAF})$ with some probability $P_{\text{trim}}$ ($\leq 0.1$). We perform this step as many VAF distributions observed in patient biopsies lack the characteristic neutral tail, even at high sequencing depth\textsuperscript{16}. By randomly trimming neutral synthetic VAF distributions, we tend to more parsimonious explanations of the data, with respect to positive selection, when assessing incomplete and potentially noisy VAF distributions. Secondly, we randomly shifted the heterozygous, diploid clonal peak (that should be centered at 50% VAF) to between 45 and 50% VAF. We perform this random perturbation of the clonal peak to avoid overestimating positive selection when patient samples have incorrect tumour purity estimates that may have led to spurious elevation in the number of subclonal mutations.

Finally, to ensure positively selected and neutrally evolving tumours were not out of distribution with each other given the alternate data generation approaches, we built an aggregate simulation framework that generated neutral and positive synthetic tumours in pairs - assigning the neutral VAF distributions with parameter-matched sequencing noise and equivalent clonal and non-clonal mutations with respect to the paired positive selection simulation.
The synthetic data generation algorithms are outlined in supplementary, and code is available at https://github.com/tomouellette/CanEvolve.jl.

Simulation parameter selection
Each stochastic simulation described above was parametrized by the mutation rate (per genome per division), the probability a mutation was a driver, the mean for the exponential selection coefficient distribution, the number of clonal mutations in the founder cell, the maximum number of driver mutation events, the final tumour population size, the sequencing depth, and the sequencing overdispersion. We chose simulation parameters that were consistent with previous studies⁷,¹³,¹⁶,³⁰ and that captured the expected qualitative and quantitative attributes of VAF distributions observed empirically (Supplementary Table S1). All non-fixed parameters were uniformly random sampled during the development of the synthetic datasets. To improve computational speed and efficiency in our stochastic simulations, we used similar simulation approximations as ref⁷. Namely, a small $N$ population size approximation (where we simulated data using a final population size of $10^3$) and a fixed growth rate (where the birth rate was set to log(2) and the death rate was set to 0). In addition, as we were focused on differentiating between neutral evolution and selection at effective sequencing depths of ~50 - 250x, we constrained our search space to 1 or 2 detectable subclones present between 10 - 40% VAF. We implemented this constraint as (i) it is extremely rare to detect 3 subclones in a one-dimensional VAF distribution as each subclone has to be >5-10% VAF (10-20% cellular fraction) for detection, (ii) most frequency-based methods or studies show limited evidence for detecting >1 subclone at 50 - 250x coverage in a single time point, one-dimensional VAF distribution¹⁶, and (iii) below greater than roughly 10% VAF subclones merge with the neutral tail and above roughly 40% VAF subclones begin to merge with the clonal peak when considering diploid regions.

Synthetic supervised learning
As outlined in the results, synthetic or simulation-based deep learning has been shown to be equivalent to amortized approximate inference under a generative model²⁸. Therefore, by optimizing a neural network using synthetic VAF distributions sampled from a stochastic generative process $p(x,z|\Theta)$ (i.e. the synthetic data generation scheme defined above), we can build fast approximate inference models for evolutionary inference. We describe the synthetic supervised workflow from feature generation to prediction below.

Input representation. For each simulation, we converted mutation frequencies into a VAF distribution (histogram) of length $k$ that had a fixed range between 2% and 50% VAF. To implicitly condition our model on mean sequencing depth (readily available from sequenced tumour biopsies), we only included mutations above a frequency cutoff based on the variance of a binomial sequencing noise model. We note that this conditioning step is arbitrary and simply acts as a way to improve model optimization during training. In general, a simple approach to conditioning a neural network on a measurable variable involvings finding a
reasonable encoding within the feature representation. For example, an alternative approach instead of using a frequency cutoff would be to concatenate the sequencing depth to the input feature vector. Overall, each input feature vector was created by counting mutations into \( k \) bins where each bin had a width \( w \) of \((50 - 2\% \text{ VAF}) / k\) and a lower frequency cutoff defined by \( f_{\text{alt}} + (2\sqrt{f_{\text{alt}}c[1 - f_{\text{alt}}]}) / c \) where \( f_{\text{alt}} \) is the minimum alternative reads to call a mutation divided by mean sequencing depth and \( c \) is mean sequencing depth. For all model development and training, we generated and concatenated two feature vectors with \( k = 64 \) and \( k = 128 \) for each simulation to capture varying levels of information depending on the sparsity of mutations in a given synthetic tumour.

**Model search.** We initially developed neural networks for three single or multi-task inference problems: (i) evolutionary mode (neutral evolution or positive selection) and number of subclones classification \((M_{ms})\), (ii) frequency prediction for a single subclone \((M_{1s})\), and (iii) frequency predictions for two subclones \((M_{2s})\). For each multi-task, we performed a random hyperparameter search using a one dimensional (1D) convolutional neural network (CNN) with task-specific fully connected branches as a base architecture. For the random search, the hyperparameters included the number of convolutional layers (1 - 20) the task-specific branch type (fully connected or global average pooling), the number of feature maps/channels for each convolutional layer (4 - 32), the convolutional kernel width for the left trunk, right trunk, and task-specific branches (1 - 17, odd), the learning rate \((10^{-7} - 10^{-3})\), and the patience for early stopping (3 - 5). To tend toward higher precision and lower recall for predicting selection, we also tuned a penalty term on the positive class in the binary cross entropy loss. Batch size was fixed to 256. Hardswish activations were used at each hidden layer. Dropout, fixed at a probability of 0.5, was added after each layer to allow for downstream application of uncertainty estimation (see Uncertainty Estimation below). We note that we also explored inferring subclone emergence time under a multiplicative fitness model, but could not effectively recover parameters likely due to a complex non-linear relationship between subclonal fitness and emergence time. However, we provide these estimates as an ‘experimental’ output in the TumE python package (links below).

**Model training.** We trained over 150 models for each evolutionary inference task(s) using an Adam optimizer, minimizing the cross-entropy loss for classification tasks \((M_{ms})\) and the L1 loss for regression tasks \((M_{1s} \text{ and } M_{2s})\), on approximately 40 million synthetic tumours simulated with parameters outlined in Supplementary Table S1. For training, each batch consisted of 20,000 unique simulations and training was stopped after 4 epochs or when early stopping, updated after each batch, was activated based on specified patience. To avoid overrepresentation of any subclone frequency during training, we re-balanced positive selection simulations before each batch to have an equal number of subclones at each frequency up to two decimal places (e.g. 0.11 or 11% VAF). For two subclone simulations, we re-balanced simulations based on the distance between subclones \((f_{\text{subclone2}} - f_{\text{subclone1}})\) and only included simulations where the distance
between subclones was >5% VAF. Note that only positive selection simulations were used to train $M_{1s}$ and $M_{2s}$.

**Model selection.** Using an independent test set of one hundred thousand simulations, we then selected the top models across each multi-task for further validation. For $M_{ms}$, we selected models that maximized the mean accuracy across the evolutionary mode, $P(\text{Selection})$, and number of subclones, $P(\text{N subclones})$, classification, and favoured models that assigned a larger penalty term to misprediction of positive selection (i.e. a lower weight to the positive class in the binary cross entropy loss). For the regression models $M_{1s}$ and $M_{2s}$, we selected models that minimized the mean absolute error between the true and predicted subclone frequency on the test set while also ensuring that predictions properly extrapolated across the entire simulated parameter range (e.g. ~10 - 40% VAF for subclone frequencies).

**Uncertainty estimation.** To capture model-based uncertainty in our estimates, we implemented a form of Bayesian approximation for deep learning called Monte Carlo (MC) dropout\(^{26,27}\). Conceptually, MC dropout captures model-based uncertainty by taking advantage of the relationship between model averaging and standard dropout - a network with dropout at every layer encodes $2^n$ possible network configurations. By keeping dropout on at test time, each prediction is a stochastic pass through a set of randomly activated neurons. More specifically and with a slight abuse of notation w.r.t to ref\(^{26}\) ignoring the variational notation, we make estimates of our target variable $y$ (e.g. subclone frequency) by performing $T$ stochastic forward passes through the network and averaging, $E(y)$, the results:

$$E(y) \approx \frac{1}{T} \sum_{t=1}^{T} \hat{y}(x, W_1^t, \ldots, W_L^t)$$

where $\hat{y}$ is the output with respect to the input data $x$ for a neural network with $L$ layers, and $W$ corresponds to a weight matrix for each layer $L$. For every stochastic pass, each $W$ is assigned a randomly sampled vector of Bernoulli random variables such that each individual neuron is inactivated with a probability equal to the dropout rate. Under this framework, MC sampling over $T$ stochastic passes through the network generates an approximate posterior for our target variables with respect to the input data.

**Making predictions.** For differentiating between neutral evolution and positive selection, $P(\text{Selection})$, and predicting the number of subclones, $P(\text{N subclones})$, in both synthetic and real patient tumours, we took a conservative, more parsimonious approach to prediction by considering the variance in the approximate posterior. For $P(\text{Selection})$, we only called positive selection if the lower bound of an 89% equal-tailed interval for the approximate posterior, computed across 50 stochastic passes through $M_{ms}$, was greater than 0.5. If the lower bound was less than 0.5, we called neutrality and zero subclones, independent of the result of $P(\text{N subclones})$. We show the utility of this strategy for mitigating model overconfidence in a synthetic toy example (Supplementary Figure 6). For the regression tasks of predicting subclone frequency
and emergence time, we estimated the true value by performing 50 stochastic passes through the networks and averaging the results, while also providing the complete approximate posterior. We describe additional considerations for making estimates in real patient tumour biopsies below.

All model development and training was done using *pytorch* v1.8.1. We provide a python package, scripts, and all trained neural network models for downloading, use, and modification at [https://github.com/tomouellette/TumE](https://github.com/tomouellette/TumE).

**Model performance in synthetic tumour sequencing datasets**

We simulated or collected 3 different datasets of synthetic tumour sequencing data to study the performance of TumE under changing parameter regimes or changes to model assumptions. The first dataset, generated by our simulation framework described above, consisted of ~2.8 million synthetic tumours simulated across varying sequencing depths and overdispersions (all parameters provided in Supplementary Table S1). Using this dataset, we compared TumE against six frequency-based summary statistics for differentiating between positive selection and neutral evolution. Four of the statistics were cancer evolution statistics developed previously\textsuperscript{12} and provided in the R package *neutralitytestr*. For each sample, the parameters of *neutralitytestr* were set as follows: ploidy = 2, cellularity = 1, read_depth = simulated mean sequencing depth, rho = simulated overdispersion (rho). Two of the statistics were common population genetic statistics, Tajima’s $D$\textsuperscript{20} and Fay and Wu’s $H$\textsuperscript{21}. Only variant allele frequencies and sequencing depth were required for input to compute these statistics. We provide an implementation of Tajima’s $D$ and Fay and Wu’s $H$ for tumour sequencing data in the github repository. We additionally evaluated a mixture model based approach MOBSTER\textsuperscript{16} for subclone detection and frequency quantification. To enable analysis of ~2.8 million synthetic tumours, we ran MOBSTER with the following parameters: $K = 1:3$, samples = 2, init = "peaks", tail = c(TRUE, FALSE), epsilon = 1e-6, maxIter = 100, fit.type = "MM", seed = 12345, model.selection = "reICL", pi_cutoff = 0.02, N_cutoff = 10. We defined the number of subclones that MOBSTER detected as follows. If a tail and 3 beta components were fit then we assigned 2 subclones, if a tail and 2 beta components or if no tail and 3 beta components were fit we assigned 1 subclone, and for all remaining fits we assigned 0 subclones or neutrality.

The second dataset was retrieved from Caravagna et al.\textsuperscript{16} and consists of synthetic tumour sequencing data from 150 tumours sequenced to 120x depth using a beta-binomial sequencing model and grown to a size of $>10^8$ at a birth rate of 1 and death rate of 0.2. The complete description is provided in the supplementary of ref\textsuperscript{16}. We used this dataset to evaluate the small N approximation and to compare TumE to existing mixture model methods. We applied both MOBSTER and a variational Bayesian mixture model sciClone\textsuperscript{24} to this dataset. MOBSTER was run under default package settings without parallel computation and with $K = 1$ to 3 beta components. sciClone was run under default package settings with copyNumberCalls fixed to 2 and maximumClusters fixed to 4. To estimate the number of selected subclones with sciClone, which doesn’t account for neutral evolutionary dynamics, we took the inferred number of
subclones and subtracted 2 (representing the neutral tail and clonal peak). Per-sample runtimes for TumE, MOBSTER, and sciClone were computed on a single machine with 16GB memory and a 2.3GHz quad-core Intel i7 processor.

The third dataset was used to evaluate variable birth and death rates on TumE estimates for predicting positive selection, determining the number of subclones, and estimating subclone frequency. The dataset consisted of ~6 million synthetic tumours, generated by our simulation framework described above, grown at variable birth and death rate combinations. Mutation rate and mean sequencing depth were both fixed to 100. Other parameters were uniformly sampled and all parameters evaluated are outlined in Supplementary Table S1.

All statistical analyses comparing methods, including computation of AUROC, correlation coefficients, and performance metrics such as precision and recall, were performed in R v4.0.3.

**Evolutionary parameter estimates in bulk sequenced single tumour biopsies**

In this study, we used diploid regions of patient tumours for evolutionary inference as we did not have access to accurate phased mutation information for copy number correction of VAFs at non-diploid sites. However, in the absence of complete whole-genome duplication, mutated diploid regions should be sufficient to capture ongoing selection, due to selective sweeps from genome-wide linkage, if a sufficient number of neutral passengers mutations are accumulated during cell division over time\(^7,16\).

In addition to only analyzing diploid regions, we only accepted tumour samples that had at least a 50x mean effective coverage (mean sequencing depth times purity). We set this cutoff as previous studies have shown that tumour genomes sequenced below 50-70x coverage are exceedingly noisy and have insufficient limits of detection relative to low-frequency mutations for proper evolutionary inference\(^16,31\).

Relative to our simulations, VAFs in bulk sequenced single tumour biopsies may also be confounded by impurity, where purity (cellularity) is defined as the percentage of cells in the biopsy that are of malignant or tumour origin. In general, low tumour purity can lead to spurious identification of subclones as it results in lower observed VAFs relative to the true underlying population VAFs. To ensure our inferences weren’t biased by impurity, we corrected all VAFs using corresponding tumour purity estimates collected from the study of origin where \(\text{VAF}_{\text{corrected}} = \frac{\text{VAF}_{\text{observed}}}{\text{purity}}\).

We also note that some purity estimates may be incorrect - in these cases updating the VAFs with incorrect purity estimates can lead to a heterozygous clonal cluster (that should be centered at approximately 50% VAF) in the subclonal frequency range (~10% - 40% VAF). To ensure clonal clusters were properly centered
at 50% VAF following purity correction, we performed additional adjustments to each patients’ VAF distribution using the following heuristic. We first computed the density for each VAF distribution and then identified all the locations where the second derivative of the density was zero i.e. peak finding. If the closest peak to 50% VAF (the theoretical diploid clonal cluster) was above 35% VAF, we considered it a misrepresented clonal peak. We made this assumption as analyses in pan-cancer datasets suggest that all tumours are initiated in somatic cells already carrying mutations\textsuperscript{10,11}. We then fit a Gaussian distribution to the identified clonal cluster of each patient VAF distribution and adjusted each VAF by multiplying by 0.5 divided by the mean of the fit. Although a Beta distribution is generally used for fitting clonal clusters in cancer genomics, a Gaussian is a reasonable approximation for adjusting VAFs based on incorrect purity estimates as it provides accurate estimates of the cluster mean, and has been used in previous subclonal clustering methods\textsuperscript{41}. Plots of patient VAF distributions before and after application of this correction are provided in (Supplementary Figures 17 & 18).

Heuristic clustering using the estimated subclone frequencies was performed either using the expected variance under a binomial sequencing noise model or, alternatively, using the subclone frequency estimates to initialize the means of a gaussian mixture model. Clustering under the binomial framework was performed as follows. Given an estimated subclone frequency $q$, all mutations within the frequency range of $q \pm (\epsilon \sqrt{qc(1-q)}) / c$ were assigned to the subclone, where $\epsilon$ scales the cluster width and $c$ is the mean sequencing depth across the tumour genome or exome. We fixed $\epsilon$ to 2 in this study.

**Transfer learning for inference in alternative synthetic data regimes**

Given a pre-trained network with weights optimized to a source domain $S$, composed of input space $X_S$, output space $Y_S$, transfer learning attempts to use pre-training to improve the performance on another target domain $T$ composed of $X_T$ and $Y_T$. We employ a variant of transfer learning called open set domain adaptation\textsuperscript{36} to take advantage of our pre-trained models for additional inference tasks. In this case, the input spaces remain constant ($X_S = X_T$, VAF distribution) but the inferred tasks are allowed to differ. Open set indicates that some tasks may overlap with the output space of both the source and target domains.

To provide a concrete use case for transfer learning in synthetic supervised learning, we aimed to infer additional evolutionary parameters such as subclone fitness, subclone emergence time, mutation rate, and subclone cellular fraction (subclone frequency * 2) using synthetic tumour sequencing data generated by an alternative cancer simulation framework TEMULATOR\textsuperscript{37}. TEMULATOR differs from our synthetic data generation scheme, which was built around a multiplicative fitness driver model, as subclones are deterministically initiated at user specified emergence times and fitnesses. To facilitate transfer between previous and new tasks, we performed architecture renovation on the pre-trained neural networks, retaining all convolutional layers while replacing existing task-specific fully-connected branches with new task-
specific fully connected branches (4 in total). To maximize the amount of information transferred to new
tasks, we combined the convolutional layers from both the $M_{ms}$ and $M_{ts}$ models described above.

We then simulated 500,000 synthetic tumours at a birth rate of 1, death rate of 0.2, and final population
size of $10^4$ (additional parameters such as mutation rate were uniformly sampled and are outlined in
Supplementary Table S3). To facilitate efficient simulation, we first fit a noisy Gaussian process (GP)
regression to the viable emergence time and fitness parameter combinations (that generated detectable
subclones between ~10-40% VAF) and used the GP to sample viable emergence times given a uniformly
sampled fitness. We made the GP noisy to facilitate parameter combinations that resulted in subclones
across the entire frequency range. The GP was fit using three kernels (RBF with length scale 100, dot-
product, and white noise) and an alpha of $10^{-6}$ in the python package *scikit-learn* v1.0. Next, we used the
simulations to re-optimize the pre-trained model weights, using an Adam optimizer to minimize the L1 loss
(mean absolute error) for predicting new evolutionary inference tasks. To ensure fair benchmarking
between networks with and without pre-trained weights, we performed a random hyperparameter search
with ~150 pre-trained and ~150 non-pretrained models, tuning the learning rate and number of fully
connected layers in the task-specific branches. Additional synthetic data used for evaluating performance
was generated under similar parameter settings. We corrected modest yet systematic overestimates in
mutation rate (~50% mean percentage error) in the final transferred model by fitting a polynomial (degree
2) ridge regression in *scikit-learn* v1.0 to the predicted mutation rates. The mutation rate adjustment was
performed using VAF distributions from 1000 synthetic tumours. We validated the correction on an
additional 100,000 synthetic tumours. All TEMULATOR synthetic tumours were generated using parameter
settings in Supplementary Table S3.

Predictions in empirical samples were performed by taking 500 Monte Carlo dropout samples and
averaging the results. Dropout was only activated at test time on the new task-specific branches. Per-base
mutation rate estimates in whole-exome sequenced MMR-GE samples were rescaled based on the 60MB
Agilent SureSelectXT Human All Exon v6 kit used in the original study. Because subclone fitness and
emergence time is impacted by final tumour size, we rescaled our estimates to a realistic tumour size of
$10^{10}$ cells, similar to ref. Previous work has shown that given subclone frequency $f_{sub}$ and an estimated
final population size $N_{end}$, the age of a tumour at time of biopsy can be estimated by $t_{end} = \log 2([1 - f_{sub}] \cdot
N_{end})$. Therefore, given that the relationship between emergence time in tumour doublings and log
population size is linear, we can generate a rescale fitness estimate $w_R$ as follows.

$$w_R = I + (w - I) \cdot \frac{t_{end} - t_s}{t_{endR} - t_{sR}}$$
where \( w \) equals subclone fitness, \( t_{\text{end}} \) indicates time at tumour biopsy or final population size in tumour doublings, and \( t_s \) indicates the time of subclone emergence in tumour doublings. \( R \) subscript indicates values rescaled to population size of \( 10^{10} \). The parameters \( f_{\text{sub}}, w, \) and \( t_s \) are all inferred. We approximate the rescaled subclone emergence time \( t_{sr} \) as \( t_s \ast \log(N_{\text{end}})/\log(N_{\text{sr}}) \).

**Data availability**

All TumE predictions in synthetic and empirical datasets, intermediate processing data, data used for generating figures, and fully trained deep learning models can be found at https://doi.org/10.5281/zenodo.5575877 repository. Whole-genome sequenced AML samples were retrieved from Griffith et al\(^{31}\). Multi-region whole-exome sequenced mismatch deficient repair gastro-esophageal samples were retrieved from von Loga et al\(^{33}\). The remaining whole-genome sequenced samples were retrieved from PCAWG\(^{11}\). We provide hosting of the electronic supplementary at https://tomouellette.gitlab.io/ouellette_awadalla_2021/.

**Code availability**

Scripts for generating figures and analyses can be found at https://doi.org/10.5281/zenodo.5575877. Code for generating synthetic tumour sequencing data can be found at https://github.com/tomouellette/CanEvolve.jl. Code for performing inference with TumE can be found at https://github.com/tomouellette/TumE.

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