Algorithms to Model Single Gene, Single Chromosome, and Whole Genome Copy Number Changes Jointly in Tumor Phylogenetics

Salim Akhter Chowdhury¹², Stanley E. Shackney³, Kerstin Heselmeyer-Haddad⁴, Thomas Ried⁴, Alejandro A. Schäffer⁵, Russell Schwartz²⁶*

¹ Joint Carnegie Mellon/University of Pittsburgh Ph.D. Program in Computational Biology, Carnegie Mellon University, Pittsburgh, Pennsylvania, United States of America, ² Lane Center for Computational Biology, Carnegie Mellon University, Pittsburgh, Pennsylvania, United States of America, ³ Intelligent Oncotherapeutics, Pittsburgh, Pennsylvania, United States of America, ⁴ Genetics Branch, Center for Cancer Research, NCI, NIH, Bethesda, Maryland, United States of America, ⁵ Computational Biology Branch, NCBI, NIH, Bethesda, Maryland, United States of America, ⁶ Department of Biological Sciences, Carnegie Mellon University, Pittsburgh, Pennsylvania, United States of America

Abstract
We present methods to construct phylogenetic models of tumor progression at the cellular level that include copy number changes at the scale of single genes, entire chromosomes, and the whole genome. The methods are designed for data collected by fluorescence in situ hybridization (FISH), an experimental technique especially well suited to characterizing intratumor heterogeneity using counts of probes to genetic regions frequently gained or lost in tumor development. Here, we develop new provably optimal methods for computing an edit distance between the copy number states of two cells given evolution by copy number changes of single probes, all probes on a chromosome, or all probes in the genome. We then apply this theory to develop a practical heuristic algorithm, implemented in publicly available software, for inferring tumor phylogenies on data from potentially hundreds of single cells by this evolutionary model. We demonstrate and validate the methods on simulated data and published FISH data from cervical cancers and breast cancers. Our computational experiments show that the new model and algorithm lead to more parsimonious trees than prior methods for single-tumor phylogenetics and to improved performance on various classification tasks, such as distinguishing primary tumors from metastases obtained from the same patient population.

This is a PLOS Computational Biology Methods article.

Introduction
In this paper, we develop new methods to advance the theory of phylogenetic inference for reconstructing evolutionary histories of cell populations in solid tumors. The work is specifically designed for use in tracking tumor evolution by gain and loss of genomic regions as assessed by multicolor fluorescence in situ hybridization (FISH), which measures the copy numbers of targeted genes and chromosomes in potentially hundreds of individual cells of a tumor. This technology was the basis of the earliest methods for phylogenetic reconstruction of single tumors [1,2], FISH remains uniquely valuable for such studies because the large number of cells that FISH can profile makes it possible to collect data on enough tumors in enough detail to build cell-by-cell phylogenies for populations of tumors and begin to study the common features of these phylogenies. In the present work, we specifically extend our previously developed inference algorithms to encompass a more complicated but realistic model of evolution of FISH probe counts, accounting for gain and loss of genetic material at the level of single gene probes, multiple probes on a single chromosome, or a probe set distributed across the whole genome. We demonstrate the value of these algorithmic improvements to more accurate phylogenetic inference and improved effectiveness of the resulting phylogenies in downstream prediction tasks.

The present work adds to the growing list of phylogenetic methods in cancer modeling, which were reviewed through 2008 in [3]. These include methods for analyzing comparative genomic hybridization (CGH) or other genetic gain/loss data in a single tumor type [4–11], for defining the cell type lineage of single tumors [12,13], for organizing a taxonomy of tumor types [14], for reconstructing a partial order of genetic changes in multiple samples from one patient [15], and for reconstructing progression from cell types inferred from bulk genomic assays [16]. Recent high-throughput sequencing studies have also used ad hoc phylogenetic methods to infer putative tumor progression scenarios, e.g., [17–20]. Like many of these methods, the present work is
algorithmic [e.g., neighbor-joining [45]] that are not designed to model profiles of copy number changes. One would expect from evolution by chromosome abnormalities that largely drive tumor evolution. Until recently, algorithms designed specifically for inferring phylogenies of single tumors from FISH data have been limited to just a few probes per cell and lacked robust, publicly available software implementations [1,2,34]. In prior work [46], we developed algorithms to find copy-number phylogenies for in principle arbitrary numbers of probes and cells. That work, however, was itself limited to a simple model in which tumor cells evolve by events of gain or loss of a single copy number of a single probe at each mutation step. In real tumors, gene copy numbers can change due to a variety of mechanisms, including:

1. Single gene duplication/loss events (SD), in which one copy of a genetic region covered by a single probe is gained or lost.
2. Chromosome duplication/loss events (CD), in which entire chromosomes are unequally distributed among daughter cells during mitosis along with potentially several probes.
3. Whole genome duplication events (GD), in which a cell fails to divide during mitosis leading to doubling of all genetic material and all probe counts.

These events are illustrated schematically in Figure 1. While more complex probabilistic models of tumor evolution have been developed for inference of small phylogenies, with approximately ten taxa per tumor corresponding to distinct biopsies (e.g., [47]), the class of inference algorithms such models require would not be expected to scale to phylogenies of hundreds of single cells per tumor such as those examined in the present work. The work presented here seeks to fill this need for scalable phylogenetic algorithms capable of fitting more realistic models of tumor-like evolution to data sets of hundreds of single cells per tumor. We improve on our prior work for inferring tumor evolutionary models considering only SD events [46] to now include CD and GD events, which are also frequently observed in tumor progression. We specifically focus on the problem of accurately inferring evolutionary distances between distinct cells in terms of maximum parsimony combinations of SD, CD, and GD events. The major contributions of the work are:

1. Algorithms to compute minimum evolutionary distances $D$ between pairs of cell states in terms of SD and CD events and in terms of SD, CD, and GD events;
2. A heuristic Steiner tree method based on the median-joining method [48] and our prior work on SD-only inference [46];
3. Software implementation of the new methods to compute $D$ and use of those methods to construct tumor progression trees;
4. Evaluation of the new methods on simulated data, which shows that they do better than the SD-only approach at recovering simulated tree topologies;
5. Application of the methods to published data on cervical cancer (CC, [49]) and breast cancer (BC, [36]);
6. Demonstration of improved ability to classify tumor types from phylogenetic features using a strategy in the spirit of the genomic progression scores (GPS) of Rahnenführer et al. [50].

The new methods are implemented in version 2 of our software FISHtrees (ftp://ftp.ncbi.nlm.nih.gov/pub/FISHtrees). The work addresses a critical need in modern cancer research for algorithms capable of inferring evolutionary trajectories of hundreds of single cells per tumor under plausible models of evolution including both

---

**Author Summary**

Cancer is an evolutionary system whose growth and development is attributed to aberrations in well-known genes and to cancer-type specific genomic imbalances. Here, we present methods for reconstructing the evolution of individual tumors based on cell-to-cell variations between copy numbers of targeted regions of the genome. The methods are designed to work with fluorescence in situ hybridization (FISH), a technique that allows one to profile copy number changes in potentially thousands of single cells per study. Our work advances the prior art by developing theory and practical algorithms for building evolutionary trees of single tumors that can model gain or loss of genetic regions at the scale of single genes, whole chromosomes, or the entire genome, all common events in tumor evolution. We apply these methods on simulated and real tumor data to demonstrate substantial improvements in tree-building accuracy and in our ability to accurately classify tumors from their inferred evolutionary models. The newly developed algorithms have been released through our publicly available software, FISHtrees.

aimed at building tree models that provide a proposed partial order on the observed cell states, a strategy motivated originally by the work of Fearon and Vogelstein, proposing a linear order for four types of events in colorectal cancer and associating each event with a tumor stage [21]. Other ordering methods have been proposed, mostly for CGH or breakpoint data [15,22–28] and, with a tumor stage [21]. Other ordering methods have been proposed, mostly for CGH or breakpoint data [15,22–28] and, with a tumor stage [21]. Other ordering methods have been proposed, mostly for CGH or breakpoint data [15,22–28] and, with a tumor stage [21].

Furthermore, a simulation study has suggested that methods based on average copy number data perform poorly when there is substantial intratumor heterogeneity [44]. Such findings suggest a need for improved methods for organizing the dozens or hundreds of observed cell states in single tumors to infer the evolutionary processes that produced them.

Despite extensive work on tumor phylogenetics, however, the study of algorithms for reconstructing tumor evolution from large numbers of single cells has lagged far behind advances in data generation. The standard in practice for single-cell tumor phylogenetics remains the use of simple generic phylogeny...
gene-specific and chromosome abnormalities that are central drivers of true tumor evolution.

Results

We used data collected from cervical cancer (CC) [49] and breast cancer (BC) [36] patients to evaluate our methods. Figure 2(A) shows a tumor progression tree inferred from one of the cervical cancer samples. For comparison, Figure 2(B) shows a progression tree inferred on the same sample using our prior SD model [46]. Visual inspection shows that large regions of the two trees are identical but that allowing CD and GD events leads to some rearrangement and a reduction in tree depth and overall size. Next we evaluate the changes induced by adding SD, CD and GD events, using simulated data to show effectiveness of the methods in finding more parsimonious solutions to the broader model and using the real CC and BC data to show the biological relevance of the improvements. We further show that our algorithms infer trees with higher accuracy than the prevailing alternative algorithms for single-tumor phylogenetic inference. Finally, we perform statistical experiments to evaluate the effects of tumor sample size on the performance of our tree building algorithm.

Simulation experiments

To measure accuracy of the methods for FISH datasets with a known ground truth, we generated a dataset of 100 trees with six probes, two of which were treated as being on the same chromosome. Each tree was generated by starting from a diploid root node and executing a branching process in which each node was recursively assigned a number of children drawn from a geometrically distributed random variable with mean 0.50. Each child was distinguished from its parent by selecting an SD, CD, or GD event with probability 0.1167 for each of the six possible SD events, 0.18 of a CD event, and 0.12 of a GD event. This process terminated when all leaf nodes had been assigned zero children by the sampling. We then generated simulated FISH data for each tree by uniformly sampling 300 cells from the nodes in this topology. The simulated data corresponds to counts of probes for each sampled cell in the tree. We applied Algorithm 3 (see Methods) to find a minimum-cost tree for each of four event models: (i) SD only, (ii) SD and CD, (iii) SD and GD, and (iv) SD, CD and GD.

We quantified the accuracy of tree inference by comparing each simulated true tree to its corresponding inferred tree derived from the sampled cells. This assessment was performed at the level of accuracy of tree edges by the following procedure:

1. We pruned the real tree so as to remove any subtree for which no cell in the tree was sampled. This step was intended to avoid penalizing for “impossible” inferences of subtrees unsupported by any data.
2. We computed a maximum matching of edges between the real subtree and the inferred tree, with each pair of edges weighted by the maximum number of nodes in agreement between the corresponding parts of the bipartitions that the two edges define [46,51]. We used the Hungarian algorithm [52] for computing the maximum matching (applying the function “Hungarian” by Alexander Melin from the Matlab Central File Exchange).
3. We calculated a reconstruction error $R$ of the inferred tree using the following formula:

$$R = \left(1 - \frac{W}{|T| \times (|P_r| + |P_i|) - W}\right) \times 100$$

where $W$ is the weight of the maximum matching, $T$ is set of taxa in common between the real and inferred trees, and $P_r$ and $P_i$ represent the sets of nontrivial bipartitions in the real and inferred trees, respectively.

Intuitively, this formula measures the fractional agreement between bipartitions of the trees relative to the total number of bipartitions. We use a matching-based formula, rather than the more familiar Robinson-Foulds metric [33], both because of its greater sensitivity to small changes in trees and because the Robinson-Foulds measure is not defined for trees with different node sets. We also note that we use a different normalization factor than in our prior work [46], normalizing essentially by the total number of edges between the two trees, to control properly for the fact that different inference methods may infer different numbers of tree edges. The reconstruction error $R$
ranges in value from 0, if the real and inferred trees are isomorphic, to an upper bound of 100 in the limit of complete disagreement.

To illustrate the meanings of the terms of the equation for $R$, we present a simple example using a hypothetical ground truth and an inferred tree presented in Figure 3(A) and Figure 3(B), respectively. The set of nontrivial bipartitions in the ground truth are

$$\{\{1,3,6,7,8,9\}, \{2,4,5\}, \{\{3,6,7,8,9\}, \{1,2,4,5\}\}$$
and the nontrivial bipartitions in the inferred tree are

\[
\{\{1,3,6,7\}, \{2,8,9\}\}, \{\{1,2,8,9\}, \{3,6,7\}\}
\]

If we apply the matching algorithm on these two sets of bipartitions, the first and second bipartitions in the ground truth tree are matched with the first and second bipartitions in the inferred tree, respectively. The weight \(W\) of the matching is 10. The number of common taxa between these two datasets is \(|T| = 7\). The total number of nontrivial bipartitions in the real and inferred trees are \(|P_f| = 3\) and \(|P_f| = 2\). Plugging these values into the equation for \(R\), we calculate \(R = 60\%\).

A comparison of the four models is presented in Figure 4. The SD model showed 17.43% reconstruction error with standard deviation (s.d.) of 5.1% across the 100 trees. The SD+CD model yielded 15.91% error with s.d. 4.59%. SD+GD yielded 12.01% error with s.d. 6.4%. The full SD+CD+GD model yielded 10.84% error with s.d. 3.88%. Collectively, the results suggest that one can reconstruct reasonably accurate trees even from the SD-only model, despite the fact that the trees were generated from a model of all three event types, although accuracy improves with each event type added. Accounting for GD events made a larger impact on reconstruction error with s.d. of all three event types, although accuracy improves with each event type added. When the probability of SD events is high (as in combination 3), the SD model results in highly accurate trees (84% with s.d. 0.1%). Account-
larger improvement in the reconstruction error in comparison to the SD+CD models, even when the CD events are very frequent (as in combinations 2 and 4). Finally, accounting for GD events in combination with SD and CD events results in the largest improvements when the probability ratio of GD events to SD+CD events is highest, as can be seen from comparison of parameter sets 1 and 2.

Next, we performed simulation tests to evaluate the effects of non-uniform distributions of cells across different levels of the trees on the performance of our tree inference method. In our initial simulation experiments described above, we assumed that observed cells were sampled uniformly across clones. In real tumors, the distribution of cells would not typically be uniform due to differences in age and fitness of clones. In order to test robustness of our method to non-uniformity of clone frequencies, we sampled the cells following a non-uniform model in which the sampling frequency of a clone varies geometrically with its depth in the tree with a parameter \( \gamma \). We used values of 1.1 and 1.3 for \( \gamma \) in our experiments. When \( \gamma = 1.1 \), 25% of the total cells are located in the first three levels of the trees, while for \( \gamma = 1.3 \), this fraction is 55%. We generated 100 trees in each case with probabilities of SD, CD and GD events fixed at 0.1167, 0.09, and 0.12. We again used SD, SD+CD, SD+GD and SD+CD+GD models to infer the tumor progression trees. We present the results from this experiment in Table 2, where we also show the results from the uniform sampling of the cells. Additionally, we report the results on the trees inferred using NJ and MP for these three different cell distributions. From the table, we can see that the reconstruction error increases with increasing \( \gamma \) for all methods. The SD+CD+GD model, however, shows the best performance among all the models for all three values of \( \gamma \) and the least loss of performance with increasing \( \gamma \).

Finally, we performed simulation experiments to understand the effects of varying the numbers of chromosomes with multiple probes. We created a simulated dataset of 100 trees with eight probes where two pairs of probes each reside on two different chromosomes and the remaining four probes reside on four separate chromosomes. The probabilities of each of the SD, CD and GD events were fixed at 0.1167, 0.09, and 0.12, respectively. We report the results from this experiment in Table 3, which compares the results from this experiment with our earlier result using only a single chromosome with two probes and four other probes located on separate chromosomes. The table shows that inclusion of the extra possible CD event results in higher accuracy for all the models except for the SD only model. The performance drop in the SD model is expected, as it would require more SD events to explain a greater number of missed CD events. The highest gain in performance is observed for SD+CD+GD model. These results show that our algorithm will tend to yield comparatively more advantage over the earlier work with more complicated scenarios of sharing probes across chromosomes, suggesting its utility will increase as improvements in technology allow for larger probe sets.

### Application to real cervical and breast cancer data

We applied the algorithm to two sets of real data:

- A set of CC [49] FISH data consisting of 47 samples organized into 16 primary samples of metastic patients, 16 paired metastasis samples from the same patients, and 15 primary samples from patients who did not progress to metastasis. Each sample consisted of 223–250 cells profiled on four FISH probes: LAMP3 (Entrez Gene Id 27074) [55], PROX1 (5629) [56], PRKAA1 (5562) [57] and CCND1 (595) [58]. All of these four genes are oncogenes, which typically show copy number gains in tumor cells. Each of the genes belongs to a distinct chromosome.

- A set of BC [36] FISH data consisting of 13 paired (from the same patient) ductal carcinoma in situ (DCIS) and invasive ductal breast carcinoma (IDC) samples with 76–220 cells per sample profiled on eight FISH probes: COX-2 (5743) [59], MYC (4609) [60], CCND1 [58], HER-2 (2064) [61], ZNF217 (7764) [62], DBC2 (23221) [63], CDH1 (999) [64] and TP53 (7157) [65]. The first five genes in this list are oncogenes and the last three genes are tumor suppressors. In tumor cells, tumor suppressors are typically associated with loss in copy numbers.

Among the eight genes in the BC dataset, DBC2 and MYC reside on chromosome 8 and HER-2 and TP53 reside on chromosome 17. The other four genes belong to distinct chromosomes. The oncogene Cyclin D1 (CCND1), which plays a role in many solid tumor types, is in both the BC and CC datasets. However, in some other tumor types, such as oral cancer, CCND1 is part of a larger region with recurrent copy number gains on chromosome 11 and other nearby genes have also been suggested to play a role in oncogenesis [66].

We evaluated the SD+CD+GD method by its effectiveness in reducing the parsimony score (total number of mutation events) of the resulting trees relative to the prior SD-only model. With the primary CC samples, the SD+CD+GD method found a lower-cost

| Probabilities of (SD,CD,GD) Events | SD   | SD+CD | SD+GD | SD+CD+GD |
|-----------------------------------|------|-------|-------|----------|
| (0.125,0.05,0.2)                  | 17.97(4.49) | 16.89(4.32) | 9.85(3.51) | 9.25(4.18) |
| (0.1,0.2,0.2)                     | 25.58(4.50) | 21.82(3.98) | 13.81(3.62) | 10.96(3.99) |
| (0.15,0.07,0.03)                  | 16.02(4.15) | 14.96(4.16) | 11.92(4.29) | 11.71(4.77) |
| (0.1,0.3,0.1)                     | 23.13(4.37) | 20.02(4.50) | 15.43(4.60) | 13.42(4.64) |
| (0.1166,0.18,0.12)               | 17.43(5.10) | 15.91(4.59) | 12.01(6.40) | 10.84(3.88) |

Mean percentage reconstruction error on 100 simulated samples are shown for four tree-building models considering (i) SD, (ii) SD+CD, (iii) SD+GD and (iv) SD+CD+GD across five different combinations of SD, CD, and GD probabilities.

doi:10.1371/journal.pcbi.1003740.10001
tree in 21 of 31 cases, a tree of equal weight in 4 cases, and a higher-cost tree in 6 cases. In each case of increased weight, the increase was by 1 and appears to result from the subtree regrafting heuristic used in handling GD events (see Methods). These results suggest that the heuristic tree search may more often yield a suboptimal result for the SD+CD+GD model than it does for the SD-only model. The benefit of the more realistic model, however, outweighs the cost of this suboptimality in a large majority of instances. For trees derived from metastatic samples, 12 of 16 trees had lower weight for the full SD+CD+GD model and the remainder all had equal weight for the two models. Metastatic data sets tend to have fewer distinct cell types than do primary trees and thus may represent an easier optimization challenge. For data sets tend to have fewer distinct cell types than do primary CD samples, trees and thus may represent an easier optimization challenge. For trees derived from metastatic samples, 12 of 16 trees had lower weight for the full SD+CD+GD model and the remainder all had equal weight for the two models. Metastatic data sets tend to have fewer distinct cell types than do primary trees and thus may represent an easier optimization challenge.

Table 2. Comparison of mean percentage reconstruction error (with standard deviation) of different phylogeny models on simulated data for different sampling distributions of the cells.

| Distribution | SD | SD+CD | SD+GD | SD+CD+GD | NJ | MP |
|-------------|----|-------|-------|----------|----|----|
| Uniform     | 17.43(5.10) | 15.91(4.59) | 12.01(6.40) | 10.84(3.88) | 43.23(4.24) | 45.21(3.86) |
| Skewed (γ = 1.1) | 22.74(4.49) | 19.09(4.47) | 14.75(4.64) | 11.92(4.64) | 47.00(3.76) | 47.38(3.72) |
| Skewed (γ = 1.3) | 29.93(7.37) | 26.35(5.66) | 18.89(7.24) | 15.36(6.78) | 50.63(5.89) | 50.32(5.74) |

Mean percentage reconstruction error on 100 simulated samples are shown for six tree-building models considering (i) SD, (ii) SD+CD, (iii) SD+GD, (iv) SD+CD+GD (v) NJ and (vi) MP when the sampling distribution of cells is varied. doi:10.1371/journal.pcbi.1003740.t002

As a final step, we repeated an approach developed in our prior work [46] to both validate the biological relevance of the trees and develop a practical application of them by treating the trees as sources of features for classification tasks applied to the CC data. For this purpose, we developed several sets of quantitative features based on inferred GD events, and the uncertainty among whether a GD event occurred prior to or after the emergence of the Steiner node may explain why the per-edge agreement is higher. Nonetheless, the data support the conclusion that inferred GD events are correct in a majority of cases.
trees. We compared these features to four features derived directly from FISH probe counts without reference to the trees:

1. Mean gain and loss of individual genes.
2. Maximum copy number of individual genes.
3. An information theoretic measure, Shannon index [67]. For each gene, each combination of gene copy number and cellular ploidy represents a species. If we denote the frequency of species $i$ among all tumors by $p_i$, then Shannon index is given by the formula $H = - \sum p_i \log_2 (p_i)$.
4. Simpson’s index [67], which is defined as $\sum p_i^2$.

We used each feature set as input to the Matlab support vector machine (SVM) classifier with a quadratic kernel using 500 rounds of bootstrap replicates per test with leave-one-out cross-validation to compute mean and standard deviation of accuracy. We used

Figure 5. Parsimony score comparison on the CC samples. Comparison of (A) Primary and (B) Metastatic CC tumor progression tree weights built considering only SD and combined SD, CD and GD models. “Total Cell Type” refers to the total number of unique probe copy number configurations in the dataset, providing a lower bound on the minimum possible parsimony score for a given data set.

doi:10.1371/journal.pcbi.1003740.g005
Matlab functions “svmtrain” and “svmclassify” for training and testing of the SVM classifier.

We then applied these methods for three classification tasks: (i) distinguishing primary samples that progressed to metastasis from their paired metastatic samples, (ii) distinguishing all primary samples from all metastatic samples, and (iii) distinguishing primary samples that metastasized from primary samples that did not metastasize. The first two tasks are relevant to identifying features that help us understand the differences in evolutionary mechanisms of primary and metastatic samples. The third is intended to model an important practical problem in cancer treatment: determining whether a given primary tumor will metastasize.

Figure 8 shows results on each task. For task (i), allowing SD+CD+GD events increased accuracy relative to SD trees from 64.31% to 80.77% for edge counts and from 81.91% to 84.63% for tree level cell count. The SD+CD+GD tree level cell count was the most effective of all features, tree-based or not. For task (ii), we similarly saw a substantial improvement in prediction accuracy for SD+CD+GD trees relative to SD trees. Classification accuracy improved from 68.87% to 84.06% for edge count features and from 82.26% to 87.79% for tree level features. In this case, both SD+CD+GD tree feature sets outperformed all other features sets, tree-based or otherwise. These results provide an indirect validation that using a more general tree model gets closer to the biological ground truth. For task (iii), we saw no improvement, with identical results for SD and SD+CD+GD trees for either feature set. All tree-based feature sets significantly outperformed all non-tree-based feature sets for this task. We conclude that the more realistic evolutionary models appear not to reveal any more information to the classifiers for predicting which primary samples will go on to metastasize than the SD trees, which were already quite effective for that task.

Dependence on data size
A key advantage of FISH for profiling tumor heterogeneity is that it makes it cost-effective to profile much larger numbers of cells than alternatives such as single-cell sequencing. To assess the practical importance of this advantage, we asked two related questions: (1) how many cells do we need per tumor to accurately reconstruct single-cell phylogenies and (2) how many tumors do we need to examine to identify reproducible, statistically significant features across trees.

We first assessed the number of cells needed per tumor by using our first simulated dataset of 100 trees described above with subsamples of varying numbers of cells per tumor, measuring reconstruction error of our SD+CD+GD algorithm with the weighted matching algorithm. The mean reconstruction errors calculated across 100 cases for subsamples of 20, 50, 100, 150 and 200 cells were 33.66% (s.d. 14.40%), 20.43% (7.97%), 15.28% (6.38%), 11.79% (4.03%), and 11.70% (4.4%) respectively. We can thus conclude that accuracy improves noticeably with increasing numbers of cells to at least 100 cells per tumor before plateauing at approximately 10% error.

We next assessed numbers of tumors needed to identify meaningful statistically significant properties of tumor classes by analysis of the 32 CC paired and primary samples. We randomly subsampled from among the 32 pairs and, for each subsample, calculated the following three tree statistics on progression trees inferred from our SD+CD+GD algorithm:

1. Shannon index based on distribution of cells across different tree levels.

Figure 6. Parsimony score comparison on the BC samples. Comparison of DCIS (id 1–13) and IDC (id 14–26) BC tumor progression tree weights built considering only SD and combined SD, CD and GD models. “Cell Types” refers to the total number of unique probe copy number configurations in the dataset, providing a lower bound on the minimum possible parsimony score for a given data set.

doi:10.1371/journal.pcbi.1003740.g006
2. Weighted mean depth of the trees.
3. Sum of differences of fractional gain and loss of each gene across the tree edges.

We then compared distributions of each statistic on primary vs. metastatic trees by a Wilcoxon signed rank test. As the samples were selected randomly, no ordering among the samples was considered. Figure 9 shows the 1-sided p-values of the three statistical tests when the number of randomly selected samples are increased from 5 to 32. The figure shows that ability to distinguish the two tumor subsets improves with increasing number of tumors. While the threshold for significance varies by statistic, each reaches weak significance ($p < 0.05$) between 10 and 24 tumors. We can thus conclude that finding reproducible features distinguishing the tree types requires on the order of tens of tumors, at least for the candidate probe sets examined here.

Taken together, these two results demonstrate that building accurate trees on a large enough scale to distinguish meaningfully primary from metastatic trees requires data sets with roughly the order of thousands of single cells (hundreds of cells per tumor for tens of tumors), a scale of data that has so far been achieved only by FISH studies of tumor heterogeneity. We note, however, that one would expect these numbers to vary depending on the degree of tumor heterogeneity, the classes of trees one wishes to distinguish, and the specific markers examined.

**Discussion**

This paper has presented novel theory and algorithms for reconstructing evolutionary trajectories of gene copy numbers in solid tumors in terms of a model of tumor evolution incorporating changes at the scale of single gene probes, full chromosomes, or all probes in the genome. We have derived algorithms to reconstruct maximum parsimony sequences of events, and thus estimates of evolutionary distance, between pairs of cells assayed by FISH probes. We have further incorporated these inferences into a method for building phylogenies of hundreds of cells in single tumors. These methods have been added to FISHtrees [46], our software for inferring tumor phylogenies from single-cell copy number data. Experimental results on simulated data confirm the ability of the new methods to improve phylogenetic inference accuracy relative to simpler models by adding CD and GD events that model chromosome-scale and whole-genome copy number changes that are frequently observed in tumor evolution. Application to observed human tumor data shows that these extended evolutionary models are able to yield more parsimonious tree reconstructions and that the resulting trees lead
to improved accuracy in prediction tasks related to diagnosis and prognosis.

In future work, we hope to extend the theory developed here to handle more realistic models and more challenging data types. One important direction will be advancing the theory developed here to improve upon the heuristic approximations used in the Steiner tree inference to better approach the goal of finding globally optimal trees for the most computationally challenging FISH data sets. The evolutionary models, likewise, might be further extended to go beyond the three mutational event types considered here to better approximate the numerous distinct mutational mechanisms by which copy number profiles of tumor cells might evolve. The data sets studied here do not include geographical information about locations of individual cells in the tumor, but other data sets for analyzing tumor heterogeneity do include such geographical information [38,68]. We expect it would be interesting to construct phylogenies with distance functions that combine spatial distance in three dimensions with combinatorial distance measures between the cell count patterns, as we have studied here. Further, while FISH for the moment retains a unique advantage in the large number of cells it can profile, one can reasonably anticipate that single-cell sequencing will eventually become practical for comparable cross-tumor studies. There would thus be value in extending the theory developed here to single-cell sequencing data, a goal that would pose substantial algorithmic challenges due to the much larger number and variety of markers it can reveal as well as the more complicated error models it would entail. Finally, we hope to make more use of these single-tumor phylogenetic models in clinically relevant prediction tasks and further explore the biological insights one can gain from more accurate tumor phylogenies.

**Methods**

Our main theoretical result is a method for inferring minimum distances between two states within a copy number phylogeny when duplication/loss of single genes (SD), duplication/loss of all genes on a common chromosome (CD), and duplication of all genes in the full genome (GD) events are possible. We first establish some mathematical results and then develop an algorithm for accurate distance computation. This algorithm then becomes a subroutine in a heuristic Steiner tree algorithm for inferring copy number phylogenies in the presence of SD, CD, and GD events. We introduce some notation required for specifying and proving the theoretical results:

1. $C(g_1, g_2, \ldots, g_d)$: A set of copy numbers of one or more genes $g_1, g_2, \ldots, g_d$, which we call a “configuration”. When $g_1, g_2, \ldots, g_d$ are clear from the context, we use $C$ as shorthand.
2. $L_1(C, C')$: $L_1$ or rectilinear distance between two configurations $C$ and $C'$.
3. $D^{sd}(C, C'), D^{cd}(C, C'), D^{gd}(C, C')$: Distance between two configurations $C$ and $C'$ when considering SD+CD (s,c,h), SD+GD (s,g), or SD+CD+GD (s,c,g) events, respectively.
4. $O(C)$, $O(C')$, $H(C)$: Operations corresponding to single chromosome (CD) events corresponding to either gain $g$, loss $l$, or either (no subscript) of all genes belonging to the same chromosome $c$ from starting configuration $C$, while keeping the copy numbers of genes on other chromosomes unchanged.
5. $O(C')$, $H(C)$: Operations corresponding to doubling ($O$) or halving ($H$) counts of all genes in configuration $C$. In the case of halving, it is assumed that all genes in $C$ have even counts.
6. even, odd configuration: A configuration (copy number profile) $C(g_1, g_2, \ldots, g_d)$ is denoted an even configuration if $g_i \mod 2 = 0$. Otherwise, it is denoted an odd configuration.
7. $G^k(C(g_1, g_2, \ldots, g_d))$: The set of “nearest even” values for each $g_i$ in $C$, i.e., if $C(g_1, g_2, \ldots, g_d) = (x_1, \ldots, x_d)$ then $G^k(C(g_1, g_2, \ldots, g_d)) = \{(y_1, \ldots, y_d) | (y_i \mod 2 = 0) \land (y_i = x_i) \lor (y_i = x_i \pm 1) \land (y_i = x_i \pm 1)\}$. For example, $G^2((7, 2)) = \{(6, 2), (8, 2), (6, 0), (8, 0), (6, 4), (8, 4)\}.
8. An operation $F$ is valid on a configuration $C(g_1, g_2, \ldots, g_d)$ if $(x_1, \ldots, x_d) = F(C(g_1, g_2, \ldots, g_d))$ satisfies $LB \leq x_i \leq UB$ for all $i = 1, \ldots, d$, given predefined lower-bound LB and upper-bound UB. Otherwise, $F$ is invalid on $C$, $LB = 0$ and $UB = 9$ is used in the software, but the theory only requires that $UB > LB$.
9. A sequence of operations $F_1, \ldots, F_k$ is boundary-sensitive on configuration $C$ if $(x_1, \ldots, x_d) = F_k(\ldots(F_1(C(g_1, g_2, \ldots, g_d))))$ satisfies $LB \leq x_i \leq UB$ for all $i = 1, \ldots, d$ and $j = 1, \ldots, k$. We use boundary-insensitive to refer to a sequence on which this condition has not been checked.

**Progression model considering SD and CD events**

We develop the theory for inference of the Steiner (unsampled or extinct cell configurations) nodes in the paths formed by the sequence of gene copy number gains and losses from an initial configuration $C'(g_1, g_2, \ldots, g_d)$ to a final configuration $C(g_1, g_2, \ldots, g_d)$. We first extend the prior theory to account for SD and CD events. Our model assumes that on division of a tumor cell, the configuration can change either by gain or loss of one copy of a single gene (SD event) or by gain or loss of one copy of each gene on a single chromosome (CD event). For example, a configuration of four genes $(2, 2, 2, 2)$ with
the first two genes on the same chromosome might evolve in a single mutational event to (3,2,2,2) by an SD event or to (3,3,2,2) by a CD event. We propose Algorithm 1, provided in Figure 10, to calculate the minimum number of steps required to transform \( C^i(g_i, g_{i+1}, \ldots, g_j) \) into \( C^i(g_i, g_{i+1}, \ldots, g_j) \) considering SD and CD events, where, without loss of generality, we assume that the genes on a common chromosome have consecutive indices \((g_i, g_{i+1}, \ldots, g_j)\) in \( C \). Algorithm 1 also identifies a minimum-length sequence of events, although this sequence is not necessarily unique. For example, if there are four genes on one chromosome and we want to get from configuration \((1,1,1,1)\) to configuration \((2,4,3,2)\), then a shortest sequence of SD and CD events would be CD to \((2,2,2,2)\), SD to \((2,3,2,2)\), SD to \((2,4,2,2)\), and SD to \((2,4,3,2)\). Other orders of the same four events are also possible.

The above example focuses on a single chromosome because as explained below, the problem of finding the shortest SD+CD path can be solved one chromosome at a time. We begin by establishing the following lemmas:

Lemma 1. A minimum-length boundary-insensitive sequence of CD and SD events cannot have both a gain of chromosome \( c_i \) and a loss of the same chromosome \( c_i \).

Proof. By contradiction. Suppose \( S \) is a sequence of events that has both a gain and a loss of the same chromosome. Then removing one gain and one loss produces a new sequence that is 2 shorter and has the same final state.

Lemma 2. For any gene \( g_i \), a minimum-length boundary-insensitive sequence of events cannot have both a gain of \( g_i \) and a loss of \( g_i \).

Proof. By contradiction. Suppose \( S \) is a sequence of events that has both a gain of \( G \) and a loss of \( G \). Then removing one gain and one loss produces a new sequence that is 2 shorter and has the same final state.

Lemma 3. The following sequence of events describes a minimum-length boundary-insensitive sequence of SD and CD events for transforming \( C^i(g_i, g_{i+1}, \ldots, g_j) \) into \( C^i(g_i, g_{i+1}, \ldots, g_j) \):

1. Perform CD events in arbitrary order starting from \( C^i \) so that each successive event decreases the \( L_1 \) distance between the intermediate configurations \( C^{md}(g_i, g_{i+1}, \ldots, g_j) \) and \( C^i(g_i, g_{i+1}, \ldots, g_j) \) until any further CD event will increase the \( L_1 \) distance. We define the final configuration reached after this step to be \( C^f(g_i, g_{i+1}, \ldots, g_j) \).
2. Perform SD events in arbitrary order starting at \( C^f(g_i, g_{i+1}, \ldots, g_j) \) so that the \( L_1 \) distance between \( C^{md}(g_i, g_{i+1}, \ldots, g_j) \) and \( C^f(g_i, g_{i+1}, \ldots, g_j) \) decreases on each step until the distance becomes zero. The total number of events required will be \( L_1(C^i, C^f) \).

Proof. Since the sequence of events is boundary-insensitive and addition is commutative, we can change the order of events.
without changing the endpoints or the cost. Therefore, we assume that all CD events precede all SD events. The construction of the above sequence of the events ensures that it uses a maximum number of possible CD events. If we denote the number of genes on the common chromosome by $k$ and the number of CD events by $c$, then the total number of events required is $L_1(C, C^*) = (k - 1)c$. If there exists a shorter sequence of events to transform $C$ to $C^*$, then that sequence must have a larger number of CD events, which is contradicted by the construction. Thus, the number of events is minimized.

The above lemmas show how to construct a minimum-length boundary-insensitive sequence of events. We now establish that this sequence can be used to derive a minimum-length boundary-sensitive sequence of events:

**Lemma 4.** For any boundary-insensitive minimum-length sequence of SD and CD events $S$ transforming $C$ to $C^*$, there exists a boundary-sensitive sequence of SD and CD events $S'$ such that $S$ and $S'$ have equal length.

**Proof.** We analyze one chromosome at a time because in this section the events on different chromosomes are independent. By Lemma 1, on any specific chromosome all the CD events are gains or all the CD events are losses. We analyze in detail the case in which all CD events are losses; the case of all gains is symmetric.

The proof is constructive. Specifically, we will show that the upper part of Algorithm 1 will transform a boundary-insensitive $S$ to a boundary-sensitive $S'$ of equal cost solely by reordering events. Without loss of generality, suppose the only CD events in $S$ are chromosome losses. There is a symmetric algorithm, shown as the lower part of Algorithm 1, for the case where all the chromosome events are gains. We add the following definition:

A gene $G$ is defined as unidirectional with respect to $S$ if there are no gains of $G$ in $S$. A gene $G$ is defined as bidirectional with respect to $S$ if $S$ includes gains of $G$. For unidirectional genes, the order of chromosome losses and gene losses can never cause a boundary to be crossed because the copy numbers are monotonically decreasing. The situations we need to avoid are:

1. A bidirectional gene $G$ has copy number UB and the next operation affecting $G$ is a gain of $G$.
2. A bidirectional gene $G$ has copy number LB and the next operation affecting $G$ is a chromosome loss.

Chromosome gains are excluded by Lemma 1 and our assumption without loss of generality that all CD events are losses. Gene losses for bidirectional genes are excluded by Lemma 2.

To prove correctness of the algorithm, we note that $S'$ can never cross LB for the unidirectional genes because their net loss equals their total loss. $S'$ can never cross UB for the bidirectional genes, because when their copy number is at LB, a gene gain must still be pending and the gene gains alternate in the first while loop until no chromosome losses or gene gains are remaining. $S'$ can never cross UB for the unidirectional genes because they have only losses. $S'$ can never cross UB for the bidirectional genes because of the test $N^b < UB$ (line 8) before any gene gain is done. Further, all the chromosome losses will be used because one chromosome loss happens on each pass through the first while loop, if any chromosome losses remain. All gene gains in $S$ will be used in the first while loop because the net change for any gene must keep its copy number below UB. All the gene losses for the unidirectional genes are used in the second while loop. The unordered set of events and total change in each gene is thus preserved between $S'$ and $S$, while $S'$ guarantees that the sequence is boundary-sensitive.

We use the preceding result to derive the main theorem of this section, which establishes a method to find a minimum-length sequence of SD and CD events transforming $C$ to $C^*$. As in the proof of Lemma 4, we can consider each chromosome separately since each SD and CD event affects only one chromosome.

**Theorem 5.** Assume we partition the genome list by chromosomes such that each chromosome $c_i \in \{e_1, \ldots, e_r\}$ corresponds to a consecutive subset of genes $g_1, \ldots, g_{|d|}$. Further define $C(g_1, g_2, \ldots, g_{|d|}) = (s_1, \ldots, s_d)$ and $C'(g_1, g_2, \ldots, g_{|d|}) = (t_1, \ldots, t_d)$. Then we can construct a minimum-length boundary-sensitive sequence of events transforming $C(g_1, g_2, \ldots, g_{|d|})$ to $C'(g_1, g_2, \ldots, g_{|d|})$ by constructing a minimum-length boundary-sensitive sequence of events $S_i$ transforming $(s_1, \ldots, s_{i-1}, s_{i+1}, \ldots, s_d)$ to $(t_1, \ldots, t_{i-1}, t_{i+1}, \ldots, t_d)$ for each chromosome $c_i$ in arbitrary order.

**Proof.** The distance function can be decomposed into individual parts for genes belonging to distinct chromosomes as follows:

$$D^{\text{ch}}(C, C') = \sum_{i=1}^{n} D^{\text{ch}}(C'(s_1, \ldots, s_{i-1}), C'(s_{i+1}, \ldots, s_d))$$

Because the distance cost can be decomposed in this way and each CD or SD event contributes to only a single term of the outer sum, we can minimize the cost of events for each chromosome independently and combine the events from distinct chromosomes in arbitrary order without changing the value of the objective function. Likewise, since these each chromosome affects a disjoint subset of genes, boundary-sensitive sequences for each chromosome will yield a boundary-sensitive sequence across all genes.

**Progression model combining SD, CD and GD events**

We now extend the theory from the prior section to include SD, CD, and GD events. We assume in the proofs and discussion below that $C^* < C$, where $<$ denotes lexicographical ordering. This assumption reduces the number of cases in several proofs. If instead, $C^* < C^*$, the proofs are identical or symmetric except that GD events may be used in the wrong direction (halving instead of doubling). The use of halving events is corrected heuristically by a procedure of subtree pruning and regrafting at line 24 of the pseudocode of Algorithm 3, described below, and in FISHtrees. We will produce the complete proof by deriving a series of lemmas for three cases that together will cover all possible $C^*$ and $C^*$.

**Lemma 6.** For an an even configuration $C^*$, if there exists an optimal sequence of copy number change events from $C^*$ to $C^*$ composed of one or more SD and CD events and a single GD event, then the following sequence of events is of minimum length:

1. SD and CD events to transform $C^*$ into $H(C^*)$, constructed as described in the first named subsection of Methods
2. A single GD event to transform $H(C^*)$ into $C^*$.

**Proof.** We prove the statement by considering the three different ways that can be used to transform $C^*$ to $C^*$ using single GD and multiple SD and CD events. The statement of the lemma presents one case and the remaining two possibilities are as follows:

1. A single GD event to transform $C^*$ into $O(C^*)$ and then multiple SD and CD events to transform $O(C^*)$ to $C^*$.
2. Multiple SD and CD events to transform $C^*$ to an intermediate configuration $C'$, a single GD event to transform $C'$ into $C'$, and multiple SD and CD events to transform $C'$ into $C^*$.  

PLOS Computational Biology | www.ploscompbiol.org 13 July 2014 | Volume 10 | Issue 7 | e1003740
We show that for either of these alternative cases, we can produce a sequence satisfying the conditions of the lemma with equal or smaller length. For the first case, we have to show that

\[ D_{ch}(C^i, H(C^i)) < D_{ch}(C^i, O^i(C^i)) \]

It can be seen that

\[ L_i(C^i, H(C^i)) = \frac{1}{2} L_i(C^i, O^i(C^i)) \]

If all genes are located on distinct chromosomes, then,

\[ D_{ch}(C^i, H(C^i)) = \frac{1}{2} D_{ch}(C^i, O^i(C^i)) \]

and the claim follows directly.

Now, assume the genes are partitioned into sets of chromosomes such that each chromosome \( c_i \in \{c_1, \ldots, c_q\} \) corresponds to a consecutive subset of genes \( g_{i,1}, \ldots, g_{i,q_i} \). We focus on a specific chromosome \( c_i \) and consider the problem of updating just genes of that chromosome from their values in \( O^i(C^i) \) to their values in \( C^i \).
Either zero or a positive even number of CD events must be performed to convert these genes from $O^M(C^e)$ to $C^s$ and along with zero or a positive even number of SD operations on each gene. If an odd number of CD operations are performed on $O^M(C^e)$, then we get an odd configuration and at least one or an odd number of SD operations must be performed on each gene of this odd configuration to convert it to the even configuration $C^e$. But a combination of single SD operations acting on each of the individual genes in $g_1, \ldots, g_{\lfloor \frac{m}{2} \rfloor}$ has the same effect as a single CD operation on chromosome $c_i$ and this combination therefore cannot be minimal. Therefore, the number of CD operations is even. If a total of $m$ CD operations and $n$ SD operations are needed to convert $C^o$ to $C^s$, then a total of $\frac{1}{2} m$ CD operations and $\frac{1}{2} n$ SD operations are needed to convert $C^o$ to $C^e$, then a total of $\frac{1}{2}$ of $m$ CD operations and $\frac{1}{2}$ of $n$ SD operations are needed to convert $C^o$ to $C^e$. So,

$$D^{ch}(C^o,H(C^e)) < D^{ch}(C^e,O^M(C^e))$$

For alternative 2, we can write the distance function as:

$$D^{ch}(C^o,C^e) = D^{ch}(C^e,C^o) + 1 = D^{ch}(O^M(C^e),C^e)$$

The distance function for our proposed optimal sequence can be written as:

$$D^{ch}(C^o,C^e) = D^{ch}(C^e,C^o) + D^{ch}(C^o,H(C^e)) + 1$$

As shown for alternative 1, we can write:

$$D^{ch}(O^M(C^e),C^e) > D^{ch}(C^e,H(C^e))$$

which implies $D^{ch}(C^o,C^e) > D^{ch}(C^e,C^o)$.  

**Lemma 7.** For an odd configuration $C^e$, if the optimal sequence of copy number change events from $C^o$ to $C^e$ is composed of one or more SD and CD events, followed by a single GD event, then the configuration from which the final set of SD and CD events takes place is a member of $G^c(C^e)$. 

**Proof.** We denote the intermediate configuration following the GD event to be $C^m$. We will show by contradiction that if there exists any optimal sequence of events for which $C^m \notin G^c(C^e)$ then there must exist another, shorter sequence of events. Define the full sequence of events from $C^o$ to $C^e$ to be $\vec{p}$, subdivided into the subsequences $\vec{p}_1, (GD), \vec{p}_2$. First, we note that if there is any duplicated event in $\vec{p}_2$ then we can construct a more parsimonious solution by replacing the duplicate in $\vec{p}_2$ with a single copy of the event in $\vec{p}_1$. Therefore, no event appears more than once in $\vec{p}_2$. There are exactly two CD and SD events that can increase the count of any given probe (SD of that probe or CD of its chromosome) and similarly exactly two events that can decrease the count of any probe. Thus, no probe’s value changes by more than $\pm 2$ in the transition from $C^m$ to $C^e$ in $\vec{p}_2$. Finally, we note that since $C^m$ immediately follows a GD event, it must be an even configuration. Together, these assertions establish that $C^m \in G^c(C^e)$ for any optimal path $\vec{p}$.

**Lemma 8.** For an odd configuration $C^e$, if the optimal sequence of copy number change events from $C^o$ to $C^e$ is composed of one or more SD and CD events and a single GD event, then the optimum sequence of events follows the following path:

1. Generate $C^m = G^c(C^e)$.
2. SD and CD events to transform $C^o$ into $H(C^m)$.
3. A single GD event to transform $H(C^m)$ into $C^m$.
4. SD and CD events to transform $C^m$ into $C^e$.

**The optimal sequence is an element of the set of sequences generated using this procedure.**

**Proof.** The proof follows from application of Lemma 6 and Lemma 7. As $C^e$ is an odd configuration, the final step cannot be a GD event. So, the last steps have to be a combination of SD and/or CD events; in that case, Lemma 7 shows that the configuration reached as a result of GD must be a member of $G^c(C^e)$, which we denote by $C^m$. Lemma 6 shows that to reach any member of $G^c(C^e)$, which are even configurations, the optimal sequence of events is to generate SD and CD events to transform $C^o$ into $H(C^m)$ first and then to perform a GD event to transform $H(C^m)$ into $C^m$. This sequence of events matches the sequence proposed in the lemma.

The above lemmas allow us to derive Algorithm 2 to transform $C^o$ to $C^e$ using a minimum-length combination of SD, CD, and GD events. The pseudocode of Algorithm 2 is presented in Figure 11. To illustrate the algorithm, suppose $C^o = (3,1)$ and $C^e = (7,5)$, where we will assume we have two probes on a single chromosome. Since $C^e$ is an odd configuration, we first generate its nearest even neighbors $G^c(C^e) = ((6,4),(6,6),(8,4),(8,6))$ and calculate $H(G^c(C^e)) = ((3,2),(3,3),(4,2),(4,3))$. The algorithm tests for two stopping conditions by which a solution can be constructed (lines 22 and 24 in Algorithm 2), neither of which applies to any of the solutions at this point. $(3,2),(3,3),(4,2),(4,3)$ are therefore considered for the next iteration. $(3,2),(3,3)$, and $(4,3)$ are odd configurations, so we generate their neighbor sets $G^c((3,2)) = \{(2,2),(2,4),(2,0),(4,0),(2,4),(4,4)\}$, $G^c((3,3)) = \{(2,2),(2,4),(2,0),(4,0),(4,2),(4,4),(6,2),(6,4)\}$. One stopping condition is satisfied for each of the elements of these neighbor sets, so $(3,2),(3,3)$, and $(4,3)$ are each considered in turn as the next candidate neighbor. $(4,2)$ is an even configuration, so we only need to consider one possible stopping condition (line 11), which it satisfies, so it is also considered as a possible next candidate neighbor. Among the four possibilities, we will conclude that using $(3,2)$ as the immediate neighbor will lead to the smallest possible number of steps when accumulating SD+CD events from $C^o$ to the candidate, a single GD event from the candidate to its double, and SD+CD events from that double to $C^e$. Following some postprocessing updates (procedure CheckSrc-Neighbor), the algorithm computes a minimum-length solution of $(3,1) = (3,2) = (6,4) = (7,5)$ and returns the corresponding length 3.

Algorithm 2 satisfies the following theorem, which constitutes the major result of this section:

**Theorem 9.** Algorithm 2 returns the minimum distance between two configurations $C^o$ and $C^e$, where $C^e < C^o$.

**Proof.** We use induction on the minimum number of steps to get from $C^o$ to $C^e$, which we denote by $M(C^o,C^e)$.

**Base case.** For the base case, we have $M(C^o,C^e) = 1$. We must consider two sub-cases: (i) $C^o = 2C^e$ and (ii) $M(C^o,C^e) = 1$. For case (i), $C^o$ is an even configuration. The condition at line 11 in Algorithm 2 fails and $\frac{1}{2} C^e = C^o$ is considered for the next iteration. In the next iteration, if $C^o$ is an even configuration then the condition at line 11 is now satisfied and $M(C^o,C^e)$ is assigned the
Figure 11. Algorithm 2 pseudocode. Algorithm 2 finds the shortest directed distance between two configurations using SD, CD, and GD events.

doi:10.1371/journal.pcbi.1003740.g011

```plaintext
1: procedure MINIMUMDISTANCE(C*, C')  // Minimum distance between configurations C* and C'
2: Require: C*, C'  // C* and C' are configurations
3: Ensure: D^s,ch,g(C*, C'), a minimum-distance sequence of events, is stored implicitly via the parent function
4:     prev ← {C'}  // prev stores the configurations to be considered in the next iteration of the Algorithm
5:     dist(C*) ← ∞  // dist(C*) stores the optimal distance between C* and C' calculated so far
6:     D^s,ch(C*, C') ← Length of the optimal path between C* and C' constructed using the procedure defined by Theorem 5.
7:   while true do
8:     nextStates ← ∅  // nextStates is a set of configurations to be considered in the next iteration
9:     for i ← 1, |prev| do  // |prev| is the size of the set prev
10:        parent ← prev(i)  // parent is the configuration from which the current configuration is reached
11:        if parent is an even configuration then
12:            prevHalf ← \frac{1}{2}(prevConf)
13:            if D^s,ch(C*, parent) ≤ (D^s,ch(C*, prevHalf) + 1) then
14:                CHECKSRCNEIGHBOR(C*, parent, srcNeighbor, dist)
15:            else
16:                nextStates ← nextStates ∪ prevHalf  // add current configuration and its parent to the nextStates
17:                dist(prevHalf) ← dist(prevConf) + 1  // update the distance
18:                parent(prevHalf) ← prevConf  // update the parent
19:        else
20:            prevNeighborSet ← G^E(prevConf)
21:            for j ← 1, |prevNeighborSet| do  // |prevNeighborSet| is the size of the set prevNeighborSet
22:                prevNeighbor ← prevNeighborSet(j)  // prevNeighbor is the next configuration to be considered
23:                prevNeighborHalf ← \frac{1}{2}(prevNeighbor)
24:                if D^s,ch(C*, prevConf) ≤ (D^s,ch(prevConf, prevNeighbor) + D^s,ch(prevConf, prevNeighborHalf)) then
25:                    CHECKSRCNEIGHBOR(C*, prevConf, srcNeighbor, dist)
26:                else if D^s,ch(C*, prevNeighbor) ≤ (D^s,ch(C*, prevNeighborHalf) + 1) then
27:                    dist(prevNeighbor) ← dist(prevConf) + D^s,ch(prevConf, prevNeighbor)
28:                    parent(prevNeighbor) ← prevConf  // update the parent
29:                    CHECKSRCNEIGHBOR(C*, prevNeighbor, srcNeighbor, dist)
30:                else
31:                    nextStates ← nextStates ∪ prevNeighbor  // add current configuration and its parent to the nextStates
32:                    parent(prevNeighbor) ← prevConf  // update the parent
33:                    dist(prevNeighbor) ← dist(prevConf) + D^s,ch(prevNeighbor, prevConf)
34:                    parent(prevNeighborHalf) ← prevNeighbor  // update the parent
35:                    dist(prevNeighborHalf) ← dist(prevNeighbor) + 1  // update the distance
36:                if nextStates == ∅ then
37:                    break  // if no new configurations are added, break the loop
38:            end for
39:        end if
40:     end for
41: end while
42: return D^s,ch,g(C*, C') ← dist(C*)  // return the minimum distance

end procedure

procedure CHECKSRCNEIGHBOR(C*, prevConf, srcNeighbor, dist)  // Checks if prevConf can become the new candidate neighbor of C*
43: testDistance ← dist(prevConf) + D^s,ch(C*, prevConf)
44: if dist(C*) > testDistance then
45:    srcNeighbor ← prevConf  // update the srcNeighbor
46:    dist(C*) ← testDistance

end procedure
```
value 1 in CheckSrcNeighbor procedure called at line 12 in the main procedure. If $C'$ is an odd configuration, then the condition at line 22 is satisfied in the first iteration of the algorithm depending on whether $C'$ is an even or odd configuration and $M(C', C)$ is assigned the value $D^{C'}(C; C') = 1$ at line 12 or 23.

**Induction step.** For the induction hypothesis, we assume that the the algorithm uses the minimum number of steps for all cases where $M(C', C) \leq m$. Then, suppose that an adversary selects an example that has complexity $M(C', C) = m + 1$. Let us assume that the penultimate configuration in the optimal solution is $C^{\text{opt}}$. If $C'$ is an even configuration, then it can be reached from $C^{\text{opt}}$ by using (i) a GD event, (ii) an SD event, or (iii) a CD event. According to the induction hypothesis, for each of these cases, Algorithm 2 uses the minimum number of $m$ steps to generate $C^{\text{opt}}$ from $C'$. If there is at least one GD event in the optimal solution, then Algorithm 2 first calculates $C^{\text{opt}} = \frac{1}{2} C'$. The induction hypothesis ensures that $M(C', C) \leq m$ and thus, Algorithm 2 returns a solution with a maximum length of $m + 1$. If there is no GD event in the optimal solution from $C'$ to $C$, then Algorithm 2 uses the procedure described in the first named subsection of Methods to calculate the optimal path from $C^{\text{opt}}$ to $C'$ and combining it with the optimal solution from $C'$ to $C^{\text{opt}}$, it returns the optimal path between $C'$ and $C$. Now, if $C'$ is an odd configuration, then going from the penultimate configuration $C^{\text{opt}}$ to $C'$ can only be achieved using either an SD or a CD event. For odd $C'$, Algorithm 2 first generates its even neighbors $CN$ which are steps $§$ from $C'$. If $C^{\text{opt}} \in CN$, the proof follows directly from the inductive hypothesis. If $C^{\text{opt}} \notin CN$, then there is a $C^* \in CN$ such that $C^{\text{opt}}$ is located on the optimal path between $C^*$ and $C'$ formed using SD and CD events only. If $k$ is the total number of genes with odd copy number values in $C'$, then $D^{C^*}(C^*, C) \sim k$ and $D^{C^*}(C^*, C^{\text{opt}}) \sim k - 1$. Using the induction hypothesis, we can write,

$$M(C', C) \leq m - k + 1$$

As Algorithm 2 uses the procedure described in the first named subsection of Methods to construct the optimal path between $C'$ and $C$, we can see that it returns a path with $M(C', C) \leq m + 1$.

**Runtime analysis of Algorithm 2**

We provide an upper bound on the runtime of Algorithm 2 as a function of the number of genes $d$ and their copy numbers. Considering all three events, where $C' < C$, the maximum
number of doublings required is \( \left\lceil \log_2 \left( \frac{C'(g_i)}{C'(g_j)} \right) \right\rceil \), where \( g_i \) denotes the copy number of the first gene where \( C'(g_i) < C'(g_j) \) and \( C'(g_i) \geq 0 \). At each stage of the algorithm, the maximum number of nodes generated as a result of a \( G^2 \) operation is \( 3^d \). \( \delta \) and \( \overline{\delta} \) SD and CD events are used to create each of those \( 3^d \) nodes in the case of an odd configuration. So, the maximum number of required \( L_i \) operations is \( \left\lceil \log_2 \left( \frac{C'(g_i)}{C'(g_j)} \right) \right\rceil \delta \). Therefore, the number of operations performed during the execution of Algorithm 2 is \( O \left( \left\lceil \log_2 \left( \frac{C'(g_i)}{C'(g_j)} \right) \right\rceil \delta \right) \).

Generating tumor phylogenies

We implemented Algorithm 2 and integrated it with our approximate median-joining-based algorithm from our prior SD-only FISHtrees [46] code. The key steps of this algorithm are summarized in Algorithm 3 (Figure 12), which we describe at a high level here. The phylogeny algorithm first relies on Algorithm 2 to derive a matrix of pairwise distances between observed cell configurations, which are treated as states on a truncated integer lattice of dimension \( d \) with a maximum value \( \langle UB \rangle \) set to 9 in the current code. For each currently repeated triplets of nodes, identifying as potential Steiner nodes those that agree in each dimension with at least one of the triplets. These Steiner nodes that lead to reduced minimum spanning tree cost are added to the node set, with the process is repeated until there is no further improvement. Finally a series of post-processing steps are performed to prune Steiner nodes that are not needed for the final tree and to apply subtree regrafting to correct for a potential source of suboptimality arising from the fact that the core phylogeny algorithm assumes symmetric distances but GD operations are asymmetric.

Inferring tumor phylogenies using Neighbor Joining (NJ) and Maximum Parsimony (MP) methods

Neighbor Joining (NJ) and Maximum Parsimony (MP) methods have been commonly used for building single-tumor phylogenies [16,54] and we therefore compared their accuracy to that of our own methods in inferring copy number phylogenies. We applied these two traditional phylogenetic tree building methods to build tumor progression trees using the individual copy number profiles as taxa and compared them with the trees built using our algorithms. We used implementations of both approaches in MEGA version 6 [69]. For NJ, we used Euclidean distances between cell copy number profiles to build the pairwise distance matrix. For MP, we treated copy number profiles of the genes in individual cells as sequences of arbitrary phylogenetic characters. We used the “Close-Neighbor-Interchange on Random Trees” search method. For the parameters “Number of Initial Trees” and “MP search level”, we used values of 10 and 1 respectively.

Acknowledgements

We thank Darawaeel Wangsa for collecting the CC data and we thank Lisa Berroa Garcia, Amanda Bradley, and Clarymara Ostiez-Melendez for help in collecting the BC data.

Author Contributions

Conceived and designed the experiments: SAC SES KHH TR AAS RS. Performed the experiments: SAC AAS RS. Analyzed the data: SAC SES KHH TR AAS RS. Wrote the paper: SAC SES KHH TR AAS RS. Designed the software used in the analysis: SAC AAS RS.

References

1. Pennington G, Smith CA, Shackney S, Schwartz R (2006) Cancer phylogenetics from single-cell assays. Technical report, Carnegie Mellon University.
2. Pennington G, Smith CA, Shackney S, Schwartz R (2005) Reconstructing tumor phylogenies from heterogeneous single-cell data. J Bioinform Comput Biol 5: 407–427.
3. Attolini CSO, Michor F (2009) Evolutionary theory of cancer. Ann NY Acad Sci 1168: 23–51.
4. Desper R, Jiang F, Kallioniemi OP, Moch H, Papadimitriou CH, et al. (1999) Inferring tree models of oncogenesis from comparative genomic hybridization data. J Comput Biol 6: 57–71.
5. Desper R, Jiang F, Kallioniemi OP, Moch H, Papadimitriou CH, et al. (2000) Distance-based reconstruction of tree models for oncogenesis. J Bioinform Comput Biol 2: 789–803.
6. Szabo A, Boucher K (2002) Estimating an oncogenic tree when false negatives and positives are present. Math Biosci 176: 219–236.
7. McGlynn KA, Edmonson MN, Michielli RA, London WT, Lin WY, et al. (2005) Learning multiple evolutionary pathways from cross-sectional data. J Comput Biol 12: 584–598.
8. Beerwinkel N, Rahnenfuhrer J, Deumier M, Hoffmann D, Kaiser R, et al. (2003) Multivariate analyses of genomic imbalances in solid tumors reveal distinct and converging pathways of karyotypic evolution. Genes Chromosomes Cancer 31: 156–171.
9. Letouzé E, Allory Y, Bilotet MA, Radovanyi F, Guyen F (2010) Analysis of the copy number profiles of several tumor samples from the same patient reveals the successive steps in tumorigenesis. Genome Biol 11: R76.
10. Subramanian A, Shackney S, Schwartz R (2012) Inference of tumor phylogenies from genomic assays on heterogeneous samples. J Biomed Biotechnol: 797612.
11. Bogojeska J, Lengauer T, Rahnenfu¨hrer J (2008) Stability analysis of mixtures of mutagenetic trees. BMC Bioinformatics 9:163.
12. Franklin D, Wasserstrom A, Izukovitz S, Stern T, Harmelin A, et al. (2008) Cell lineage analysis of a mouse tumor. Cancer Res 68: 5924–5931.
13. Shahrabi Farahani H, Lagergren J (2013) Learning oncogenetic networks by inference from single-cell assays. Technical report, Carnegie Mellon University.
14. Liu J, Bandyopadhyay N, Ranka S, Baudis M, Kalveci T (2009) Inferring progression scores for GGH data. Bioinformatics 25: 2209–2215.
15. Notta F, Mallighan GC, Wang J, Pecoil P, Doulou S, et al. (2010) Evolution of human BCR-ABL1 lymphoblastic leukaemia-initiating cells. Nature 469: 362–367.
16. Bennington G, Smith CA, Shackney S, Schwartz R (2007) Reconstructing tumor phylogenies from single-cell assays. Technical report, Carnegie Mellon University.
17. Hoffl A, Gisselsson D, Mandahl N, Johansson B, Mertens F, et al. (2001) A phylogenetic analysis identifies heterogeneity among hepatocellular carcinomas. Hepatology 36: 1341–1348.
18. Newton MA (2002) Discovering combinations of genomic aberrations associated with hepatocellular carcinoma and the driving mutations revealed by cell-population genetic analytic whole-of-gene data. Proc Natl Acad Sci USA 105: 12042–12047.
19. Pennington G, Smith CA, Shackney S, Schwartz R (2006) Cancer phylogenetics from single-cell assays. Technical report, Carnegie Mellon University.
20. Fearon E, Vogelstein B (1990) A genetic model for colorectal tumorigenesis. Cell 61: 759–767.
21. Newton MA (2002) Discovering combinations of genomic aberrations associated with cancer. J Am Stat Assoc 97: 931–942.
22. Bilke S, Chen QR, Westerman F, Schwab M, Catchpoole D, et al. (2005) Learning multiple evolutionary pathways from cross-sectional data. J Comput Biol 12: 584–598.
23. Newton MA (2002) Discovering combinations of genomic aberrations associated with cancer. J Am Stat Assoc 97: 931–942.
24. Hjelm M, Ho¨glund M, Lagergren J (2006) New probabilistic network models to infer subtypes and algorithms for oncogenesis. J Comput Biol 13: 853–865.
25. Hou Y, Song L, Zhu P, Zhang B, Tao Y, et al. (2012) Single-cell single-exome sequencing and monoclonal evolution of a JAK2-negative myeloproliferative neoplasm. Cell 148: 873–883.
26. Letouzé E, Allory Y, Bilotet MA, Radovanyi F, Guyen F (2010) Analysis of the copy number profiles of several tumor samples from the same patient reveals the successive steps in tumorigenesis. Genome Biol 11: R76.
27. Subramanian A, Shackney S, Schwartz R (2012) Inference of tumor phylogenies from genomic assays on heterogeneous samples. J Biomed Biotechnol: 797612.
28. Bean E, Vogelstein B (1990) A genetic model for colorectal tumorigenesis. Cell 61: 759–767.
29. Hjelm M, Gisselsson D, Mandahl N, Johansson B, Mertens F, et al. (2001) Multivariate analyses of genomic imbalances in solid tumors reveal distinct and converging pathways of karyotypic evolution. Genes Chromosomes Cancer 31: 136–171.
30. Hou Y, Song L, Zhu P, Zhang B, Tao Y, et al. (2012) Single-cell single-exome sequencing and monoclonal evolution of a JAK2-negative myeloproliferative neoplasm. Cell 148: 873–883.
31. Letouzé E, Allory Y, Bilotet MA, Radovanyi F, Guyen F (2010) Analysis of the copy number profiles of several tumor samples from the same patient reveals the successive steps in tumorigenesis. Genome Biol 11: R76.
32. Heselmeyer-Haddad K, Chaudhri N, Stoltzfus P, Cheng JC, Wilber K, et al.
31. Janocko LE, Brown KA, Smith CA, Gu LP, Pollice AA (2001) Distinctive
30. Purdom E, Ho C, Grasso CS, Quist MJ, Cho RJ, et al. (2013) Methods and
48. Bandelt H, Forster P, Rohland K, et al. (2012) Single-cell genetic analysis of ductal carcinoma in situ and invasive breast cancer reveals enormous tumor heterogeneity, yet conserved genomic imbalances and gain of MYC during progression. Am J Pathol 181: 1807–1822.
47. Sottoriva A, Spiteri I, Shibata D, Curtis C, TavARES S (2013) Single-molecule analysis of multiple receptor tyrosine kinase genes in glioblastoma. Cancer Cell 20: 810–817.
46. Chowdhury SA, Shackney SE, Heselmeyer-Haddad K, Ried T, Scha¨ffer AA, et al. (2012) Evolutionary pathways in BRCA1-associated breast tumors. Cancer Discov 2: 503–511.
45. Saitou N, Nei M (1987) The neighbor-joining method: a new method for reconstructing phylogenetic trees. Mol Biol Evol 4: 406–425.
44. Sprouffske K, Pepper JW, Maley CC (2011) Accurate reconstruction of the temporal order of mutations in neoplastic progression. Cancer Prev Res 4: 1135–1144.
43. Urbschat S, Rahnenfu¨hrer J, Henn W, Fedien W, Wemmert S, et al. (2011) Clonal cytogenetic progression within intratumorally heterogeneous meningio-
42. Ding L, Raphael BJ, Chen F, Wendl MC (2013) Advances for studying clonal evolution inferred by single-cell sequencing. Nature 497: 90–94.
41. Nowell PC (1976) The clonal evolution of tumor cell populations. Science 194: 23–28.
39. Xu X, Hou Y, Yin X, Bao L, Tang A, et al. (2012) Single-cell exome sequencing estimates of rearrangement phylogeny for cancer genomes. Genome Res 22: 1135–1144.
38. Gerlinger M, Rowan AJ, Horswell S, Larkin J, Endesfelder D, et al. (2012) Intra-tumor heterogeneity of receptor tyrosine kinases EGFR and PDGFRα amplification in glioblastoma defines subpopulations with distinct growth factor response. Proc Natl Acad Sci USA 109: 3041–3046.
37. Navin N, Kendall J, Troge J, Andrews P, Rodgers L, et al. (2011) Tumour evolution inferred by single-cell sequencing. Nature 477: 90–94.
36. Heselmeyer-Haddad K, Berroa Garcia LY, Bradley A, Ortiz-Melendez C, Lee WJ, et al. (2012) Single-cell genetic analysis of ductal carcinoma in situ and invasive breast cancer reveals enormous tumor heterogeneity, yet conserved genomic imbalances and gain of MYC during progression. Am J Pathol 181: 1807–1822.
35. Szerlip NJ, Pedraza A, Chakravarty D, Azim M, McGuire J, et al. (2012) Intratumoral heterogeneity of receptor tyrosine kinases EGFR and PDGFRα amplification in glioblastoma defines subpopulations with distinct growth factor response. Proc Natl Acad Sci USA 109: 3041–3046.
34. Martins FC, De S, Almenroth V, Gonen M, Park SY, et al. (2012) Evolutionary pathways in BRCA1-associated breast tumors. Cancer Discov 2: 503–511.
33. Snuderl M, Fazlollahi L, Le LP, Nitta M, Zhelyuzkova BH, et al. (2011) Mosaic amplification of multiple receptor tyrosine kinase genes in glioblastoma. Cancer Cell 20: 810–817.
32. Heselmeyer-Haddad K, Chaudhri N, Stoltzfus P, Cheng JC, Wilber K, et al.
31. Janocko LE, Brown KA, Smith CA, Gu LP, Pollice AA (2001) Distinctive
30. Purdom E, Ho C, Grasso CS, Quist MJ, Cho RJ, et al. (2013) Methods and