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Computational tools to detect signatures of mutational processes in DNA from tumours: A review and empirical comparison of performance

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Abstract

Mutational signatures refer to patterns in the occurrence of somatic mutations that might be uniquely ascribed to particular mutational process. Tumour mutation catalogues can reveal mutational signatures but are often consistent with the mutation spectra produced by a variety of mutagens. To date, after the analysis of tens of thousands of exomes and genomes from about 40 different cancer types, tens of mutational signatures characterized by a unique probability profile across the 96 trinucleotide-based mutation types have been identified, validated and catalogued. At the same time, several concurrent methods have been developed for either the quantification of the contribution of catalogued signatures in a given cancer sequence or the identification of new signatures from a sample of cancer sequences. A review of existing computational tools has been recently published to guide researchers and practitioners through their mutational signature analyses, but other tools have been introduced since its publication and, a systematic evaluation and comparison of the performance of such tools is still lacking. In order to fill this gap, we have carried out an empirical evaluation of the main packages available to date, using both real and simulated data.

Among other results, our empirical study shows that the identification of signatures is more difficult for cancers characterized by multiple signatures each having a small contribution. This work suggests that detection methods based on probabilistic models, especially EMu and bayesNMF, have in general better performance than NMF-based methods.

Introduction

Cancers contain mutant cells that originate from a DNA modification in a single normal cell that is then propagated through cell divisions and that accumulates with further DNA modifications finally leading to abnormal, cancerous cells [1]. Such changes that include Single
Nucleotide Variants (SNVs), insertions or deletions, Copy Number Variation (CNVs) and chromosomal aberrations, have been termed as "somatic mutations" to distinguish them with those inherited and transmitted from parents (germline mutations). With the development and the improvement of sequencing technologies collectively referred to as High-Throughput Sequencing (HTS) and the availability of cancer exome and genome data from most human cancers, much has been learnt about these mutations and the set of genes (oncogenes and tumour suppressor genes) that operate in human cancers. Beyond the relatively small number of "driver" mutations, those that confer a large selective advantage for tumour development and progression, the vast majority of mutations confer weaker selective advantage or are truly neutral in that they do not affect cancer cells' survival. Together, passenger and driver mutations constitute a record of all cumulative DNA damage and repair activities occurred during the cellular lineage of the cancer cell [2].

Among all types of somatic mutations, a particular focus has been placed on single base substitutions that have been classified in six types according to the mutated pyrimidine base (C or T) in a strand-symmetric model of mutation. Such 6 substitutions (C \( \rightarrow \) A, C \( \rightarrow \) G, C \( \rightarrow \) T, T \( \rightarrow \) A, T \( \rightarrow \) C and T \( \rightarrow \) G) may be further classified in different types when considering the sequence pattern in which they are located (sequence context). For practical reasons, the sequence context is typically defined using the 5' and 3' bases proximal to the mutated base, that results in substitutions being classified in 96 types (6^4). In this paper we adopt this convention and use "mutations" to refer to these 96 types of contextualized substitutions.

It has been hypothesised that specific patterns of mutations are the results of the action of particular underlying mutational process. To identify such patterns, computational models, such as matrix decomposition algorithms or probabilistic models, have been developed. The first of such methods was published in 2013 by Alexandrov and colleagues [2,3], and, as for most of all the other models that followed, is based on the idea that a mutational signature can be seen as a probability distribution of the 96 types of mutations. Mutational signatures contribute to the total mutational burden of a cancer genome, commonly referred to as mutational "catalogue" or "spectrum" in the recent computational biology literature.

Such original methodological framework, applied to tens of thousands of genomes and exomes from 40 different cancers types from large data repositories such as TCGA (The Cancer Genome Atlas), has led to the identification of 30 mutational signatures characterized by a unique probability profile across the 96 mutation types. These validated mutational signatures are listed in a repertory on the COSMIC (Catalogue Of Somatic Mutations In Cancer) website [3] and have been widely used as references. More recently, Alexandrov et al. have introduced an updated set of signatures identified from an even larger collection of both exome and whole-genome sequences (including the sequences from the PCAWG project) using two different methods (a new version of the original framework and a Bayesian alternative) [4]. The new repertory includes 49 new mutational signatures based on single base substitutions as in COSMIC, and also mutational signatures built in the context of other types of mutations such as double base substitutions (11 signatures), clustered based substitutions (4 signatures) and small insertions and deletions (17 signatures).

After the introduction of the original framework, several other mathematical methods and computational tools have been proposed to detect mutational signatures and estimate their contribution to a given catalogue. These methods can be grouped in two categories with different goals. The first class of methods aims to discover novel signatures while the second class aims to detect the known and validated mutational signatures in the mutational catalogue of a given sample. The approaches used in the first class are referred to as “de novo” (or “signature extraction”) while those in the second class as “refitting” (or “signature fitting”). All methods

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have been implemented in open source tools, mainly R packages, but some of them are available through command line, the Galaxy project or a web interface.

Signatures identified with de novo methods can be compared to reference signatures (for instance those listed in COSMIC) through measures such as cosine [5] or bootstrapped cosine similarity [6], which is a distance metric between two non-zero vectors. In this step of the analysis, extracted signatures are matched to the most similar reference signature, provided that their similarity is greater than a fixed threshold.

Recently, Baez-Ortega and Gori published a paper that discussed the mathematical models and computational techniques for mutational signature analysis [7]. However, to our knowledge no systematic evaluation of the performance of the existing methods has been conducted and the issue of the choice of an appropriate cosine similarity threshold when matching a newly extracted signature to the most similar counterpart in a reference set has not been addressed yet.

We use simulated and real-world cancer sequences in order to evaluate and compare for the first time the methods available to date, and assess their ability to accurately detect the underlying mutational signatures in mutational catalogues.

**Mathematical definition of mutational catalogues, spectra and signatures**

In this section we briefly review the mathematical framework and notations based on the original model developed by Alexandrov and colleagues [2]. The data structure that is used as input by most current methods is the mutational catalogue, or spectrum, of a genome (or exome), with the counts for each of the 96 different mutation features, or types. In most approaches, the notion of mutation type refers to base substitutions within the trinucleotide context described above.

The mutational catalogue of a genome (or exome) \( g \) is defined as the vector \( \left( m_1^g, \ldots, m_K^g \right)^T \), where each \( m_k^g \) is the number of mutations of type \( k \) found in the genome and \( K \), the number of possible mutation types, is equal to 96, and the superscript \( T \) denotes the transpose of a matrix so that vectors are thought as column vectors. Note that in this setting, information about mutation locations in the sequence is lost. The catalogue is built by comparing the sequence to a reference sequence in order to detect mutations and then by simply counting the occurrences of each type. The reference sequence can either be a standard reference (e.g. the assembly GRCh38) or a sequence from a “normal” tissue from the same individual (e.g. DNA from blood or from normal tissue surrounding tumours when available). For the purposes of the present work we will use the generic term “samples” for both genomes and exomes as the concepts and models used may be applied to both.

The basic idea underlying all computational models proposed is that the mutational catalogue of a sample results from the combination of all the mutational processes operative during lifetime, and therefore it can be seen as the weighted superposition of simpler mutational signatures, each uniquely corresponding to a specific process. The weight is larger if the process has a larger role in the final catalogue of mutations: for example, mutagens that last longer, are more intense, generate poorly repaired DNA lesions, mutate more genes, or also act as selection pressures favouring mutant cells. Here we adopt the term “signature” as it is the most commonly used in the recent computational cancer biology literature, although this term could be misleading as it implies unique backward identification with a mutagen or a cancer type that in some cases might not be possible.

Formally, the signature of a mutational process \( n \) is a vector \( p_n = (p_1^n, \ldots, p_K^n)^T \), where each \( p_k^n \) represents the probability that the mutational process will induce a mutation of type \( k \). In
other words, \( p_k^n \) is the expected relative frequency of type \( k \) mutations in genomes exposed to \( n \). Note that \( \sum_{k=1}^{K} p_k^n = 1 \) and \( 0 \leq p_k^n \leq 1 \) for all \( k \).

The intensity of the exposure to a mutational process \( n \) in a sample \( g \) is measured by the number of mutations \( e^n_g \) that are due to \( n \). For this reason, \( e^n_g \) is referred to as the “exposure” of \( g \) to \( n \). It is important to note that the term “exposure” does not refer here to the exposure to a mutagen \( \text{per se} \), because it also includes the likelihood that an unrepaired DNA lesion will cause a mutation. The expected number of mutations of type \( k \) due to the process \( n \) in sample \( g \) is therefore \( p_k^n e^n_g \). If sample \( g \) has been exposed to \( N \) mutational processes, then the total number of mutations of type \( k \) is

\[
m_k^g = \sum_{n=1}^{N} p_k^n e^n_g + \epsilon_k^g
\]

where \( \epsilon_k^g \) is an error term reflecting sampling variability and non-systematic errors in sequencing or subsequent analysis.

Matrix notation is effectively used when dealing with several samples and signatures. In this situation, the collection of \( G \) samples is represented by the \( K \times G \) matrix, with catalogues in columns:

\[
M = \begin{pmatrix}
m_1^1 & m_1^2 & \ldots & m_1^G \\
m_2^1 & m_2^2 & \ldots & m_2^G \\
\vdots & \vdots & \ddots & \vdots \\
m_K^1 & m_K^2 & \ldots & m_K^G
\end{pmatrix},
\]

the \( N \) signatures are represented by the \( K \times N \) matrix

\[
P = \begin{pmatrix}
p_1^1 & p_1^2 & \ldots & p_1^N \\
p_2^1 & p_2^2 & \ldots & p_2^N \\
\vdots & \vdots & \ddots & \vdots \\
p_K^1 & p_K^2 & \ldots & p_K^N
\end{pmatrix},
\]

and the exposures by the \( N \times G \) matrix

\[
E = \begin{pmatrix}
\epsilon_1^1 & \epsilon_1^2 & \ldots & \epsilon_1^G \\
\epsilon_2^1 & \epsilon_2^2 & \ldots & \epsilon_2^G \\
\vdots & \vdots & \ddots & \vdots \\
\epsilon_N^1 & \epsilon_N^2 & \ldots & \epsilon_N^G
\end{pmatrix}
\]

Eq (1) then becomes

\[
M \approx P \times E
\]

where we omitted the error term.

**de novo extraction vs. refitting**

*de novo* signature extraction methods aim at estimating \( P \) and \( E \) given \( M \). Non-negative matrix factorization (NMF) is an appealing solution to this unsupervised learning problem, because, by definition, all involved matrices are non-negative [8]. NMF identifies two matrices \( P \) and \( E \) that minimize the distance between \( M \) and \( P \times E \). In particular, NMF finds an approximated solution to the non-convex optimization problem

\[
\arg\min_{P \geq 0, E \geq 0} \| M - P \times E \|_F^2,
\]
where the Frobenius matrix norm of the error term is considered. NMF requires the number of signatures \( N \), an unknown parameter, to be predefined. An approach for selecting this parameter consists in obtaining a factorization of \( M \) for several of its values and then choosing the best \( N \) with respect to some performance measure such as the reconstruction error or the overall reproducibility. NMF is at the core of the Wellcome Trust Sanger Institute (WTSI) Mutational Signature Framework, the first published method for signature extraction [2]. An alternative to numerical approaches based on NMF is given by statistical modelling and algorithms. With these latter approaches, the number of mutations of a given type can be modelled by a Poisson distribution

\[
m_k^g \sim \mathcal{P} \left( \sum_{n=1}^{N} p_n^g e_n^g \right)
\]

where mutational processes are assumed to be mutually independent. This latter independence hypothesis simplifies the mathematics but does not necessarily hold in practice, where mutation processes are likely to interfere with each other (e.g. distinct defective DNA repair processes). In order to estimate \( E \) and \( P \), it has been proposed to consider \( E \) as latent data and \( P \) as a matrix of unknown parameters and to apply an expectation-maximization algorithm [9] or use Bayesian approaches [10–12]. One important advantage of statistical approaches is the availability of model selection techniques for the choice of \( N \). We refer to the recent review [7] for a comprehensive overview of the mathematical and computational aspects of existing methods, including alternative techniques not mentioned above.

The refitting approaches consider that the signatures \( P \) are known and the goal is to estimate \( E \) given \( M \) and \( P \). Refitting can be done for individual mutational catalogues (i.e. individual samples) and, from a linear algebra perspective, can be seen as the problem of projecting a catalogue living in the \( K \)-dimensional vector space (the space spanned by all mutation types) onto its subset of all linear combinations of the given mutational signatures having non-negative coefficients (the cone spanned by the given signatures).

A current practice consists in first performing a de novo extraction of signatures followed by a comparison of the newly identified signatures with the reference signatures (e.g. COSMIC signatures) by means of a similarity score, typically cosine similarity ranging from 0 (completely different) to 1 (identical) [6,10]. Finally, a “novel” signature is considered to reflect a specific reference signature if the similarity is larger than a fixed cut-off. If similarity is observed with more than one reference signature, the one with the largest value of similarity is chosen, see S1 Fig. This assignment step crucially depends on the choice of the cut-off \( \hat{h} \) that has been so far inconsistent in the literature with some studies using a value of 0.75 [11] whereas others 0.80 [12,13]. Another difficulty is that different signatures might have very close cosine similarity, as it happens also between COSMIC signatures, see S2 Fig, so that a unique assignment is not always possible. This shows that mutational signatures are a useful mathematical construct that, however, might have biological ambiguous meaning.

**Overview of the available tools**

A similar number of de novo and refitting methods exist and all of them are available as open source tools, mainly as R packages, or web interfaces (Table 1). The typical input of the tools is a file including the mutation counts but some tools derive the mutation counts from ad-hoc input files that may include for each individual a list of mutated bases, their position within the genome and the corresponding bases from a reference genome. The typical format of such input files is MAF, VCF or less common formats such as MPF and MFVF. For biologists or those who are not familiar with programming, a set of tools were also developed and provided
## Table 1. Available tools for the detection of mutational signatures.

<table>
<thead>
<tr>
<th>Software</th>
<th>Available platform/model</th>
<th>Input files</th>
<th>Additional features</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>de novo approaches</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>EMu [9] <a href="https://github.com/andrej-fischer/EMu">https://github.com/andrej-fischer/EMu</a></td>
<td>Command line/EM algorithm</td>
<td>- Mutation counts file</td>
<td>- An improved version has been recently implemented in SigProfiler</td>
</tr>
<tr>
<td>pmsignature [16] <a href="https://github.com/friend1ws/pmsignature">https://github.com/friend1ws/pmsignature</a></td>
<td>R/mixed-membership model</td>
<td>- Mutation Position Format -Mutation Feature Vector Format</td>
<td>- Selection of the optimal number of signatures</td>
</tr>
<tr>
<td>mutSignatures [20] <a href="https://cran.r-project.org/web/packages/mutSignatures/index.html">https://cran.r-project.org/web/packages/mutSignatures/index.html</a></td>
<td>R/NMF</td>
<td>Mutation counts file</td>
<td>- Selection of the optimal number of signatures - Data pre-treatment with the function get. legor96.hyper reduces the influence of hypermutated catalogues</td>
</tr>
<tr>
<td>Helmsman [21] <a href="https://github.com/carjed/helmsman">https://github.com/carjed/helmsman</a></td>
<td>Python/ NMF and PCA</td>
<td>- Variant Call Format - Mutation Annotation Format</td>
<td>- Able to run in parallel and designed for large datasets - Connection to external packages (in R) - may generate mutational catalogues from sequence data</td>
</tr>
<tr>
<td>SigProfiler [2,4] <a href="https://fr.mathworks.com/matlabcentral/fileexchange/38724-sigprofiler">https://fr.mathworks.com/matlabcentral/fileexchange/38724-sigprofiler</a></td>
<td>Matlab/ NMF</td>
<td>Mutation counts file</td>
<td>- Further development of the original framework - Two steps: 1) extraction of a minimal set of signatures, 2) estimation of their contributions to individual samples</td>
</tr>
<tr>
<td>SparseSignatures [22] <a href="https://bioconductor.org/packages/release/bioc/html/SparseSignatures.html">https://bioconductor.org/packages/release/bioc/html/SparseSignatures.html</a></td>
<td>R/ NMF with Lasso-penalized cost function</td>
<td>Mutation counts file</td>
<td>- Integration of DNA replication error signature - Sparse signature matrix - Number of signatures estimated with cross-validations - Scalable to large datasets</td>
</tr>
</tbody>
</table>

(Continued)
with user-friendly interfaces (see also Table 1). Some tools include additional features such as the possibility to search for specific patterns of mutations (e.g. APOBEC-related mutations [5]) and differential analysis [14].
**de novo approaches**

Most tools that have been developed to identify mutational signatures were based on decomposition algorithms including NMF or a Bayesian version of NMF \[14,17\]. The original method developed by Alexandrov et al. was based on NMF and was implemented in MATLAB \[2\] and is available also as an R package developed independently \[20\]. An updated and elaborated version named SigProfiler, was proposed recently for extracting a minimal set of signatures and estimating their contribution to individual samples \[4\]. The latter article also discusses an alternative method based on Bayesian NMF, called SignatureAnalyzer, that led to the identification of 49 reference signatures. Another tool that utilises NMF is maftools that is one of the few \(\text{de novo}\) tools that allows systematic comparison with the 30 validated signatures in COSMIC by computing cosine similarity and assigning the identified signatures to the COSMIC one with the highest cosine similarity \[5\].

Other tools such as SomaticSignatures \[15\] or the recent Helmsman \[21\] allows the identification of mutational signatures through Principal Component Analysis (PCA) in addition to NMF. For the sake of our formal comparison of the tools performance, we have tested SomaticSignatures only with the NMF implementation because with PCA the factors are orthogonal and the values inside the matrix can potentially be null or negatives, which is a deviation from the paradigm postulating that catalogues are the superposition of positively weighted signatures. We note, however, that PCA could be a promising way to explore complex situations in which mutational processes interfere with each other (e.g. relatively error free repair processes competing with error prone repair processes). Developed in the Python language, Helmsman allows the rapid and efficient analysis of mutational signatures directly from large sequencing datasets with thousands of samples and millions of variants.

A promising recent method called SparseSignatures \[22\] proposes an improvement of the traditional NMF algorithm based on two innovations, namely the default incorporation of a background signature due to DNA replication errors and the enforcement of sparsity in identified signatures through a Lasso penalty. This latter feature allows the identification of signatures with well-differentiated profiles, thus reducing the risk of overfitting.

In addition to decomposition methods, an approach based on the Expectation Maximization (EM) algorithm has been proposed to infer the number of mutational processes operative in a mutational catalogue and their individual signatures. This approach is implemented in the EMu tool \[9\], where the underlying probabilistic model assumes that input samples are independent and the number of mutational signatures is estimated using the Bayesian Information Criterion (BIC).

Another tool that uses a probabilistic model named mixed-membership model is pmsignature \[16\]. This tool utilises a flexible approach that at the same time reduces the number of estimated parameters and allows to modify key contextual parameters such as the number of flanking bases.

The latter feature may be particularly useful as the standard and most commonly used methods based on trinucleotides may not be the most adequate to detect specific mutational processes that lead to larger-scale substitution patterns. Evaluating the impact of limiting to trinucleotides or estimating the gain in performance associated with the extension of the context sequence to two flanking bases, is difficult and beyond the scope of our work. However, it is worth noting that trinucleotide-based methods have been able to identify several signatures associated with defective DNA mismatch repair and microsatellite instability (i.e. signatures 6, 14, 15, 20, 21, 26 and 44 of COSMIC v3) \[4\]. It is important to note that for the purpose of our comparison with the other tools, we set to one the number of flanking bases, and therefore to 96 mutation types.
EMu, signeR and pmsignature (and the refitting tool deconstructSigs) have been designed
to take into account the distribution of triplets in a reference exome or genome for example
from a sequence of normal tissue in the same individual. This is done by “normalizing” the
input mutational catalogues with respect to the distribution of triplets in the reference exome
or genome using an “opportunity matrix”.

**Refitting with known mutational signatures**

In addition to the identification of novel mutational signatures, we are often interested in eval-
uating whether a signature observed in an individual tumour is one of an established set of sig-
natures (e.g. the COSMIC signatures). This task is performed by “refitting tools” that aim to
search for the “best” combination of established signatures that explains the observed muta-
tional catalogue by projecting the latter (i.e. mapping) into the multidimensional space of all
non-negative linear combinations of the \( N \) established signatures.

The deconstructSigs [23] tool searches for the best linear combination of the established sig-
natures through an iterative process based on multiple linear regression aimed at minimizing
the distance between the linear combination of the signatures and the mutational catalogue.
All the other tools minimise the distance through equivalent approaches based on quadratic
programming [24,25,28], non-negative least square [26] linear combination decomposition
[27] and simulated annealing [25].

**A faster implementation.** We propose an alternative implementation of the decomposi-
tion performed in [26,27] based on a simple geometric framework. Finding the linear decom-
position of the input catalogue \( M \) on a set of given signatures minimizing the distance can be
seen as the problem of projecting \( M \) on the geometric cone whose edges are the reference sig-
natures. We propose to solve this problem by applying the very efficient R package called cone-
proj [34]. More details about our algorithm, which we called MutationalCone, together with
the R code implementing it, can be found in the S1 File.

**Combining de novo and refitting procedures**

Sigfit [29] is a recently introduced R package for Bayesian inference based on two alternative
probabilistic models. The first of such models is a statistical formulation of classic NMF where
signatures are the parameters of independent multinomial distributions and catalogues are
sampled according to a mixture of such distributions with weights given by the exposures,
while the second model is a Bayesian version of the EMu model. An interesting innovation of
Sigfit is that it allows the fitting of given signatures and the extraction of undefined signatures
in the same Bayesian process. As argued by the authors, this unique feature might be helpful in
cases where the small sample of catalogues makes it difficult to try to identify new signatures
or when the aim is to study the heterogeneity between the primary tumour and metastasis in
terms of the signatures they show.

In this work we empirically evaluate the methods that have been already presented in a peer
review published paper and for which an implementation in R is available. To this aim, we
adopt the COSMIC set as reference for the analysis of simulated and real mutational catalogues
because we evaluate tools that were developed at the time when COSMIC was the only avail-
able database of reference.
Data and experimental settings

Real cancer data from TCGA

In order to evaluate the performance of the available algorithms on real data, we used the exome sequences in The Cancer Genome Atlas (TCGA) repository (https://cancergenome.nih.gov/) for four cancer types: breast cancer, lung adenocarcinoma, B-cell lymphoma, and melanoma.

Mutation Annotation Format (MAF) files with the whole-exome somatic mutation datasets from these cohorts were downloaded from the portal gdc.cancer.gov on 6 March 2018. Data were annotated with MuSE [35] and the latest human reference genome (GRCh38). Mutational catalogues from these cohorts were obtained by counting the number of different mutation types using maftools [5]. The distribution of the number of mutations for each sample and separately for each cancer type is depicted in Fig 1.

According to the COSMIC website, 13 and 7 signatures have been found for breast cancer and lung adenocarcinoma respectively, 6 for B-cell lymphomas and 5 for melanoma.

Simulated data

The first key assumption of our model of simulated mutational catalogues is that the number of mutations in a sample $g$ that are induced by process $n$ follows a zero-inflated Poisson (ZIP) distribution. According to this two-component mixture model, $e^g_n$ is either 0 with probability $\pi$ or is sampled according to a Poisson distribution $\mathcal{P}(\lambda_n)$ with total probability $1-\pi$. Such a model depends on two parameters: the expectation $\lambda_n$ of the Poisson component, and the probability $\pi$ of extra “structural” zeros. The ZIP model allows for frequent zeroes and is therefore more suitable for modelling a heterogeneous situation where some samples are not exposed to a given mutational process ($e^g_n = 0$) while some others are ($e^g_n > 0$). Realistically, the mutation counts due to process $n$ in each of the $G$ samples, $e_1^n, \ldots, e_G^n$, are assumed to be independent and identically distributed according to a ZIP model where the expectation of the Poisson component is specific to $n$:

$$e^g_n \sim \text{ZIP}(\lambda_n, \pi) \text{, for all } n = 1, \ldots, N.$$ 

Note that the expected number of mutations in sample $g$ due to process $n$ is $(1-\pi)\lambda_n$. This flexibility given by process-specific average counts is the second important characteristic of our model and reflects the possibility that the mutagenic actions of different processes are intrinsically different with respect to their intensity. Obviously, it would have been possible to do one step further and allow for parameters $\lambda_{ng}$ specific to both processes and samples, thus representing the realistic situation in which the exposures of different samples to the same process have different duration or intensity (e.g. smokers/non-smokers). However, this would have resulted in too many parameters to tune, thus making it difficult to interpret the results of our simulation study. For the same reason we considered one fixed value of $\pi$.

The parameter $\lambda_n$ depends on both the average total number $r$ of mutations in a sample and the relative contribution of $n$. We therefore imposed the parameterization $\lambda_n (1-\pi) = q_n r$, where $q_n$ is the average proportion of mutations due to the process $n$.

When taking a unique value of $r$, this model produces realistic simulations even though it underrepresents extreme catalogues with very large or small total numbers of mutations (see Part C of S3 Fig). While considering a specific value of $r$ for each sample, or group of samples, would definitely make it possible to obtain a more realistic distribution of simulated catalogues (see Part B of S3 Fig), such multidimensional parameter would complicate unnecessarily our empirical assessment of mutational signature detection methods by introducing too many
specifications. We therefore decided to consider a unique \( r \) for each set of simulations. This formulation allowed us to study empirically the performance of a given signature detection method as a function of the average number of mutations \( r \) while fixing the average proportion of mutations due to each mutational process \( q_n \), according to different profiles that mimic real cancer catalogues. Interestingly, the ZIP model appeared to be more appropriate to represent mutational catalogues than the pure Poisson model used in previous publications [9,14] (see Part D of S3 Fig).

We adopted the following simulation protocol:

1. We chose \( N \) signatures from the COSMIC database, thus obtaining the matrix \( P \).
2. For each sample \( g \) and process \( n \) we sampled \( e^e_n \) from a ZIP distribution with parameters \( \lambda_n = q_n r / (1 - \pi) \) and \( \pi \) and obtained \( E \). Here \( q_n \), \( r \) and \( \pi \) are fixed parameters to set.
3. We computed the product \( P \times E \). In order to obtain the final simulated catalogue \( M \), we added some noise to the latter matrix by taking \( m_k^g \sim P((P \times E)_k^g) \).

We generated four alternative sets of simulated catalogues, referred to as Profiles 1, 2, 3 and 4, each set mimicking a particular cancer: breast cancer, lung adenocarcinoma, B-cell lymphoma and melanoma. In order to do so, for each tumour type, we applied MutationalCone to the corresponding TCGA datasets and we calculated the mean contribution across all samples of each signature known to contribute to the specific cancer type \( q_n \). Signatures with \( q_n = 0 \) do not contribute to the final catalogue and were not in the matrix \( P \). S4 Fig depicts the resulting four sets of configurations \( (q_1, \ldots, q_N) \) used for the simulations. Profiles 3 and 4 are characterised by one dominant signature, Profiles 2 by two signatures with similar large contributions and Profile 1 by several signatures with small effects. We fixed the relative frequency of structural zero contributions to the catalogues to \( \pi = 0.6 \) in all simulations. We chose this value because it leads to a small number of hypermutated catalogues, as it is often encountered in practice. Finally, we varied the number of \( r \) from as little as 10 to as much as 100,000 mutations. This allowed us to study the performance of methods on a large spectrum of catalogues:
from a limited number of mutations as in exomes, to a very large number, as in whole cancer genome sequences.

For each of the four tumour types and for each value of \( r \), we simulated a catalogue matrix with \( G \) samples. Fig 2 shows examples with \( G = 30 \) for three values of \( r \).

### Methods for measuring the performance of algorithms

All methods for identifying signatures find solutions to the minimization problem (2). A straightforward way to measure the accuracy of the reconstructed catalogue is, therefore, to calculate the Frobenius norm of the reconstruction error

\[
||M - \hat{M}||_F^2 = \sum_{g=1}^{G} \sum_{n=1}^{N} (m_{g}^n - \hat{m}_{g}^n)^2,
\]

where \( \hat{M} = \hat{P} \times \hat{E} \) is the matrix of catalogues reconstructed from the estimated signature and exposure matrices. Some of the algorithms involve stochastic steps such as resampling and/or random draws of initial parameters. For these algorithms, one simple way to assess the robustness of the estimates is to look at the variability of the reconstruction error when the same catalogues are analysed several times with the same algorithm.

With regards to bayesNMF, it is known that the performance of its principal function might be poor in presence of hypermutated catalogues that mask the detection of signals from less mutated catalogues [17]. For this reason, we pre-treated the catalogues to be analysed by this tool and replaced hypermutated catalogues by synthetic non-hypermutated catalogues to maintain the original mutational distribution catalogues using the standalone get.lego96.hyper function that can be found in the bayesNMF script.

In order to make decisions about whether an extracted signature is the same as validated signatures (e.g. COSMIC signatures) a cut-off for cosine similarity needs to be defined. We decided to apply six different cut-offs and consider as “new” all identified mutational signatures for which the maximal cosine similarity is lower than the cut-off value.

### Specificity and sensitivity for de novo extraction and assignment

In most applications, signature extraction is done in two steps: first, signatures are found using a de novo extraction tool and then for the extracted signatures a cosine similarity with each of the COSMIC signatures is calculated. In order to measure the performance of both these steps combined, we used simulated catalogues and computed false and true positive rates and false and true negative rates. In a simulated catalogue, the set of true signatures \( p_1, \ldots, p_N \) that do contribute to the catalogue are known, thus allowing the comparison of the latter to the estimated signatures \( \hat{p}_1, \ldots, \hat{p}_N \). Note that, for the sake of simplicity, we explicitly look for the same number of signatures that we used to simulate the catalogues, not addressing questions about model selection performance.

Estimated signatures that belong to the set of “true signatures” are considered as true positives, while all “true signatures” that are not extracted count as false negatives. False positives are all estimated signatures that do not have a match in the set of “true signatures”. This can happen for two reasons: the estimated signature is assigned to a COSMIC signature not used to build the catalogue, or it is not sufficiently similar to any COSMIC signature. This last situation usually takes place when setting a very high cosine similarity threshold \( h \). In this case, signatures that have maximal cosine similarity lower than the cutoff, will be termed as “new”. Finally, true negatives are all COSMIC signatures not used for the simulation, nor estimated. From these four measures, we compute specificity (number of true negatives divided by the total number of negatives) and sensitivity (number of true positives divided by the total number of positives).
Fig 2. Sample of simulated catalogues. Catalogues are simulated according to the model described in the text. Tuning parameters are the average number of mutations, here $r = 10^2, 10^4$ and $10^5$, and the cancer type-specific average contribution of each signature (also see S4 Fig). Coloured bars indicate the contribution of each signature to the total mutational load of each sequence.

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In our empirical study, for each simulation setting described in the Simulated data section (that is for each profile given by a choice of proportions \((q_1, \ldots, q_N)\) and for a choice of total number of mutations \(r\)) we built 50 replicates, each made of a matrix of \(G\) samples. Then we extracted signatures from all replicates with a given tool. Importantly, we set the expected number of signatures to be the same as the number of signatures used in the simulation, thus not addressing the problem of model selection. Then we compared the extracted signatures to the COSMIC signatures using a cosine similarity threshold \(h\). Finally, we computed specificity and sensitivity and obtained Monte-Carlo estimates based on the means over all replicates.

**Bias of refitting procedures.** Refitting algorithms assume that the matrix of signatures is known, and return the exposure estimates \(\hat{e}^n_g\), i.e. estimates of the contribution of each signature \(e^n_g\). A simple way to assess the performance of the refitting method is then to look at the bias of such estimates, by comparing them to the true exposures. In order to do so, we simulated 50 replicates each consisting of one lung adenocarcinoma-mimicking catalogue \(g\) (Profile 3) with an average number of mutations set to \(r = 10^4\). Then, for each process \(n\), we obtained Monte-Carlo estimates of the bias \(E[\hat{e}^n_g - e^g_n]\) by averaging the differences \(\hat{e}^n_g - e^g_n\) over all replicates. A global measure of performance that considers all exposure estimates is given by the mean squared error (MSE), that is the expected value of the loss function \(\sum_{g=1}^{30} (\hat{e}^n_g - e^g_n)^2\). We obtained Monte-Carlo estimates of the mean squared error by averaging the loss function values across all 50 replicates and calculate asymptotic confidence intervals.

Analyses were performed using R 3.5.1 on a Linux Ubuntu (14.04) machine with 32 GB RAM and Intel Xeon E5-2630 v4 CPUs @ 2.2 GHz x 16.

**Comparisons of algorithm performance**

**Performance of de novo tools**

**Fig 3** shows the distribution of the reconstruction error when a given computational tool is applied several times to the same real trinucleotide matrix. Reconstruction errors show limited variability due to stochastic steps in the algorithms and no variability whatsoever for maftools. All methods under evaluation are roughly equivalent in terms of their ability to properly reconstruct the initial matrix of mutational catalogues. This is not surprising, given that all methods are meant to solve the optimization problem in (2).

In general, the error value appears to depend on the cancer dataset. This is expected because the four datasets differ with regards to the number of samples, their total number of mutations and the number of operating mutational signatures, making the decomposition more or less difficult.

In order to study the robustness to the presence of outliers, we also estimated the reconstruction error of each method (not only bayesNMF) after pre-treating all datasets with the function get.lego96.hyper. The results reported in S5 Fig show that the performance of each method improves after pre-treating the samples, especially for Melanoma and Breast cancer datasets that are characterised by a few samples with an extremely high number of mutations. For the Melanoma dataset, the gain in performance is considerable for bayesNMF and maftools.

We used realistic simulations to evaluate the performance of each method for de novo extraction followed by a classification step in which the extracted signatures are assigned to the most similar COSMIC signature.

**Figs 4 and 5**, respectively show the specificity and sensitivity of such two-stage procedure as functions of the number of samples \(G\) in each catalogue and the cosine similarity cut-off \(h\),
while Figs 6 and 7 show the specificity and sensitivity as functions of the number of mutations in each catalogue and \( h \).
We do not see very large differences in the tools’ specificity with respect to the number of samples (Fig 4). For Profiles 2, 3 and 4 the specificity of all methods is close to 1 even for small sample sizes, while for Profile 1, that is characterised by small contributions from several signatures (see S4 Fig), the specificity is close to 1 starting from 50 samples. The sensitivity of most of the algorithms increases with the sample size (Fig 5) and this trend is more evident for Profile 1. Methods based on NMF (maftools, SomaticSignatures, mutSignatures) have lower sensitivity, while methods based on probabilistic models perform better, with the notable exception of signeR. Most of the differences between tools are observed for Profile 4, with some methods
(EMu, bayesNMF, pmsignature) having a sensitivity close to 1 and the others having lower and more variable sensitivities.

Specificity increases with the average number of mutations (Fig 6). For Profiles 2, 3 and 4 it is close to 1 starting from as low as 1000 mutations, while for Profile 1 it is only for at least 10,000 mutations that we observe a specificity close to 1 for most of the methods, with the notable exception of bayesNMF that performs well even for lower numbers of mutations. Sensitivity increases with the average number of mutations with a large variability according to the cancer profile and method (Fig 7). Sensitivity is high for cancer profiles characterised by one predominant signature (Profiles 3 and 4) or two strong signatures (Profile 2) but may vary.
become relatively low for datasets characterized by small contributions by several signatures (Profiles 1). This indicates that signatures that act together with other signatures and have small effects may be more difficult to identify.

Specificity and sensitivity slightly deteriorate for higher cut-off values. This is expected because by setting a higher cut-off, the number of found signatures that are not similar enough to COSMIC signatures increases. Because these estimated signatures are considered as novel, they are false positives (that is found signatures not used for simulations), leading to a greater number of false positives and therefore to a lower specificity. Moreover, if the cut-off is too stringent, the number of false negatives will be high because some signatures used for the

Fig 6. Simulation study: Specificity of extraction methods and mapping on COSMIC signatures as the average number of mutations and the cosine cut-off \( h \) vary. Specificity is estimated from 50 replicates each made of \( G = 30 \) catalogues. The model used to simulate realistic replicates according to the four Profiles and the estimation methods are described in the section Data and experimental settings.

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simulations are correctly found but do not score a high enough cosine similarity and therefore count as false negatives. This will make the resulting sensitivity low.

Fig 8 shows the running time when tools are applied to real lung datasets with a varying number of samples. While all methods show a fast-growing running time with increasing number of samples, SomaticSignatures and maftools are much faster than the others for more than 100 samples, making it possible to analyse large number of samples in few seconds. For example, for two hundred samples, the slowest method (signeR), the running time is 913.72s while for the fastest (maftools), the value is 5.97s.
Performance of refitting tools

Fig 9 shows the distribution of the differences between the estimated and true contribution of all \( n \) signatures \( e_1^g, \ldots, e_30^g \) for the different refitting methods under evaluation. Sample catalogues were simulated mimicking Lung cancer profiles (Profile 3), with signatures 1, 2, 4, 5, 6, 13 and 17 actually contributing as shown in S4 Fig. All methods give almost identical results.

By comparison with the true exposure profile given in S4 Fig, it is clear that all refitting methods provide good estimates of the contributions of all but signatures 4, 5 and 17 and to a lesser extent signature 6. Moreover, all methods correctly estimate a zero contribution for signatures 3 and 16 even though these are very similar to signature 5, see S2 Fig.

Interestingly, we note that signatures 2 and 13 (both attributed to APOBEC activity) are in general well identified by all methods. This finding is in line with previous claims about the stability of these two signatures.

In terms of running time, deconstructSigs and SignatureEstimation based on simulated annealing are more than two orders of magnitude slower than the other methods (Fig 10). All other methods run in a fraction of second. As expected, the running time increases linearly with the number of samples. MutationalCone, our custom implementation of the solution to the optimization problem solved by YAPSA and MutationalPatterns outperforms all other methods. The second fastest method is SignatureEstimation based on Quadratic Programming. As example, for two hundred samples, the execution time of deconstructSigs is 86.148s and for MutationalCone is 0.028s.

Discussion

In this work we complement and expand a recent review of the available methods to identify mutational signatures [7] by empirically comparing their performance, using both real world TCGA data and simulated data. To our knowledge, this is the first formal comparison of both de novo discovery and refitting of mutational signatures. The research about mutational signatures is very active and in rapid development, with preprints about new methods regularly...

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Since the publication of the first work about mutational signatures in 2013 [2], multiple algorithms have been developed, leading to similar but not identical results, a source of concern for researchers interested in this type of analysis. Conceptually, this is not surprising: mutational signatures are naturally defined in terms of non-negative matrix factorization, a well-known ill-posed problem (a unique solution does not exist). Although this limitation has cast doubts on the biological validity of mutational signatures, this has been somehow validated using experimental and computational approaches by Zou et al. [36]. Similarly, the term “signature” itself is somehow controversial as it implies a unique correspondence with mutagens, while in this context signatures are mathematically defined as basic signals each contributing to the final sequence mutational burden. Sufficiently detailed tumour catalogues and mutagen spectra might yield patterns that are unique to a tumour type or mutagen, and therefore become “true” signatures that allow backward inference from the tumour to the mutagen. Mutational signatures data in combination with epidemiological information may provide useful insights to identify the causes of cancer [37,38]. The utility of the current models of substitution mutational signatures is also shown in a recent experimental work based on a human induced pluripotent stem cell (iPSC) line that provides evidence for the possibility to identify the agents responsible for some specific mutational signatures [39]. In such work, Kucab and...
colleagues compared iPSCs treated and untreated with 79 known or suspected environmental carcinogens and identified specific substitution mutational signatures for around half of such carcinogens. Some of such signatures were similar to those identified in human tumour DNA.

An original aspect of this paper is the empirical performance study and in particular the probabilistic model of mutational catalogues used to generate simulations. This model is based on the zero-inflated Poisson distribution that allows for sparse contribution of signatures and thus makes it possible to build mutation count data that are more realistic than the pure Poisson model previously considered [9,14]. As argued in the Simulated data section, our model underrepresents the few samples with extremely large total mutation counts. Because catalogues of this type might hamper the detection of signals from less mutated samples in the same dataset [7], it is likely that our results slightly overestimate the methods performance in the presence of hypermutated samples. However, our main objective was the comparison of the different methods, and this is not affected by this systematic bias. In our model, a larger number of hypermutated catalogues could be obtained by lowering the value of \( \pi \), the parameter that controls the relative frequency of structural zeroes in the zero-inflated Poisson model. As discussed, it would have been possible to consider even more realistic models, however these would have led to results that depend on too many parameters thus making the interpretation harder. For instance, the zero inflated negative binomial model is a more flexible model and looks a promising method to build realistic synthetic samples, including hypermutated ones. We leave this interesting perspective to future work. Alternative models that were recently proposed are based on the negative binomial distribution [22] and on the Dirichlet distributions for the exposures and signatures and the multinomial distribution for the catalogues [29]. We suggest that developers should assess their new methods on simulations based on realistic models such as ours or the latter. The advantage of simulations over real data is that the underlying model generating the synthetic data is known and can be compared to the estimation provided by the method being evaluated. For this reason, we decided not to simulate catalogues from real data using the bootstrap: this would have produced almost real samples but without the possibility to evaluate the performance of methods according to different parametric scenarios. We strongly believe that the mutational signature research could benefit from the development of public realistic datasets that can be used to benchmark old and new detection tools.

Another innovative aspect of our work is the assessment of the performance of de novo approaches. This is in general a slippery problem given the non-supervised learning nature of NMF and concurrent methods. We addressed it by building confusion matrices after comparing the “true” signatures used for simulations with the newly identified signatures in terms of their cosine similarity.

Finally, we introduced MutationalCone, a new implementation of existing refitting tools that is based on a very simple geometric model and proved to be faster than all other methods. The R code implementing MutationalCone can be found in the S1 File.

The results of our study can be helpful to guide researchers through the planning of mutational signature analysis. In particular, we show that the performance of de novo methods depends on the complexity of the analysed sequences, the number of mutations and to a lesser degree the number of samples analysed. It was somehow expected that the performance of the methods for a cancer in which multiple, concomitant, signatures are present is poorer than for a cancer with a single or predominant signature, particularly when the concomitant signatures are similar and have a low contribution. However, we demonstrated that in the latter case it is mainly sensitivity and not as much specificity that significantly decreases. An intuitive reason for this result is that a signature with low impact is difficult to detect and therefore will be wrongly considered as a “negative”; several such signatures will then imply a large number of
false negatives, i.e. low sensitivity. Indeed, recent evidence shows that the majority of cancers harbour a large number of mutational signatures [4] and therefore belong to the latter scenario.

With regards to the mutation number, we observe that with the number of mutations that could be found in some cancer exomes the performance is generally poor (i.e. low specificity and sensitivity). This problem is likely to be mitigated if counts were normalized by the expected number of each type’s trinucleotides in the analysed region under healthy condition, that is if an opportunity matrix was provided. We do not address this important aspect in our comparison study as only a few methods can incorporate opportunity matrices.

We showed that when comparing identified signatures with COSMIC signatures, the choice of a cosine similarity cut-off has a relatively small impact on the overall performance. If the aim is to identify novel signatures it would be preferable to choose a lower value (0.75 or less). On the contrary, if the aim is to assess the presence of known signatures in mutational catalogues (cancer genomes or exomes), we recommend turning to refitting methods. For well-studied cancers, refitting approaches are a faster and more powerful alternative to de novo methods, even with just one input sample. As the COSMIC database has been built and validated by analysing tens of thousands of sequences of most cancer types, we recommend borrowing strength from previous studies and using refitting tools when performing standard analysis not aimed at the discovery of new signatures.

Our simulation study seems to indicate that de novo probabilistic methods EMu and bayesNMF have an overall better performance as they achieve better sensitivity and specificity with a fair running time. However, in order to assess the robustness of results, due to the variability of outcomes and the presence of hypermutated samples, we recommend to systematically perform a sensitivity analysis based on the application of one or more alternative methods based on different algorithms.

Our analysis also reveals that if the dataset under consideration contains catalogues with a very large number of mutations, all methods achieve better performance by replacing such outliers with the bayesNMF pre-treatment function get.lego96.hyper. Interestingly, the mutation profiles of the synthetic datasets simulated with the ZIP model resemble the profiles of datasets after such pre-treatment, see S6 Fig.

Not all the de novo methods we evaluated offer the possibility to automatically choose the number of signatures to be found. For instance, the popular SomaticSignatures only provides a graphical visualisation of the residual sum of squares for several choices of the number of signatures; the user can choose the optimal number by identifying the inflexion point. For this reason, we did not address this crucial aspect in our empirical assessment. Similarly, we only considered mutation types defined by the trinucleotide motifs, as currently only pmsignature [16] can consider more than one flanking base on each side of the substitution.

A more recent set of 49 signatures have been estimated from a new large dataset including thousands of additional sequences. The article introducing this new set has not been published yet but the new set of signatures is available on the COSMIC website. In the future it will be interesting to evaluate the performance of the different tools using the new set of signatures.

**Supporting information**

**S1 Fig. Comparison of newly identified signatures with COSMIC signatures.** Signatures a-g were identified in a de novo extraction using the maftools R package from the TCGA Lung Adenocarcinoma cohort of 563 cancer genomes. The novel signatures were then compared to the 30 signatures validated in the COSMIC database in terms of cosine similarity. Each signature is then assigned to the most similar COSMIC signature provided that their cosine

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similarity is above a fixed threshold. For instance, signature f is matched to signature 5 at a cut-off of 0.75 but is considered as a completely new signature if the cut-off is at 0.80. Also note that a unique assignment can be controversial: for instance, signature g is similar both to signatures 12 and 26.

(TIFF)

S2 Fig. Cosine similarity plot of COSMIC signatures. Some COSMIC signatures are very similar to others. For instance, signature 8 is similar to signatures 3, 4 and 5.

(TIFF)

S3 Fig. Simulations of 563 lung adenocarcinoma catalogues according to different models. Part a: Real catalogues from the TCGA lung adenocarcinoma cohort. Parts b and c: Catalogues sampled from the ZIP model described in the main text. The relative contribution $q_n$ of each signature $n$ is the mean of the relative contributions of $n$ in all samples as estimated by maftools. In part b simulated and real catalogues are in a 1 to 1 correspondence: for each simulated sample $g$, the total number of mutations $r_g$ in the corresponding real catalogue is taken. In part c all samples are simulated according to $r = 306$, the average total number of mutations in the real data. The latter example illustrates the parametric model used for the simulation study. Part d: Catalogues sampled according to the Poisson model $e_n^{r_g} \sim P(\lambda_n)$, where $\lambda_n$ is the mean number of mutations due to $n$ in the real samples as estimated by maftools.

(TIFF)

S4 Fig. Choice of parameters $q_n$ in the simulations. Four different configurations ($q_1, \ldots, q_{30}$) were considered for simulating realistic data. Each configuration represents the average share of mutations due to the different COSMIC signatures and was chosen to mimic real exposure profiles for four cancer types: estimates were obtained from Breast Cancer (Profile 1), Lymphoma (Profile 2), Lung Adenocarcinoma (Profile 3) and Melanoma (Profile 4) TCGA cohorts.

(TIFF)

S5 Fig. Reconstruction errors and their variability: Effect of pre-treating the samples to moderate the effects of hypermutated catalogues. Each dataset was analysed with or without data pre-treatment with the bayesNMF get.lego96.hyper function. The profiles of the pre-treated catalogues are shown in S6 Fig.

(TIFF)

S6 Fig. Barplot with the number of mutations in each sample after pre-treatment to moderate the effects of hypermutated catalogues. Datasets represented in Fig 1 were pre-treated using the bayesNMF function get.lego96.hyper.

(TIFF)

S1 File. Mutational cone. Description of our original method for signature refitting and the corresponding R script.

(DOCX)

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