Do you cov me? Effect of coverage reduction on species identification and genome reconstruction in complex biological matrices by metagenome shotgun high-throughput sequencing [version 1; peer review: 2 not approved]

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Abstract
Shotgun metagenomics sequencing is a powerful tool for the characterization of complex biological matrices, enabling analysis of prokaryotic and eukaryotic organisms in a single experiment, with the possibility of de novo reconstruction of the whole metagenome or a set of genes of interest. One of the main factors limiting the use of shotgun metagenomics on wide scale projects is the high cost associated with the approach. However, we demonstrate that—for some applications—it is possible to use shallow shotgun metagenomics to characterize complex biological matrices while reducing costs. Here we compared the results obtained on full size, real datasets with results obtained by randomly extracting a fixed number of reads. The main statistics that were compared are alpha diversity estimates, species abundance, and ability of reconstructing the metagenome in terms of length and completeness. Our results show that a classification of the communities present in a complex matrix can be accurately performed even using very low number of reads. With samples of 100,000 reads, the alpha diversity estimates were in most cases comparable to those obtained with the full sample, and the estimation of the abundance of all the present species was in excellent agreement with those obtained with the full sample. On the contrary, any task involving the reconstruction of the metagenome performed poorly, even with the largest simulated subsample (1M reads). The length of the reconstructed assembly was sensibly smaller than the length obtained with the full dataset, and...
the proportion of conserved genes that were identified in the metagenome was drastically reduced compared to the full sample. Shallow shotgun metagenomics can be a useful tool to describe the structure of complex matrices, but it is not adequate to reconstruct de novo—even partially—the metagenome.

**Keywords**
high-throughput sequencing, metagenome, metagenomics, next generation sequencing, alpha diversity, complex matrices
Introduction

Shotgun metagenomics offers the possibility to assess the complete taxonomic composition of biological matrices and to estimate the relative abundances of each species in an unbiased way\(^1,2\). It allows for agnostic characterization of complex communities containing eukaryotes, fungi, bacteria, and also viruses, using both DNA and RNA as a starting material. In addition, the whole metagenome approach can be used not only to simply identify DNA and RNA virus in a complex matrix, but also to study the genetic diversity in virus populations\(^3,4\), and to identify potential adventitious agents in biopharmaceutical manufacturing\(^5\).

Metagenome shotgun high-throughput sequencing has progressively gained popularity in parallel with the advancing of next-generation sequencing technologies\(^6,7\), which provide more data in less time at a lower cost than previous sequencing techniques. This allows the extensive application to study the most various biological mixtures such as environmental samples\(^8,9\), gut samples\(^10–13\), skin samples\(^15\), clinical samples for diagnostics and surveillance purposes\(^16–19\), food ecosystems\(^20,22\) and drugs manufactured using biological sources as vaccines\(^22,23\). The aim of whole metagenome approaches is not only to study the taxonomic composition of biological substrates but also to identify which genes and metabolic pathways are present with the aim to understand functional capacities in the studied microbiota\(^24\). Recently the approach has been also used to analyze the ensemble of genes that may encode antibiotic resistance in various microbial ecosystems (i.e. soil), which are defined as the resistome\(^24\).

Another, more traditional approach currently used to assign taxonomy to DNA sequences is based on the sequencing of target conserved regions. Metabarcoding method relies on conserved sequences to characterize communities of complex matrices. These include the highly variable region of 16S rRNA gene in bacteria\(^2\), the nuclear ribosomal internal transcribed spacer (ITS) region for fungi\(^2\), 18S rRNA gene in eukaryotes\(^24\), cytochrome c oxidase sub-unit I (COI or cox1) for taxonomical identification of animals\(^9\), rbcl, matK and ITS2 as the plant barcode\(^9\). Considering the large amount of genetic diversity within and between virus families, a universal metabarcoding approach is not applicable to detect virus nucleic acids in complex biological samples.

The selection of conserved regions has the advantage of reducing sequencing needs, since it does not require sequencing of the full genome, just a small region. On the other hand, given the currently used approaches, characterization of microbial and eukaryotic communities requires different primers and library preparations\(^32\). In addition, several studies suggested that whole shotgun metagenome sequencing is more effective in the characterization of metagenomics samples compared to target amplicon approaches, with the additional capability of providing functional information regarding the studied sample\(^33\).

Current whole shotgun metagenome experiments are performed obtaining several million reads\(^30,31\). However, obtaining a broad characterization of the relative abundance of different species, might easily be achieved with lower number of reads.

To test this hypothesis, we performed sequencing using whole metagenomics approach of seven samples derived from different complex matrices to characterize their composition, and subsequently tested the accuracy of several measures when downsampling the number of reads used for analysis including the performance of de novo assembly in the ability to reconstruct both entire genomes and genes.

Methods

Samples description and DNA extraction

The following samples were used in the present work: two samples collected from a live attenuated virus vaccine (B1 and B2), two horse fecal samples (F1 and F2), and three food samples (M1, M2, and M3). Biological medicines were two different lots of live attenuated MPRV vaccine (Prorix Tetra, Glaxo SmithKline) widely used for immunisation against measles, mumps, rubella and chickenpox in infants. Lyophilised vaccines were resuspended in 500 μl sterile water for injection and DNA extracted using Maxwell® 16 Instrument and the Maxwell® 16 Tissue DNA Purification Kit (Promega, Madison, WI, USA) according to the manufacturer’s instructions.

Horse feces from two individuals were collected as follows: 100 mg of starting material stored in 70% ethanol were processed for DNA extraction using the QIAamp PowerFecal DNA Kit (QIAGEN GmbH, Hilden, Germany), according to the manufacturer’s instructions.

Food samples were raw materials of animal and plant origin, used to industrially prepare bouillon cubes. DNA extractions from those three samples were performed starting from 2 grams of material each, using the DNeasy mericon Food Kit (QIAGEN GmbH, Hilden, Germany), according to the manufacturer’s instructions.

DNA purity and concentration were estimated using a NanoDrop Spectrophotometer (NanoDrop Technologies Inc., Wilmington, DE, USA) and Qubit 2.0 fluorometer (Invitrogen, Carlsbad, CA, USA).

Whole metagenome DNA library construction and sequencing

DNA library preparations were performed according to manufacturer’s protocol, using the kit Ovation® Ultralow System V4 1–96 (Nugen, San Carlos, CA). Library prep monitoring and validation were performed both by Qubit 2.0 fluorometer (Invitrogen, Carlsbad, CA, USA) and Agilent 2100 Bioanalyzer DNA High Sensitivity Analysis kit (Agilent Technologies, Santa Clara, CA).

Cluster generation, template hybridization, isothermal amplification, linearization, blocking and denaturization and hybridization of the sequencing primers was then performed on Illumina cBot and flowcell HiSeq SBS V4 250 cycle kit, loaded on
HiSeq2500 Illumina sequencer producing 125bp paired-end reads (for samples B1, B2, M1, M2 and M3) and 250bp paired-end reads (for samples F1 and F2).

The CASAVA Illumina Pipeline version 1.8.2 was used for base-calling and de-multiplexing. Adapters were masked using cutadapt\textsuperscript{34}. Masked and low quality bases were filtered using erne-filter version 1.4.6\textsuperscript{35}.

Bioinformatics analysis
The bioinformatics analysis performed in the present work are summarized in Figure 1.

Different read lengths among samples may constitute an additional confounder in analysis. To obtain homogeneous read length across samples, reads sequenced belonging to F1 and F2 were trimmed to a length of 125 bp using fastx-toolkit version 0.0.13 before subsequent analysis.

Reduction in coverage was simulated by randomly sampling a fixed number of reads from the full set of reads. Subsamples of 10,000, 25,000, 50,000, 100,000, 250,000, 500,000 and 1,000,000 reads were extracted from the raw reads using seqtk version 1.3. To assess variability, subsampling was performed 5 times for each sample and for each read abundance.

Classification of reads against the NCBI nt database downloaded on May 2018 was performed using Kraken 2 version 2.0.6-beta\textsuperscript{36} to estimate species abundance and Shannon diversity index. A simplified representation of species composition was obtained using Krona version 2.6\textsuperscript{37}.

Chao1\textsuperscript{38} species richness and Shannon’s diversity\textsuperscript{39} were estimated using the R package vegan version 2.4.2\textsuperscript{40}.

Assembly of the metagenome was performed using megahit version 1.1.2\textsuperscript{41}. Completeness of the assembly was assessed using BUSCO version 3.0.2\textsuperscript{42}. The proportion of the reconstructed genes was measured as the proportion of genes that were fully reconstructed, plus the proportion of genes that were partially reconstructed. BUSCO analysis was performed on prokaryotic database for all the samples with the exception of M1 (mostly composed by fungi) for which the fungal database was used. Samples B1 and B2 were also compared against the eukaryotic BUSCO database; results for the prokaryotic database are reported.

Unless otherwise specified, all the analysis were performed using R 3.3.3\textsuperscript{43}.

Results
Sample composition and downsampling
Summary statistics for the full samples included in the study are shown in Table 1.

The number of reads obtained in the samples selected for the present study ranged from slightly more than 1 million

![Figure 1. Workflow of the main bioinformatics analysis performed in the present work.](image-url)
Table 1. Summary statistics for the full samples included in the study.

| Sample | N reads   | N species | Singletons |
|--------|-----------|-----------|------------|
| B1     | 11,031,061| 2508      | 1299       |
| B2     | 3,830,083 | 4598      | 1795       |
| F1     | 12,472,553| 29661     | 14750      |
| F2     | 10,780,450| 25608     | 12374      |
| M1     | 1,898,011 | 3207      | 1469       |
| M2     | 1,558,975 | 9638      | 3377       |
| M3     | 1,867,879 | 5567      | 1999       |

N species, number of species identified in the sample including species identified by one or more reads; Singletons, number of species identified in the sample by only one read.

(sampling M2) to more than 12 million (sample F1). Our subsampling, ranging from 10,000 to 1,000,000 reads, led to a reduction in size of 36% (1,000,000 out of 1,558,975) in M2 to 0.08% of the original size (10,000 reads out of 12,472,553) in F1.

Samples used in this study had different levels of species composition (Figure 2). Some samples, such as M1, B1 and B2 were dominated by a single species, while others, in particular fecal samples, showed high heterogeneity in species composition.

Diversity and species richness

Figure 3 shows the variation of the value of Chao1 estimator, representing the estimated number of species in each sample when varying the number of reads used for the estimation, from the smallest number on the left, to the full dataset on the right. The value of Chao1 estimator for the full dataset is plotted on the right side of the plot, at the rightmost fecal samples F1 and F2 had an estimated number of species greater than 40,000, much higher than all the other samples, for which less than 20,000 species were estimated (less than 10,000 B1, B2, M1 and M3).

The effect of downsampling on the estimated number of species has different effects in different samples. For most samples, even a robust downsampling led to only a slight reduction in the estimated species richness. However, for samples F1 and F2, which were characterized by a high number of overall species and rare species, the downsampling led to a significant reduction in the estimated species richness.

Shannon’s diversity index is a widely used method to assess the biological diversity of ecological and microbiological communities. Figure 4 depicts the effect of subsampling on the Shannon’s diversity index. The effect of subsampling on Shannon’s diversity index is smaller than the effect on the estimated number of species. The variation in Shannon diversity index with subsampling is negligible for all samples, even reducing the number of reads from the full size to 100,000 or less.

Figure 5 shows the correlation in species abundance estimation between the full dataset and a reduced dataset of 100,000 reads. The linear correlation coefficient between the two datasets is >0.99 in all the replicates. The plot is in log-log scale to emphasize differences in low abundance species. Only species with frequencies lower than 0.01% (i.e. species represented in 1 read out of 10,000) show some effect of subsampling on the relative abundance estimation. All the seven samples share a similar behavior.

In Figure 6 we show the results obtained by reducing the number of sampled reads to 10,000 reads per sample. Similar to what we observed for larger subsamples, the linear correlation coefficient between species abundance estimate in the full and the reduced dataset was high in all the samples (r>0.95) and in all the replicated subsampling. The abundance of species with frequency greater than 1/1000 (0.1%) is correctly estimated in the subsamples, while for rare species the estimate is not precise. Species with frequencies <0.01% are by definition absent in the subsample obtained with 10,000 reads, and were arbitrarily set to a frequency of 0.001% to provide the reader with an idea on their abundance and distribution in the original sample.

Metagenome reconstruction

While characterizing and measuring species present in a complex matrix is an important task, some studies aim at reconstructing (partially or entirely) the metagenome via a de novo approach. We thus investigated the effect of coverage reduction on this task. We reconstructed de novo the metagenome of the full and reduced datasets, and compared the reconstructed genomes. Results are summarized in Figure 7. As expected, the size of the assembly is strongly influenced by the read number. Assemblies obtained using the full set of reads had a size ranging from slightly more than 1 Mb (sample B1) to nearly 100 Mb (F1 and F2). A decrease in the number of reads used for the assembly lead to a steady decrease in assembly size in all samples, although with different slopes. Assembly sizes obtained using 1,000,000 reads ranged from less than 1 Mb (F1 and F2) to slightly more than 10 Mb (M1), and those obtained using 100,000 reads ranged from less than 100 Kb (F1 and F2) to less than 1 Mb (all the remaining samples).

However, the total assembly length is not necessarily a sufficient measure to describe assembly goodness and completeness. Since we are interested in assessing the completeness of the reconstructed metagenome, we used BUSCO to report the proportion of genes covered by any given assembly. Figure 8 reports the proportion of metagenome completeness estimated by BUSCO from full and from the reduced dataset obtained by randomly sampling 1,000,000 reads. The prokaryotic BUSCO dataset was used for all samples with the exception of sample M1, composed prevalently by a mushroom, for which the fungal BUSCO database was used. The full samples F1 and F2 reconstructed a fairly complete proportion of the BUSCO genes (>90%), while the reduced dataset reconstructed less than 20%.

Similar trends can be observed with other datasets. Given the lower number of reads sequenced in other samples, the performance in reconstructing the BUSCO genes was generally poor, but reducing to 1 million reads led to a further decrease in performance, suggesting that this is a clearly suboptimal number of reads. Samples B1 and B2 show a very poor performance.
Figure 2. Graphical representation of the composition of the seven studied samples.
because the prokaryotic organisms in the sample are very rare contaminants. Being derived from fetal human cell cultures, a large portion of the metagenome is constituted by human sequences, but given the very small ability in reconstructing de novo a genome as large as the human one, the proportion of reconstructed BUSCO genes is very low (<5% both for prokaryotic and eukaryotic BUSCO genes).

Discussion
The aim of the present work was to assess the reliability of low-depth shotgun metagenome sequencing for the characterization of complex matrices, as follows: 1) determining diversity and species richness in complex matrices; 2) estimating abundance of the species present in the complex matrix, and 3) reconstructing de novo the genome of the species present in the samples.
Figure 5. Scatterplot of species abundance estimated using the full set of reads and a set composed of 100,000 reads. Data for all the five replicates of the subsampling are plotted. Each point (colored by sample of origin) represents a given species. The position on the X axis represents the relative abundance of the species in the full dataset, and the position on the Y axis represents the relative abundance of the species in the samples obtained by randomly sampling 100,000 reads. Both axis are plotted in log scale to facilitate visualization of low abundance species.

Figure 6. Scatterplot of species abundance estimated using the full dataset of reads and a dataset composed of 10,000 reads. Data for all the five replicates of the subsampling are plotted. Each point (colored by sample of origin) represents a given species. The position on the X axis represents the relative abundance of the species in the full dataset, and the position on the Y axis represents the relative abundance of the species in the samples obtained by randomly sampling 10,000 reads. Both axis are plotted in log scale to facilitate visualization of low abundance species.
We selected seven heterogeneous complex samples, sequenced at varying coverage (ranging 1 to 12 million reads). Shotgun metagenomics experiments—often aiming at reconstructing de novo the studied metagenome—have a tendency to generate a very high number of reads per sample. Compared to such studies, all of our samples have relatively shallow coverage of the metagenome, and we tested if even lower coverage could still provide reliable answers to the three main questions listed above.

We used Chao1 as an indicator of species richness and Shannon’s diversity index as an estimator of species diversity, and we measured their variation when reducing the number of reads used for the experiment.

An important detail to be considered here is the fact that the two indices behave differently in the full and the reduced samples. We provide an explanation regarding the reasons of this difference.

Chao1 estimator is obtained as

\[ S_{\text{Chao1}} = S_{\text{obs}} + \frac{f_1(f_1 - 1)}{2(f_2 + 1)} \]

Figure 7. Total length of the de novo metagenome assembly in each sample as a function of the number of reads. X and Y axes are in log scale. Shaded areas represent the confidence limits of resampling experiments. “Full” represents the values obtained with the full set of reads (number of reads per sample listed in column 2 of Table 1).

Figure 8. Completeness of the BUSCO genes in the full dataset (darker colors) and in the largest of the reduced datasets (lighter colors). Error bars are based on the five replicate experiments performed for each sample.
Where \( S_{obs} \) is the number of observed species in the sample, \( f_j \) is the number of species observed once, and \( f_2 \) is the number of species observed twice.

Shannon diversity index is estimated as

\[
H = -\sum_{i=1}^{N} p_i \ln(p_i)
\]

Where \( N \) is the total number of species and \( p_i \) is the frequency of the species \( i \).

Thus, the Chao1 index is heavily affected by the number of rare species that are identified and not from the relative frequencies of the most abundant species, while the Shannon diversity index is affected more by variation in the frequencies of highly abundant species than by the disappearance of rare species.

Samples F1 and F2 are characterized by a very large number of observed species (29,661 and 25,608, respectively), while all the other samples have lower number of species, ranging from 2508 in B1 to 9638 in M2. Chao1 captures this differences, showing that F1 and F2 have greater diversity estimates. The Shannon diversity index, on the contrary, relies not only on the number of observed species, but also on the frequency distribution, and for a given number of species reaches its maximum for equifrequent species. Therefore, samples that have a relatively high number of common species with comparable frequencies tend to have high Shannon’s diversity indices.

As an example the number of species with a frequency greater than 0.1% was 23 in sample F1 and was 55 in sample M2. Thus, in spite of a much lower number of species in M2 compared to F1, the Shannon diversity is higher in M2 than in F1. Given the differences in behavior between the two indices in certain conditions, we decided to use both of them to have a more complete information on sample diversity when decreasing coverage. Our results show that a substantial reduction of coverage can be safely achieved without compromising the ability of estimating species richness and abundance (Figure 3 and Figure 4), although the estimated number of species is moderately affected by coverage reduction.

We then set out to assess the changes in the estimated relative frequency of each individual species when reducing the number of sequenced reads. Accurate estimate of the relative abundance of each species is an important task when the aim is a) to detect species with a relative abundance above any given threshold, b) to differentiate two samples based on different abundance of any given species composition, or c) to cluster samples based on their species composition. Our results show that even reducing sequencing to 100,000 reads, species abundances as low as 0.01% can be reliably estimated.

The last questions to which we sought to answer is if a reduction in the sequencing coverage would have a deleterious effect on the ability of de novo assembling the metagenome. Our results show that downsampling had a strongly negative effect on the total length of the reconstructed metagenome and on the proportion of BUSCO genes reconstructed with the metagenome assembly.

BUSCO is widely used for assessing the completeness of genome and transcriptome assemblies for individual organisms, and has benchmark datasets for several lineages. It is possible that using BUSCO for assessing completeness of a metagenomics assembly, including both eukaryotic and prokaryotic organisms, results in an underestimation of the completeness of the reconstruction. However, the aim of the present work is not the absolute estimation of the completeness of the metagenomics assembly, but rather the relative variation observed when using a subsample of reads. Our results indicate that even using 1,000,000 reads is clearly suboptimal in terms of fully sampling the genes present in the complex matrices. This observation needs to be taken into account in the phase of experimental design. Our conclusions also affect research aimed at reconstruction of an interesting part of the meta-genome, such as genes involved in antibiotic resistance\(^7\). The decrease in performance observed in the reconstruction of BUSCO genes will be likely observed for the reconstruction of other gene categories. Researchers aiming at a de novo reconstruction of the metagenome (although partial) must keep in mind that several millions of reads are needed to attain reliable results.

In the present work we tested the feasibility of using metagenome shotgun shallow high-throughput sequencing to analyze complex samples for the presence of eukaryotes, prokaryotes and virus nucleic acids with the aim of monitoring, diagnosis, surveillance, quality control and traceability.

We show that, if the aim of the experiment is a taxonomical characterization of the sample or the identification and quantification of species present in it, then a low-coverage WGS is a good choice. On the other hand, if one of the aims of the study relies on de novo assembly, then a higher number of reads is required. We do not provide here a suggestion on the number of reads that are needed when the aim is the (partial) reconstruction of the meta-genome, as it depends on several factors (number of species in the sample, their genome size, and their abundance, length of the sequencing reads, quality of the DNA) and this estimation needs to be performed for each experiment based on detailed understanding of the experiment aims and of sample characteristics.

**Data availability**

Raw reads are available at NCBI Sequence Read Archive. Samples F1 and F2 are available under accession number SRP163102: https://identifiers.org/insdc.sra/SRP163102; samples B1 and B2 are available under accession number SRP163096: https://identifiers.org/insdc.sra/SRP163096; and samples M1, M2 and M3 are available under accession number SRP163007: https://identifiers.org/insdc.sra/SRP163007.

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Jose F. Cobo Diaz

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The authors proposed and evaluated the influence of reduce sequencing effort (amount of sequences) for a whole metagenome shotgun analysis, using the Illumina platform, in the species composition and diversity index of the communities studied. Although the idea and hypothesis are good, some problems were found in the experimental design and data analysis.

According to the questions proposed in the peer review form, it is not a new method, only the adaptation of a current methodology to optimize the cost and increase the potential numbers of samples analyzed per run of Illumina platform. Although the introduction is clearly explained, the reasons for use shotgun sequencing, mainly to analyze viruses data and functional data for all the organism, no emphasis on such points was done in the results and discussion. The samples used (vaccines, horse fecal samples and food samples) and the introduction remark the detection of pathogens as the main objective of the approach used, including viruses, which can not be screened by amplicons approaches, like metabarcoding sequencing. I suggest adapting the text and manuscript to focus on pathogens (mainly viruses) found along the sub-samples taken for each sample. At that point, some contaminated samples (or not contaminated samples mixed with known amounts DNA from pathogen viruses) have to be used to determine the lowest pathogen concentration that could be detected for each shotgun sequencing coverage proposed.

Many problems were found with the methodology employed, mainly the parameters used in each step and/or software employed for data filtering and analysis, which are critical for the results, which can have strong variations depending of the parameters used. Hence, the methodology proposed does not allow any replication of the method used. Moreover, there are some mistakes for species designation in the study, with at least 2508 species found in vaccine samples indicating big problems along read filtering and data analysis, because this number of species is often found in more complex systems, such as soils samples from agricultural fields. Moreover, go to species classification using some taxonomical markers, such ITS or 16SrRNA, is risky with sequences lower than 400 bp, and sometimes with bigger sequences. In the current manuscript, the use of non
taxonomical marker sequences and 150 bp lengths increase enormously the number of sequences not correctly assigned to species level, and in several cases also for higher taxonomical levels (genus, family...). Therefore, I suggest to clarify how the species assignment was done, because it looks like that each gene-species was considered as one species, and each gene found for a single species was counted as a new species.

Alpha diversity indexes employed are not the best ones, in my opinion, to describe or compare the sub-samples proposed in this manuscript. The chao1 index, an estimator of richness, has a strong influence on the number of singletons obtained in the samples, which due to the complexity of the samples-data tends to be high. Shannon index is influenced by both richness (number of taxa) and evenness (equability, Pielou index), and the reduction of richness due to the loss of rare taxa has a strong influence on this index. I propose to use the number of observed taxa instead of estimated taxa, and any evenness index, like the Pielou index, instead of the Shannon index. Moreover, the use of a coverage index, such Good's coverage index, could be useful to compare the loss of information associated to sampled size or coverage.

In conclusion, although the raw data can contains some important information, the manuscript has to be improved with new “pathogen contaminated” samples, and be re-written to focus on the detection of pathogens in the samples, which due to the low abundance of the samples could not be detected depending of the shotgun coverage.

Is the rationale for developing the new method (or application) clearly explained?
No

Is the description of the method technically sound?
Partly

Are sufficient details provided to allow replication of the method development and its use by others?
No

If any results are presented, are all the source data underlying the results available to ensure full reproducibility?
Yes

Are the conclusions about the method and its performance adequately supported by the findings presented in the article?
Partly

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: microbial ecology, metabarcoding sequencing, NGS data analysis, bacterial communities, fungal communities

I confirm that I have read this submission and believe that I have an appropriate level of expertise to state that I do not consider it to be of an acceptable scientific standard, for
reasons outlined above.

Reviewer Report 27 November 2018

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Alejandro Sanchez-Flores

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The authors propose and evaluate a whole metagenome shotgun analysis via a low sequencing yield approach, using the Illumina platform.

In general, the idea and hypothesis are good, but the experimental design itself lacks important controls and there are many variables that are not analyzed and that can potentially bias the results.

My main concern is that the used samples have many variables and despite using a "replicate" for each case, samples within the same type were very different. Also the nature of each sample could have an effect in the DNA isolation, in particular for the vaccine ones. Also, regarding the vaccines, it is not clear to me, if what they are looking for is DNA of potential contaminants, since all viruses in the vaccine are ssRNA. That would be my guess, but is not clear from the text.

The main problem is that to test the influence of the sequencing yield, it would be extremely important to know the initial DNA concentration of each organism in the sample. Therefore, a mock metagenome or controlled sample would be much better as a reference to compare real life cases. In real life cases, the presence of certain organisms detected by the presence of its DNA, is not necessarily an indicator of the availability of alive organisms. Depending on the case, the presence of just the organism DNA could be an indicator of contamination which in the case of vaccines could be really bad. However, in the case of food material, finding DNA of pathogens, has to be associated with microbiology tests. However, with low sequencing yield, is very probable that very DNA in low amounts will be missed, even if this is not changing diversity indexes such as Chao1 and Shannon.

Finally, the main difference where low yield has a significant impact can be observed in the fecal samples. This is expected since among all the tested samples, fecal ones are the most diverse and sub-sampling will really affect them as observed in Figure 3.

Since the composition of each sample is not known a priori, then there are some factors that can contribute to biases. As mentioned, the DNA concentration but also its integrity (fragmentation) will affect the library construction; the cited kit requires DNA amplification which will have a bias towards GC rich genomic regions; library size was not described and was not mentioned if the samples were pooled with other libraries with different insert sizes, which affect
not only the sequencing quality but the yield.

In terms of bioinformatics analysis, it will be required to put the parameters used for each program, in case someone wants to reproduce this. For Kraken2, it is important to know what is the kmer size to index the database. For MEGAHIT assembly it will be important to know the kmer and step sizes used. For the completeness assessment, the authors used BUSCO, but apparently they are using the whole assembly to assess the completeness. This is not correct, since they must first separate in bins which genomes they have really reconstructed and then they can assess the completeness of them. Probably they can report the an average completeness value for all the reconstructed genomes. By doing the binning they can have a better analysis of what was really reconstructed and how complete it was.

The use of Krona in Figure 2 is not very convenient. The whole point of a Krona graph is that is interactive. If authors want to provide the Krona data to be downloaded it would be possible and recommended. Having said that, I recommend to use bar plots to represent the relative abundance and composition of the samples at a given taxa level.

Again, the idea is very good but the work needs to be improved before indexing.

**Is the rationale for developing the new method (or application) clearly explained?**
Yes

**Is the description of the method technically sound?**
No

**Are sufficient details provided to allow replication of the method development and its use by others?**
Partly

**If any results are presented, are all the source data underlying the results available to ensure full reproducibility?**
Yes

**Are the conclusions about the method and its performance adequately supported by the findings presented in the article?**
No

*Competing Interests:* No competing interests were disclosed.

*Reviewer Expertise:* Genomics, Transcriptomics, Metagenomics, Bioinformatics

I confirm that I have read this submission and believe that I have an appropriate level of expertise to state that I do not consider it to be of an acceptable scientific standard, for reasons outlined above.
We are grateful for the constructive comments. We agree with all of them and we are planning corrective actions, listed below.

My main concern is that the used samples have many variables and despite using a "replicate" for each case, samples within the same type were very different.

The observation is correct. Actually, the diversity of the samples was sought by purpose in order to be able to generalize the conclusions of our paper. The fact that diversity estimate and species abundance estimation remain reliable even with strong down-sampling for all of the samples is encouraging us to think that this is a general (although not necessarily universal) observation. The same is true for the observation that de-novo assembly quickly loses accuracy when decreasing the number of sequenced reads. Maybe this wasn't made clear enough in the paper, and we will clarify it.

Also the nature of each sample could have an effect in the DNA isolation, in particular for the vaccine ones.

Quantities of DNA isolated from vaccine samples (B1 and B2) were estimated to be ~2 µg using Qbit fluorimeter. However, we will provide a table with all the details about quantity, concentration, quality and size of starting DNA for all samples used in the study.

Also, regarding the vaccines, it is not clear to me, if what they are looking for is DNA of potential contaminants, since all viruses in the vaccine are ssRNA. That would be my guess, but is not clear from the text.

The vaccine composition declared by the producer is the following:
Live attenuated viruses: Measles (ssRNA) Swartz strain, cultured in embryo chicken cell cultures; Mumps (ssRNA) strain RIT 4385, derived from the Jeryl Linn strain, cultured in embryo chicken cell cultures; Rubella (ssRNA) Wistar RA 27/3 strain, grown in human diploid cells (MRC-5); Varicella (dsDNA) OKA strain grown in human diploid cells (MRC-5).
By DNA-seq we expected to find Varicella (dsDNA) OKA strain DNA (which was found and confirmed by variant analysis with respect to AB097932.1 Human herpesvirus 3 DNA, sub strain vOka). In addition, we found also human and chicken DNA. For human's, we confirmed MRC-5 cell origin by mitochondrial genome variant analysis.
Genotyping analyses gave us confidence on the validity of the obtained results, even though they were beyond the scope of this work.
To identify vaccine's ssRNA viruses we extracted RNA and performed RNA-seq from the same B1 and B2 samples. This aspect also goes beyond the scope of this work.

The main problem is that to test the influence of the sequencing yield, it would be extremely important to know the initial DNA concentration of each organism in the sample. Therefore, a mock metagenome or controlled sample would be much better as a reference to compare real life cases.

A mock community experiment is already on-going by using '10 Strain Staggered Mix
Genomic Material (ATCC® MSA-1001™). Of course, the data obtained will be integrated in the analysis results.

In real life cases, the presence of certain organisms detected by the presence of its DNA, is not necessarily an indicator of the availability of alive organisms. Depending on the case, the presence of just the organism DNA could be an indicator of contamination which in the case of vaccines could be really bad. However, in the case of food material, finding DNA of pathogens, has to be associated with microbiology tests.

We agree with the observation of the reviewer. However, the aim of this work is to determine if low-pass whole genome sequencing can be an appropriate approach to broadly describe a complex matrix; finding and confirming contaminants in vaccines or DNA pathogens in food samples was beyond of the scope of the paper.

However, with low sequencing yield, is very probable that very DNA in low amounts will be missed, even if this is not changing diversity indexes such as Chao1 and Shannon. Finally, the main difference where low yield has a significant impact can be observed in the fecal samples. This is expected since among all the tested samples, fecal ones are the most diverse and sub-sampling will really affect them as observed in Figure 3.

We agree with the reviewer; we add some thoughts just to clarify. We indeed observed that extremely rare species (with frequencies lower than 1/10000) are lost when subsampling to the most extreme levels. When subsampling to 100K reads we are losing species with a frequency around 1/100,000 (very approximate estimate). However, the effect of losing such species on the global sample diversity as estimated by Shannon diversity index is negligible (see Figure 4, in which we show that reduction in sequencing depth has no dramatic effect on Shannon's diversity index). The situation is different for the Chao 1 estimator. This is expected and is due to the way Chao1 is computed: this estimator relies heavily on the number of singletons (i.e. species represented by only one read). By subsampling, singletons (i.e. the rarest species) are very likely to be lost. The same phenomenon can be inferred by looking at Figures 5 and 6. Those represent a scatterplot of the relative abundance of species in full sample and reduce samples (100K and 10k reads, respectively). The plots are shown in log log scale to emphasize differences for low-frequency species. Only low-frequency species have some variation in frequency estimation. However, even when sampling only 10K read, species with frequency around 0.1% (i.e. 1/1000) are appropriately quantified. All of these observations led us to conclude that coverage reduction doesn't prevent a satisfactory characterization of complex matrices (with the only exception of Chao 1 estimator).

Since the composition of each sample is not known a priori, then there are some factors that can contribute to biases. As mentioned, the DNA concentration but also its integrity (fragmentation) will affect the library construction; the cited kit requires DNA amplification which will have a bias towards GC rich genomic regions; library size was not described.

The Nugen Ovation® Ultralow System V4 kit used is a standard kit for NGS library preparation (https://www.nugen.com/sites/default/files/DS_v2-Ovation_Ultralow_V2.pdf).
It is a standard protocol widely used by the scientific community to perform DNA-seq also from low input DNA quantities (1 ng), even if in our case input DNA was of moderate quantity. Mock community experiment will shed light on eventual biases. DNA concentration and integrity as well as input DNA quantities used in library construction and libraries insert size will be reported in the version 2 of the paper.

*It was not mentioned if the samples were pooled with other libraries with different insert sizes, which affect not only the sequencing quality but the yield.*

Samples were sequenced in different runs and pooled with other libraries of similar insert sizes. The number of reads obtained per sample reflects and respects their quantities, *i.e.* nmols that were loaded on the sequencer.

*In terms of bioinformatics analysis, it will be required to put the parameters used for each program, in case someone wants to reproduce this. For Kraken2, it is important to know what is the kmer size to index the database. For MEGAHIT assembly it will be important to know the kmer and step sizes used.*

All these details will be provided in the version 2 of the paper.

*For the completeness assessment, the authors used BUSCO, but apparently they are using the whole assembly to assess the completeness. This is not correct, since they must first separate in bins which genomes they have really reconstructed and then they can assess the completeness of them. Probably they can report an average completeness value for all the reconstructed genomes. By doing the binning they can have a better analysis of what was really reconstructed and how complete it was.*

This is a good point. While our aim was to estimate the total proportion of BUSCO genes that were reconstructed, irrespective of the species of the organism to which they belong, we understand that a practical application is likely to require separating the reconstructed genomes. We will integrate our analysis by binning the reconstructed genomes.

*The use of Krona in Figure 2 is not very convenient. The whole point of a Krona graph is that is interactive. If authors want to provide the Krona data to be downloaded it would be possible and recommended. Having said that, I recommend to use bar plots to represent the relative abundance and composition of the samples at a given taxa level.*

We will either provide a link to interactive krona graphs and/or bar plots reporting the relative abundance and composition of the samples.

*Competing Interests:* No competing interests were disclosed
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